The Maryland Medical Protocols for Emergency Medical Services

Maryland Institute for Emergency Medical Services Systems

Effective July 1, 2022
The complete “Maryland Medical Protocols for Emergency Medical Services” is also available on the Internet at www.MIEMSS.org. Protocols are occasionally amended during the year. Please check the MIEMSS website to be sure you have the most up-to-date version. The edition date appears on the lower portion of the page.
Welcome Letter

To All Maryland EMS Clinicians:

Re: The Maryland Medical Protocols for Emergency Medical Services – 2022 additions, updates and revisions

EMS clinicians will be able to download the full protocol document from the MIEMSS website at www.miemss.org and will receive a single copy of the 2022 pocket protocols.

The EMS Board has approved these protocols for implementation on July 1, 2022. Prior to July 1, all EMS clinicians must complete the Maryland EMS Update: 2022, which will highlight the new material. The protocol update will be available for continuing education credit on the MIEMSS Online Training Center (www.emsonlinetraining.org) in May, 2022.

With multi-disciplined input, we have updated the protocol content to reflect best practices and evidence-based medicine. The list of these changes is for reference only, and the information contained in the full protocol book is the official medical reference for EMS clinicians in Maryland.

A summary of significant protocol changes:

- **Acetaminophen:** All patients ages 3 months and older may now receive acetaminophen for treatment of fever (EMS-documented temperature of 100.4 or greater) or mild to moderate pain.
- **Cardiac Arrest (Adult):** The Adult PEA/Asystole algorithm has been refined to provide focused guidance on the management of patients in asystole, PEA with a narrow QRS, and PEA with a wide QRS. Calcium chloride and sodium bicarbonate should only be given for PEA with a wide QRS. The medical consultation requirement for sodium bicarbonate administration in adults has been removed.
- **Critically Unstable Patient (Adult):** A critically unstable patient protocol has been added to the General Patient Care section. This protocol emphasizes resuscitation of unstable patients on-scene prior to initiating transport.
- **Direct to Triage:** This protocol defines stable patients who may be suitable for direct transport to triage upon arrival to the receiving emergency department.
- **Droperidol:** This medication replaces haloperidol for treatment of patients who are 13 years of age or older presenting with moderate agitation.
- **Extraglottic Airways:** This addition to the Procedures section allows for the use of extraglottic/supraglottic airways including the King LTS-D, LMA, Air-Q, and iGel. An EMS operational program must select one of these airway devices and carry sizes appropriate for all patients from newborn through adult. These devices serve a critically important role in assisting ventilation and oxygenation to patients with compromised airways.
- **Induced Hypothermia:** Based on recent high-level scientific evidence, induced hypothermia has been removed.
- **Lateral Uterine Displacement:** For pregnant patients with hypotension or cardiac arrest, specific guidance has been added to provide left lateral uterine displacement with a goal of increasing venous return.
- **Tranexamic Acid (TXA):** TXA has been added to the protocols for treatment of traumatic hemorrhagic shock in patients 15 years of age and older.
- **Ventricular Assist Device (VAD):** Specific guidance has been added for the management of unstable patients with VADs.
- **Optional Supplemental Protocol:** Hydrofluoric Acid: Due to increased risk of HF exposure from electric car batteries, this OSP has been added to allow for calcium gluconate via IV, topical and nebulized routes. Interested jurisdictions should submit an OSP application to the State EMS Medical Director.

Timothy Chizmar, MD, FACEP, FAEMS
State EMS Medical Director

Theodore R. Delbridge, MD, MPH, FACEP, FAEMS
Executive Director

www.miemss.org
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A. GENERAL PROVISIONS

The goal of prehospital emergency medical services is to deliver a viable patient to appropriate definitive care as soon as possible. Optimal prehospital care results from a combination of careful patient assessment, essential prehospital emergency medical services, and appropriate medical consultation.

The Maryland Medical Protocols were developed to standardize the emergency patient care that EMS clinicians, through medical consultation, deliver at the scene of illness or injury and while transporting the patient to the closest appropriate hospital. These protocols will help EMS clinicians anticipate and be better prepared to give the emergency patient care ordered during the medical consultation.

Maryland has highly trained and dedicated basic and advanced life support personnel who may need online medical consultation only for complicated or extended resuscitative patient care. These protocols are a form of “standing orders” for emergency patient care intervention in a patient who has a life-threatening illness or injury. It remains the responsibility of the EMT, CRT-(I), or paramedic to obtain online medical consultation when appropriate. If it is genuinely impossible or inappropriate (i.e., when rendering emergency care to a patient who has a life-threatening injury or medical condition) to obtain online medical consultation, the EMT/CRT-(I)/paramedic may render emergency patient care in accordance with these protocols in an effort to save a patient’s life or limb. Whenever such emergency lifesaving patient care is rendered, the EMT/CRT-(I)/paramedic must document the treatment rendered and the reason online medical consultation could not be obtained on the Patient Care Report (PCR) and on an additional narrative. In addition, the “exceptional call” area on the PCR must be marked, and the clinician must immediately notify the EMS Jurisdiction. The EMS Jurisdiction must notify the State EMS Medical Director within five (5) days of the incident. This general provision applies throughout these protocols.

Requests for additions, deletions, or exceptions must be submitted through the State EMS Medical Director’s Office of the Maryland Institute for Emergency Medical Services Systems.

Unless otherwise specified, a mandate with a stated year but no date shall be interpreted as taking effect on the protocol implementation date for that year.

THE GENERAL PATIENT CARE SECTION AND THE ALGORITHMS MUST BE FOLLOWED IN THE SPECIFIC SEQUENCE NOTED.

FOR ALL OTHER TREATMENT PROTOCOLS, THE OUTLINE FORMAT IS STRICTLY FOR RAPID AND UNIFORM REFERENCE AND DOES NOT IMPLY OR DIRECT A MANDATORY SEQUENCE FOR PATIENT CARE.

THE GENERAL PATIENT CARE SECTION SHALL APPLY TO ALL PATIENT ENCOUNTERS UNLESS OTHERWISE NOTED IN ANY SPECIFIC TREATMENT PROTOCOL.
1.1 GENERAL PROVISIONS (continued)

IF AN EMERGENCY MEDICAL RESPONDER IS DISPATCHED AS AN EMS UNIT, OR FOR PURPOSES RELATED TO MEDICAL ASSISTANCE, OXYGEN AND AED TREATMENT MAY BE UTILIZED, WHEN APPROPRIATE AND APPLICABLE, PROVIDED THE EMERGENCY MEDICAL RESPONDER IS JURISDICTIONALLY AUTHORIZED TO USE AN AED AND/OR THE EMERGENCY MEDICAL RESPONDER HAS BEEN EDUCATED AND TRAINED TO PROVIDE OXYGEN AND/OR AED THERAPY.

THE EMERGENCY MEDICAL RESPONDER SHALL DOCUMENT ALL PATIENT CARE.
## General Information – IMPORTANT NUMBERS

### B. IMPORTANT NUMBERS

1. Commercial Ambulance Licensing and Regulation  
   Office: (410) 706-8511  
   Fax: (410) 706-8552

2. Critical Incident Stress Management  
   (800) 648-3001

3. Office of Licensure and Certification  
   Office: (800) 762-7157  
   Fax: (410) 706-2367

4. Regional Programs  
   a) Region I ( Allegany and Garrett Counties)  
      Office: (301) 895-5934  
      Fax: (301) 687-0129
   
   b) Region II (Washington and Frederick Counties)  
      Office: (301) 791-2366  
      Fax: (301) 791-9231
   
   c) Region III (Baltimore City, Anne Arundel,  
      Baltimore, Carroll, Harford, and Howard Counties)  
      Office: (410) 706-3996  
      Fax: (410) 706-8530
   
   d) Region IV (Caroline, Cecil, Dorchester, Kent,  
      Queen Anne’s, Somerset, Talbot, Wicomico,  
      and Worcester Counties)  
      Office: (410) 822-1799  
      Fax: (410) 822-0861
   
   e) Region V (Calvert, Charles, Montgomery,  
      Prince George’s, and St. Mary’s Counties)  
      Office: (301) 474-1485  
      Fax: (301) 513-5941

5. State EMS Medical Director  
   Office: (410) 706-0880  
   Fax: (410) 706-0880

6. SYSCOM (Administrative)  
   (800) 648-3001

7. EMRC  
   a) Consult Line (Region I)  
      (301) 722-0494  
   b) Consult Line (Region III)  
      (800) 492-3805  
   c) Consult Line (Region IV)  
      (877) 963-6963  
   d) Consult Line (Region V)  
      (877) 840-4245
POISON INFORMATION CENTER RECOMMENDATIONS SHOULD BE SOLICITED IN CONJUNCTION WITH MEDICAL CONSULTATION, BUT MEDICATION ORDERS CAN ONLY BE ACCEPTED FROM AN APPROVED BASE STATION.

1. Poison Control Centers
   a) Maryland Poison Center/University of Maryland School of Pharmacy, Baltimore (800) 222-1222
   b) National Capital Poison Center, Washington, DC (800) 222-1222

2. In-Patient Hospice Facilities
   a) Gilchrist Center–Towson (443) 849-8200
   b) Gilchrist Center Baltimore–Joseph Richey House (410) 523-2150
   c) Stella Maris Hospice (410) 560-9695
   d) Casey House (Montgomery/PG Hospice) (240) 631-6800

3. Tissue Donation
   a) Maryland Donor Referral Line Living Legacy (Majority of Maryland) (800) 923-1133
   b) Washington Regional Transplant Community (Charles, Montgomery, and Prince George’s Counties) (703) 641-0100
C. MARYLAND TRAUMA AND SPECIALTY REFERRAL CENTERS

Adult Trauma Centers

**Primary Adult Resource Center**
- R Adams Cowley Shock Trauma Center (UM), Baltimore

**Level I**
- The Johns Hopkins Hospital Adult Trauma Center (JHM), Baltimore

**Level II**
- Capital Region Medical Center (UM), Largo
- Johns Hopkins Bayview Medical Center (JHM), Baltimore
- Sinai Hospital of Baltimore (LifeBridge)
- Suburban Hospital (JHM), Bethesda

**Level III**
- Meritus Medical Center, Hagerstown
- Peninsula Regional (TidalHealth), Salisbury
- Western Maryland (UPMC), Cumberland

**Out-of-State Centers**
- Christiana Hospital (ChristianaCare), Newark, DE
- Washington Hospital Center (MedStar), Washington, DC

Specialty Referral Centers

**Eye Trauma**
- Wilmer Eye Institute/The Johns Hopkins Hospital (JHM), Baltimore

**Hand/Upper Extremity Trauma**
- The Curtis National Hand Center for Treatment of the Hand and Upper Extremity/Union Memorial Hospital (MedStar), Baltimore

**Hyperbaric Medicine**
- Center for Hyperbaric Medicine/R Adams Cowley Shock Trauma Center (UM), Baltimore

**Neurotrauma (Head and Spinal Cord Injuries)**
- Neurotrauma Center/R Adams Cowley Shock Trauma Center (UM), Baltimore

**Pediatric Trauma**
- Johns Hopkins Children’s Center (JHM), Baltimore
- Children’s National Medical Center, Washington, DC

**Burns**
- Adult Burn Center/Johns Hopkins Bayview Medical Center (JHM), Baltimore
- Adult Burn Center/Washington Hospital Center (MedStar), Washington, DC
- Pediatric Burn Center/Johns Hopkins Children’s Center (JHM), Baltimore
- Pediatric Burn Center/Children’s National Medical Center, Washington, DC

Health Care Facility Codes may be found online at www.MIEMSS.org, under the Protocols tab.
### Specialty Referral Centers

#### Perinatal Referral Centers
- Anne Arundel Medical Center (Luminis), Annapolis
- Franklin Square Medical Center (MedStar), Baltimore
- Frederick Health Hospital, Frederick
- Greater Baltimore Medical Center, Towson
- Holy Cross Hospital, Silver Spring
- Howard County General Hospital (JHM), Columbia
- Johns Hopkins Bayview Medical Center (JHM), Baltimore
- Mercy Medical Center, Baltimore
- Saint Agnes Hospital (Ascension), Baltimore
- Shady Grove Medical Center (Adventist), Rockville
- Sinai Hospital of Baltimore (LifeBridge)
- The Johns Hopkins Hospital (JHM), Baltimore
- University of Maryland Medical Center (UM), Baltimore

#### Primary Stroke Centers
- Anne Arundel Medical Center (Luminis), Annapolis
- Atlantic General Hospital, Berlin
- Baltimore Washington Medical Center (UM), Glen Burnie
- CalvertHealth Medical Center, Prince Frederick
- Capital Region Medical Center (UM), Largo
- Carroll Hospital Center (LifeBridge), Westminster
- Charles Regional Medical Center (UM), La Plata
- Doctor's Community Hospital (Luminis), Lanham
- Frederick Health Hospital, Frederick
- Good Samaritan Hospital (MedStar), Baltimore
- Greater Baltimore Medical Center, Towson
- Harbor Hospital (MedStar), Baltimore
- Harford Memorial Hospital (UMUCH), Havre De Grace
- Holy Cross Hospital, Germantown
- Holy Cross Hospital, Silver Spring
- Howard County General Hospital (JHM), Columbia
- Mercy Medical Center, Baltimore
- Meritus Medical Center, Hagerstown
- Midtown Campus (UM), Baltimore
- Montgomery Medical Center (MedStar), Olney
- Northwest Hospital (LifeBridge), Baltimore
- Peninsula Regional (TidalHealth), Salisbury
- Saint Agnes Hospital (Ascension), Baltimore
- Saint Joseph Medical Center (UM), Baltimore
- Saint Mary’s Hospital (MedStar), Leonardtown
- Shore Medical Center at Easton (UMSRH)
- Southern Maryland Hospital (MedStar), Clinton
- Union Hospital (ChristianaCare), Elkton
### General Information – MARYLAND TRAUMA AND SPECIALTY REFERRAL CENTERS (continued)

<table>
<thead>
<tr>
<th>Primary Stroke (continued)</th>
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<tbody>
<tr>
<td>• Union Memorial Hospital (MedStar), Baltimore</td>
</tr>
<tr>
<td>• Western Maryland (UPMC), Cumberland</td>
</tr>
<tr>
<td>• White Oak Medical Center (Adventist), Silver Spring</td>
</tr>
<tr>
<td>• Upper Chesapeake Medical Center (UMUCH), Bel Air</td>
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</table>

<table>
<thead>
<tr>
<th>Primary Stroke, Thrombectomy-Capable</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Franklin Square Medical Center (MedStar), Baltimore</td>
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<tr>
<td>• Shady Grove Medical Center (Adventist), Rockville</td>
</tr>
<tr>
<td>• Sinai Hospital of Baltimore (LifeBridge)</td>
</tr>
<tr>
<td>• Suburban Hospital (JHM), Bethesda</td>
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</table>

<table>
<thead>
<tr>
<th>Comprehensive Stroke</th>
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<tbody>
<tr>
<td>• Johns Hopkins Bayview Medical Center (JHM), Baltimore</td>
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<tr>
<td>• The Johns Hopkins Hospital (JHM), Baltimore</td>
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<tr>
<td>• University of Maryland Medical Center (UM), Baltimore</td>
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<table>
<thead>
<tr>
<th>Cardiac Interventional</th>
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<td>• Anne Arundel Medical Center (Luminis), Annapolis</td>
</tr>
<tr>
<td>• Baltimore Washington Medical Center (UM), Glen Burnie</td>
</tr>
<tr>
<td>• Capital Region Medical Center (UM), Largo</td>
</tr>
<tr>
<td>• Carroll Hospital Center (LifeBridge), Westminster</td>
</tr>
<tr>
<td>• Christiana Hospital (ChristianaCare), Newark, DE</td>
</tr>
<tr>
<td>• Franklin Square Medical Center (MedStar), Baltimore</td>
</tr>
<tr>
<td>• Frederick Health Hospital, Frederick</td>
</tr>
<tr>
<td>• Holy Cross Hospital, Silver Spring</td>
</tr>
<tr>
<td>• Howard County General Hospital (JHM), Columbia</td>
</tr>
<tr>
<td>• Johns Hopkins Bayview Medical Center (JHM), Baltimore</td>
</tr>
<tr>
<td>• Kent General (Bayhealth), Dover, DE</td>
</tr>
<tr>
<td>• Meritus Medical Center, Hagerstown</td>
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<tr>
<td>• Nanticoke (TidalHealth), Seaford, DE</td>
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<tr>
<td>• Peninsula Regional (TidalHealth), Salisbury</td>
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<tr>
<td>• Saint Agnes Hospital (Ascension), Baltimore</td>
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<tr>
<td>• Saint Joseph Medical Center (UM), Baltimore</td>
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<td>• Shady Grove Medical Center (Adventist), Rockville</td>
</tr>
<tr>
<td>• Shore Medical Center at Easton (UM)</td>
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<tr>
<td>• Sinai Hospital of Baltimore (LifeBridge)</td>
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<td>• Suburban Hospital (JHM), Bethesda</td>
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<tr>
<td>• The Johns Hopkins Hospital (JHM), Baltimore</td>
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<td>• Union Memorial Hospital (MedStar), Baltimore</td>
</tr>
<tr>
<td>• University of Maryland Medical Center (UM), Baltimore</td>
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<tr>
<td>• Washington Hospital Center (MedStar), Washington, DC</td>
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</tr>
<tr>
<td>• Upper Chesapeake Medical Center (UMUCH), Bel Air</td>
</tr>
</tbody>
</table>
Maryland Sexual Assault Forensic Examination (SAFE) Hospitals
SAFE hospital programs recognized by the Maryland Coalition Against Sexual Assault (MCASA)

• Anne Arundel Medical Center (Luminis) (Adult)
• Baltimore Washington Medical Center (UM) (Pediatric and Adult)
• CalvertHealth Hospital (Adult)
• Capital Region Medical Center (UM) (Pediatric and Adult)
• Carroll Hospital Center (LifeBridge) (Pediatric and Adult)
• Charles Regional Medical Center (UM) (Pediatric and Adult)
• Chestertown Medical Center (UMSRH) (Adult)
• Dorchester Medical Center (UMSRH) (Pediatric and Adult)
• Easton Medical Center (UMSRH) (Pediatric and Adult)
• Frederick Health Hospital (Pediatric and Adult)
• Garrett Regional Medical Center (WVU) (Pediatric and Adult)
• Greater Baltimore Medical Center (Pediatric and Adult)
• Harford Memorial Hospital (UMUCH) (Adult)
• Howard County General Hospital (JHM) (Pediatric and Adult)
• Mercy Medical Center (Adult)
• Meritus Medical Center (Pediatric and Adult)
• Peninsula Regional (TidalHealth) (Pediatric and Adult)
• Saint Mary’s Hospital (MedStar) (Pediatric and Adult)
• Shady Grove Medical Center (Adventist) (Pediatric and Adult)
• Union Hospital (ChristianaCare) (Adult)
• University of Maryland Medical Center (UM) (Pediatric)
• Western Maryland (UPMC) (Pediatric and Adult)

Maryland Emerging Infectious Disease (EID) Treatment and Assessment Hospitals
EID hospitals are reviewed, approved, and recognized as such by the Maryland Department of Health Infectious Disease Epidemiology and Outbreak Response Bureau.

Maryland Primary Ebola/Special Pathogen Treatment Hospital
• The Johns Hopkins Hospital (JHM)

Maryland Alternate Ebola/Special Pathogen Treatment Hospital
• University of Maryland Medical Center (UM)

Maryland Ebola/Special Pathogen Assessment Hospitals
• Anne Arundel Medical Center (Luminis)
• Frederick Health Hospital
• Holy Cross Hospital
• Peninsula Regional (TidalHealth)
• Southern Maryland Hospital Center (MedStar)
D. PROTOCOL KEY

1. Basic Life Support Level Care

2. Advanced Life Support Level Care

3. Requires Medical Consultation

4. Pediatric Care
   NOTE: ALL CLINICIANS (BLS and ALS) SHOULD CHECK ALL PEDIATRIC SECTIONS FOR NECESSARY CARE.

<table>
<thead>
<tr>
<th>Description</th>
<th>Age</th>
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<tbody>
<tr>
<td>Newly Born</td>
<td>Up to 1 hour</td>
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<tr>
<td>Neonate</td>
<td>1 hour to 28 days</td>
</tr>
<tr>
<td>Infant</td>
<td>&gt; 28 days to 1 year</td>
</tr>
<tr>
<td>Toddler</td>
<td>1 to &lt; 2 years</td>
</tr>
<tr>
<td>Preschooler</td>
<td>2 to 4 years</td>
</tr>
<tr>
<td>School-Age</td>
<td>5 to 12 years</td>
</tr>
<tr>
<td>Adolescent</td>
<td>13 to 18th birthday</td>
</tr>
</tbody>
</table>

5. Caution/Warning/Alert
E. PROTOCOL USAGE FLOW DIAGRAM

Response

Scene Arrival + Size Up

Personal Protective Equipment

Patient Approach

Initial Assessment

History + Physical Exam

Withhold Resuscitation

YES

OPTION A/B

Palliative Care Protocol

NO

DNR/MOLST

Detailed + Ongoing Assessment

Assign Clinical Priority

Determine and Provide Care According to Treatment Protocol

Disposition: Determine Receiving Facility + Mode of Transportation

Transport the Patient when Appropriate

Communications: Consult / Notify Receiving Facility

Transfer of Care / Rendezvous: Transfer Patient to Receiving Facility

Complete Documentation

YES

NO

Termination of Resuscitation Efforts

LEGEND

General Patient Care Section
Refer to Specific Protocols
F. PROTOCOL VARIATION PROCEDURE

If an error or variance occurs (i.e., any act or failure to act, in practice or judgment, involving patient care that is not consistent with established protocol, whether or not it results in any change in the patient’s status or condition):

1. The EMS clinician must:
   a) Notify the consulting physician via radio as soon as the error or variance is discovered, if prior to arrival at the receiving hospital,
   b) Monitor the patient’s condition very closely for any changes,
   c) Notify the receiving physician upon arrival, and
   d) Notify the local EMS jurisdiction or licensed commercial ambulance service and Program Medical Director within 24 hours of the incident.

2. The EMS Operational Program Quality Assurance Officer, in accordance with COMAR 30.03.04.02 B(6), must:
   a) Within 5 days of being made aware of the incident, submit written notification of the incident to the:
      (1) Local EMS jurisdiction,
      (2) Program Medical Director,
      (3) MIEMSS Office of Integrity, and
      (4) State EMS Medical Director.
   b) Within 14 days of the written notification of the incident, initiate a Medical Review Committee QA investigation.
   c) Within 30 days of the written notification of the incident, forward to MIEMSS' Office of Integrity and State EMS Medical Director the written results of the Medical Review Committee QA investigation and recommendations.
G. INABILITY TO CARRY OUT PHYSICIAN ORDER

Occasionally, a situation may arise in which a physician’s order cannot be carried out (e.g., the clinician feels the administration of an ordered medication would endanger the patient, a medication is not available, or a physician’s order is outside the protocol). If this occurs:

1. The EMS clinician must:
   a) Immediately notify the consulting physician as to the reason the order cannot be carried out.
   b) Document on the patient care report what was ordered, the time it was ordered, and the reason the order could not be carried out.
   c) As soon as practical following the call, notify the local EMS jurisdiction of the incident.

2. Public Service EMS Operational Programs must:
   a) Within **5 days** of being made aware of the incident, submit written notification of the incident through the local EMS jurisdiction and Program Medical Director to the Regional Medical Director with a copy to the State EMS Medical Director. The MIEMSS Regional EMS Coordinator shall be notified at the discretion of the Regional Medical Director.
   b) Within **14 days** of the written notification of the incident, initiate a QA investigation under the authority of the Medical Review Committee.
   c) Within **30 days** of the written notification of the incident, forward to the MIEMSS Office of Integrity and State EMS Medical Director written results of the Medical Review Committee QA investigation and recommendations.

3. Licensed Commercial Programs must:
   a) Within **5 days** of being made aware of the incident, submit written notification of the incident through the commercial Program Medical Director to the Director of the State Office of Commercial Ambulance Licensing and Regulation with a copy to the State EMS Medical Director.
   b) Within **14 days** of the written notification of the incident, initiate a QA investigation under the authority of the Medical Review Committee.
   c) Within **30 days** of the written notification of the incident, forward to the Program Medical Director and to the Director of the State Office of Commercial Ambulance Licensing and Regulation and State EMS Medical Director written results of the Medical Review Committee QA investigation and recommendations.
H. PHYSICIAN ORDERS FOR EXTRAORDINARY CARE NOT COVERED BY MARYLAND PROTOCOL

1. **ALL** of the following criteria MUST be present for EMS clinicians to proceed with an order under this section:

   a) During the consultation, both the consulting physician and the EMS clinician must agree that the patient’s condition and extraordinary care are not addressed elsewhere within these medical protocols and that the order is absolutely necessary to maintain the life of the patient.

   b) The EMS clinician must feel capable of correctly performing the care directed by the consulting physician, based on the instructions given by the consulting physician.

   c) The EMS clinician must inform the consulting physician of the effect of the treatment and notify the receiving physician of the treatment upon arrival at the hospital (if the receiving physician is different than the consulting physician).

   d) The EMS clinician must inform the EMS Operational Program Medical Director as soon as practical after the call.

2. If an EMS clinician receives an order for care that is not covered by Maryland protocols and does not feel comfortable with it or does not agree that it is absolutely necessary to maintain the life of the patient, they shall proceed with the “Inability to Carry Out Physician Order” section.

3. This extraordinary care protocol is intended to address the potential rare, unusual or unforeseen situations not specifically addressed within protocols. This extraordinary care protocol is neither a *carte blanche* for any and all actions nor a device to avoid or circumvent protocols. In all situations, emergency health care clinicians, both EMS and on-line physicians providing medical direction, are accountable for their actions in discharging their patient care responsibilities.

### EXTRAORDINARY CARE CHECKLIST

- Identify the need for extraordinary care with physician consult and EMS clinician acceptance.
  - Care is not covered elsewhere in the protocols.
  - Care is absolutely necessary to maintain the life of the patient.
- Immediately upon delivery of patient, EMS clinician must notify the receiving physician of the effect of treatment rendered.
- Notify the EMS Operational Program Medical Director as soon as practical after the call.
The General Patient Care section shall apply to all patient encounters unless otherwise noted in any specific treatment protocol.

A. RESPONSE
   Review the dispatch information and select appropriate response.

B. SCENE ARRIVAL AND SIZE-UP
   1. Consider Body Substance Isolation (BSI).
   2. Consider Personal Protective Equipment (PPE).
   3. Evaluate the scene safety.
   4. Determine the number of patients.
   5. Consider the need for additional resources.

C. PATIENT APPROACH
   1. Determine the Mechanism of Injury (MOI)/Nature of Illness (NOI).
   2. If appropriate, begin triage and initiate Mass Casualty Incident (MCI) procedures.

D. INITIAL ASSESSMENT
   Rapidly develop a general impression of the patient on first contact:
   1. Identify the critically unstable patient – any patient in extremis or with imminent risk for deterioration to arrest:
      a) New onset of altered mental status (AVPU not alert)
      b) Airway compromise
      c) Acute respiratory distress
      d) Signs of poor perfusion
      e) Any other patient judged by the clinician to be in extremis or at risk for deterioration to cardiac arrest
   2. If you have identified a critically unstable patient:
      a) STOP ALL MOVEMENT OF PATIENT
      b) DO NOT INITIATE TRANSPORT
      c) PROCEED TO CRITICALLY UNSTABLE PATIENT PROTOCOL IMMEDIATELY

For pediatric patients, use the Pediatric Assessment Triangle.

- Appearance
- Work of Breathing
- Circulation to Skin
3. Assess mental status
   a) Alert
   b) Responds to Verbal stimuli
   c) Responds to Painful stimuli
   d) Unresponsive

4. Airway
   a) Stabilize cervical spine when appropriate
   b) Open and establish airway using appropriate adjunct.
   c) Place patient in appropriate position.
   d) Suction airway as needed, including tracheostomy tubes.
   e) If a patent airway cannot be established, the patient must be transported to the closest appropriate hospital-based emergency department or designated free-standing emergency medical facility. EMS clinicians should remain available to assist with patient transfer, if the hospital determines such a transfer is appropriate.
   f) In infants and young children, inspiratory stridor is an indication of upper airway foreign body or partial airway obstruction. Request ALS rendezvous. Transport the patient rapidly and with caution. Have foreign body airway removal equipment ready for immediate use in case the patient's airway becomes obstructed.

5. Breathing
   a) Determine if breathing is adequate and assess oxygen saturation (SpO₂) with pulse oximeter.
      (1) If patient's ventilations are not adequate, provide assistance with 100% oxygen using Bag-Valve-Mask (BVM).
         (i) For all ages except neonates, deliver 1 breath every 5 seconds (8–12 breaths/min).
         (ii) For a neonate, deliver 1 breath every 3 seconds; higher rates may be required.
      (2) The decision to oxygenate will be based on the patient’s clinical condition.
         (i) If the patient has SpO₂ less than 94%, administer supplemental oxygen, titrated to SpO₂ level of 94%.
         (ii) Supplemental oxygen is not needed if SpO₂ greater than or equal to 94% unless the patient is in respiratory distress, acutely dyspneic, or suffering from suspected CO poisoning. Patients in severe respiratory distress may benefit from high-flow oxygen from a nonrebreather (NRB).
         (iii) Unless in respiratory distress, avoid administration of high-flow oxygen to patients presenting with the following conditions:
            (a) STEMI / angina
            (b) CVA / stroke
            (c) Post-arrest
         (iv) CO exposure: Apply 100% oxygen via NRB mask. Maintain SpO₂ at 100%.
      (3) Utilize continuous ETCO₂ waveform monitoring in all intubated patients.
      (4) Measure carbon monoxide level with a co-oximeter, if appropriate and available.
b) Hyperventilate the head-injured patient only if signs/symptoms of herniation are present, including posturing, loss of pupillary light response, dilation of one or both pupils, vomiting, hypertension, bradycardia, and/or irregular respirations.

(1) If hyperventilating, use the following rates:
   (i) Adult (including adolescent 13 years of age or older): 20 breaths per minute
   (ii) Child (1-12 years of age): 30 breaths per minute
   (iii) Infant (less than 1 year of age): 35 breaths per minute

(2) Use ETCO$_2$ monitoring.
   (i) Maintain ETCO$_2$ between 35-40 mmHg for any patient with significant head injury
   (ii) For patients with significant head injury and signs of herniation, adjust ventilations to achieve ETCO$_2$ of 30-35 mmHg.

6. Circulation
   a) Assess pulse.

   (1) Patients within the first hour after delivery, refer to Newly Born protocol.
   (2) Patients from one hour after birth up to those who have not reached their 13th birthday, refer to the Universal Algorithm for Pediatric Emergency Cardiac Care for BLS.
   (3) Patients 13 years of age or greater, refer to the Universal Algorithm for Adult Emergency Cardiac Care for BLS.
   (4) If pulseless, immediately initiate high-quality continuous CPR.
      (i) Ensure frequent clinician rotations and minimal interruptions (less than 10 seconds).
      (ii) Mechanical CPR devices may be used, if available, for patients 13 years of age and older only.
      (iii) Perform CPR while preparing for rhythm analysis and defibrillation.
b) Assess for and manage profuse bleeding, using a method appropriate for the patient’s injuries:
   (1) Direct pressure
   (2) Wound packing
   (3) Hemostatic gauze
   (4) Tourniquet or junctional tourniquet (with jurisdictional training)
c) Assess skin color, temperature, and capillary refill.

7. Disability
   a) Assess for pulse, motor and sensory function in all extremities
   b) Assess GCS for trauma patients
   c) Determine the need for Spinal Motion Restriction.
      (1) Patients who have a blunt trauma with a high-energy mechanism of injury that has potential to cause spinal cord injury or vertebral instability and one or more the following should receive spinal motion restriction.
          - Midline spinal pain, tenderness, or deformity
          - Signs and symptoms of new paraplegia or quadriplegia
          - Focal neurological deficit
          - Altered mental status or disorientation
          - Distracting injury: Any injury (e.g., fracture, chest, or abdominal trauma) associated with significant discomfort that could potentially distract from a patient’s ability to accurately discern or define spinal column pain or tenderness.
      (2) In addition to the above indicators for adults, the below apply to children who have not yet reached their 15th birthday.
          - Neck pain or torticollis
          - High-impact diving incident or high-risk motor vehicle crash (head on collision, rollover, ejected from the vehicle, death in the same crash, or speed greater than 55 mph)
General Patient Care (GPC) (continued)

- Substantial torso injury
- Conditions predisposing to spine injury

d) If NO to all of the above, transport as appropriate.
e) Infant or child car seats may not be used as a spinal immobilization device for the pediatric patient.
f) If patient is unable to communicate or appropriately respond to the above questions, apply Spinal Motion Restriction protocol.

8. Exposure
To assess patient’s injuries, remove clothing as necessary, considering condition and environment.

9. Assign Clinical Priority
   a) Priority 1 — Critically ill or injured person requiring immediate attention; unstable patients with life-threatening injury or illness.
   b) Priority 2 — Less serious condition yet potentially life-threatening injury or illness, requiring emergency medical attention but not immediately endangering the patient’s life.
   c) Priority 3 — Non-emergent condition, requiring medical attention but not on an emergency basis.
   d) Priority 4 — Does not require medical attention.
   e) In the event of a multiple casualty incident, the Simple Triage and Rapid Treatment (START and/or JumpSTART) technique will be instituted for rapid tagging and sorting of patients into priority categories for both treatment and transport.

10. Normal Vital Signs Chart

<table>
<thead>
<tr>
<th>AGE</th>
<th>ESTIMATED WEIGHT</th>
<th>HEART RATE</th>
<th>RESPIRATORY RATE</th>
<th>SYSTOLIC B/P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature</td>
<td>Less than 3 kg</td>
<td>160</td>
<td>Greater than 40</td>
<td>60</td>
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<tr>
<td>Newborn</td>
<td>3.5 kg</td>
<td>130</td>
<td>40</td>
<td>70</td>
</tr>
<tr>
<td>3 mo.</td>
<td>6 kg</td>
<td>130</td>
<td>30</td>
<td>90</td>
</tr>
<tr>
<td>6 mo.</td>
<td>8 kg</td>
<td>130</td>
<td>30</td>
<td>90</td>
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<tr>
<td>1 yr.</td>
<td>10 kg</td>
<td>120</td>
<td>26</td>
<td>90</td>
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<tr>
<td>2 yrs.</td>
<td>12 kg</td>
<td>115</td>
<td>26</td>
<td>90</td>
</tr>
<tr>
<td>3 yrs.</td>
<td>15 kg</td>
<td>110</td>
<td>24</td>
<td>90</td>
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<tr>
<td>4 yrs.</td>
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<td>100</td>
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<td>6 yrs.</td>
<td>20 kg</td>
<td>100</td>
<td>20</td>
<td>95</td>
</tr>
<tr>
<td>8 yrs.</td>
<td>25 kg</td>
<td>90</td>
<td>20</td>
<td>95</td>
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<tr>
<td>10 yrs.</td>
<td>35 kg</td>
<td>85</td>
<td>20</td>
<td>100</td>
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<tr>
<td>12 yrs.</td>
<td>40 kg</td>
<td>85</td>
<td>20</td>
<td>100</td>
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<td>14 yrs.</td>
<td>50 kg</td>
<td>80</td>
<td>18</td>
<td>110</td>
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<tr>
<td>ADULT</td>
<td>Greater than 50 kg</td>
<td>80</td>
<td>18</td>
<td>120</td>
</tr>
</tbody>
</table>
E. HISTORY AND PHYSICAL EXAMINATION/ASSESSMENT

1. Conduct a Focused Examination/Detailed Examination/Ongoing Assessment.

2. Collect and transport documentation related to patient’s history (example: Emergency Information Form, Medic Alert, EMS DNR/MOLST, or jurisdictional form).

3. Clinicians should obtain and document a contact telephone number for one or more individuals who have details about the patient’s medical history so that the physician may obtain and validate additional patient information.

4. Obtain an EKG when appropriate.

ALL HEALTH CARE CLINICIANS ARE OBLIGATED BY LAW TO REPORT CASES OF SUSPECTED CHILD OR VULNERABLE ADULT ABUSE AND/OR NEGLECT TO EITHER THE LOCAL POLICE OR ADULT/CHILD PROTECTIVE SERVICE AGENCIES. DO NOT INITIATE REPORT IN FRONT OF THE PATIENT, PARENT, OR CAREGIVER (MD CODE, FAMILY LAW, § 5-704). UNDER MARYLAND LAW, EMS CLINICIANS ARE PROTECTED FROM LIABILITY IF THEY MAKE A REPORT OF CHILD/VULNERABLE ADULT ABUSE AND NEGLECT IN GOOD FAITH (COURTS AND JUDICIAL PROCEEDINGS ARTICLE § 5-620).

F. TREATMENT PROTOCOLS

1. Refer to ALL appropriate protocols.

2. Patients who have had an impaled conducted electrical weapon used on them will be transported to the nearest appropriate facility without dart removal (exception: Tactical EMS). ANY conducted electrical weapon dart impalement to the head, neck, hands, feet, or genitalia must be stabilized in place and evaluated by a physician.

3. Clinicians may assist the patient or primary caregiver in administering the patient’s prescribed rescue medication.
   a) BLS clinicians may assist with the administration of the patient’s fast-acting bronchodilator MDI and sublingual nitroglycerin.
   b) ALS clinicians may administer the patient’s prescribed benzodiazepine for seizures, Factor VIII or IX for Hemophilia A or B, or reestablish IV access for continuation of an existing vasoactive medication.
   c) Clinicians should obtain on-line medical direction to administer other prescribed rescue medications not specifically mentioned in The Maryland Medical Protocols for Emergency Medical Services (e.g., hydrocortisone (Solucortef) for adrenal insufficiency). The rescue medication must be provided by the patient or caregiver and the label must have the patient’s name and the amount of medication to be given.

4. For patients with fever documented by EMS as greater than 100.4 F (38 C), clinicians may treat with acetaminophen.

DO NOT ADMINISTER ORAL MEDICATIONS (EXCEPT ORAL GLUCOSE) TO PATIENTS WITH AN ALTERED MENTAL STATUS.
5. For pediatric patients
   a) Pediatric section of the treatment protocol will be used for children who have not reached their 15th birthday (trauma) or their 18th birthday (medical), except as otherwise stated in the treatment protocol.
   b) Medication dosing
      (1) Pediatric doses apply to patients weighing less than 50 kg.
      (2) For pediatric patients equal to or greater than 50 kg, utilize adult dosing.
   c) The developmental age of the infant/child must be considered in the communication and evaluation for treatment.
   Destination consideration:
      For those patients who are 18 years of age or older who receive specialized care at a pediatric facility, consider medical consultation with a Pediatric Base Station for patient destination.
   d) Infants and children must be properly restrained prior to and during transport.
   e) A parent/guardian/care taker may remain with a pediatric patient during transport, but must be secured in a separate vehicle restraint system at all times during transport.
   f) For patients with fever documented by EMS as greater than 100.4 F (38 C), clinicians may treat with acetaminophen.
G. COMMUNICATIONS

1. Communications with and through EMRC/SYSCOM are recorded. In addition, as part of the quality assurance and quality improvement process, communications with hospitals are frequently recorded. Therefore, you should assume that all your communications among EMS clinicians, hospitals, public safety communications centers, and EMRC/SYSCOM are being recorded.

ANY PATIENT WHOM THE CLINICIAN IDENTIFIES AS MEETING ANY “SPECIALTY” ALERT (E.G., TRAUMA, STEMI ALERT, STROKE ALERT, SEPSIS ALERT) REQUIRES A HOSPITAL NOTIFICATION, AND WHEN INDICATED BY PRIORITY OR NEED FOR INTERVENTION WILL HAVE ONLINE MEDICAL CONSULTATION THROUGH EMRC ON A RECORDED LINE (RADIO OR PHONE).

2. All Priority 1 patients require online medical consultation through EMRC on a recorded line (radio or phone).

3. All Priority 2 patients who need further therapeutic intervention(s) that require on-line medical consultation approval shall perform on-line medical consultation through EMRC on a recorded line (radio or phone).

4. For Priority 2 patients who have persistent symptoms but who do not need therapeutic intervention(s) requiring on-line medical consultation approval, clinicians shall notify the receiving facility with an “information only call” through EMRC on a recorded line (radio or phone).

5. For Priority 2 patients whose symptoms have resolved and Priority 3 patients whose vital signs are within normal limits, notification may be made through EMRC on a recorded line (radio or phone) or through an EOC/EMS communication system in accordance with the standard operating procedures of the local jurisdiction.

ONLINE MEDICAL CONSULTATION MAY BE OBTAINED AT ANY TIME FOR ANY PATIENT, IF DESIRED BY THE PREHOSPITAL EMS CLINICIAN. PEDIATRIC AND SPECIALTY CONSULTATION IS ENCOURAGED FOR TRAUMA AND MEDICAL PATIENTS. CONSULTATION WITH PEDIATRIC AND SPECIALTY CENTERS SHALL OCCUR SIMULTANEOUSLY WITH A BASE STATION CONSULT.

6. If medical consultation is genuinely unavailable, or if the time necessary to initiate consultation significantly compromises patient care, the clinician shall proceed with additional protocol directed care, so long as transport will not be significantly delayed. “Exceptional Call” must be indicated on the Patient Care Report (PCR).

7. Core essentials for communications:
   a) Assigned patient priority (1 to 4)
   b) Age
   c) Chief complaint
   d) Clinician impression
   e) Pertinent patient signs and symptoms (e.g., HR, RR, BP, Pulse Ox, and GCS) (be specific—do not use within normal limits or stable in description)
   f) Pertinent physical findings
   g) ETA
General Patient Care (GPC) (continued)

In addition, for specialty center patients:

**Trauma**
- a) Patient Trauma Decision Tree Category (Alpha, Bravo, Charlie, Delta)
- b) Number of victims if more than one
- c) Describe mechanism

**Stroke**
- d) Last known well time
- e) Specific neurological findings (sensory, motor, cognitive)
- f) Upon positive assessment using the Cincinnati Stroke Scale, a STROKE alert shall be made and the LAMS score will be included in the consult.

**STEMI**
- g) 12-Lead interpretation
- h) Duration of symptoms

**CONSIDER ACTIVATION OF THE GO-TEAM FOR SERIOUSLY INJURED PATIENTS WHO REQUIRE A PROLONGED EXTRICATION AND WHO MEET THE INDICATIONS FOR GO-TEAM ACTIVATION.**

8. Mass Casualty Incident (MCI) Communications
   - a) When a local jurisdiction declares an MCI, it is extremely important to maximize patient care resources and reserve EMS communications for emergent situations. Except for extraordinary care interventions, EMS clinicians may perform all skills and administer medications within protocol during a declared MCI. When the MCI condition is instituted, the Exceptional Call box must be checked on the PCR.
   - b) During an MCI, the EMS Officer-in-Charge (OIC) shall designate an EMS Communicator who shall establish appropriate communications.
   - c) Reference the *Multiple Casualty Incident/Unusual Event* Protocol.

**H. REASSESSMENT**
1. Reassess unstable patients frequently (recommended every 5 minutes).
2. Reassess stable patients at a minimum of every 15 minutes.
3. Reassess patients being discharged to home or long-term care at the beginning and end of the transport or more frequently, at the clinician’s discretion.
I. DISPOSITION

1. Destination
   
a) Priority 1 patients shall be triaged according to Maryland Medical Protocols to the closest appropriate hospital-based emergency department, designated trauma, or designated specialty referral center. Critically unstable patients in need of immediate life-saving interventions that cannot be provided in the field shall, with the approval of EMS system medical consultation, be diverted to the closest facility (including freestanding emergency medical facility) capable of immediately providing those interventions.

b) Priority 2 patients shall be triaged according to the Maryland Medical Protocols to the closest appropriate hospital-based emergency department, designated trauma or designated specialty referral center unless otherwise directed by EMS system medical consultation. Stable Priority 2 patients may be referred to a freestanding emergency medical facility.

c) Stable Priority 3 or 4 patients who do not need a time-critical intervention may be transported to the local emergency department or freestanding emergency medical facility.

d) Patients Under Investigation (PUI) for an Emerging Infectious Disease (EID) at a residence should be transported directly to an Assessment Hospital unless total transport time is no longer than 45 minutes greater than transport to the nearest Frontline Hospital ED. If transport time is longer than 45 minutes greater than transport to the nearest Frontline Hospital ED, the patient must be transported to the closest appropriate Frontline hospital. Priority 1 and Priority 2 patients with unresolved symptoms that cannot be managed outside the hospital should be taken to the closest Frontline Hospital. Receiving hospital notification of all suspected PUI patients should be done as early as possible to allow for hospital staff to prepare. Helicopter transport is NOT indicated for the PUI patient.

e) For Priority 2 and Priority 3 patients not meeting a specialty center destination care protocol, the EMS clinician should ask if the patient has had a hospital admission (inpatient service) within the last 30 days. If the answer is yes, the EMS clinician should transport (repatriate) the patient to that hospital as long as that hospital is not more than 15 additional minutes further than nearest hospital (or greater if allowed for by the EMS Operational Program).

2. Mode of transport (air, land, water)
   
a) Medevac patients with indications for specialty referral center should be flown to the appropriate type of specialty center if not more than 10–15 minutes further than the closest trauma center. (Patients with an airway, breathing, or circulatory status who would be jeopardized by going an additional 10–15 minutes should go to the closest trauma center.)

b) Consider utilization of a helicopter when the patient’s condition warrants transport to a trauma or specialty referral center and the use of a helicopter would result in a clinically significant reduction in time compared with driving to a trauma/specialty center.

ALL REQUESTS FOR SCENE HELICOPTER TRANSPORTS SHALL BE MADE THROUGH SYSCOM. FOR TRAUMA DECISION TREE CATEGORY CHARLIE OR DELTA, RECEIVING TRAUMA CENTER MEDICAL CONSULTATION IS REQUIRED WHEN CONSIDERING WHETHER HELICOPTER TRANSPORT IS OF CLINICAL BENEFIT.
c) If the time of arrival at the trauma or specialty referral center via ground unit is less than 30 minutes, there will generally be no benefit in using the helicopter, especially for Trauma Decision Tree categories Charlie and Delta.

d) Refer to the Trauma Decision Tree when considering use of aeromedical transport. Provide SYSCOM with the patient’s category (Alpha, Bravo, Charlie, or Delta).

e) On-line medical direction should be obtained from the local trauma center and the specialty referral center when transport to the specialty center would require more than 10–15 minutes additional transport time.

(1) Pediatric Trauma Patients: Indications as per the pediatric section of the Trauma protocols.

(2) Spinal Trauma Patients: Indications as per Spinal Motion Restriction protocol.

(3) Burn Patients: Indications as per Burn protocol. Special note: Isolated burn patients without airway injury or other associated trauma should normally be flown to a burn center, regardless of the location of the closest trauma center.

(4) Hand Injury Patients: Indications as per Hand Trauma protocol. Special note: Medevac patients with appropriate indications for hand center referral should normally be flown to the hand center, regardless of the location of the closest trauma center.

3. Status

   Evaluate the need for emergent versus non-emergent transportation.

**DO NOT WAIT ON-SCENE FOR ADVANCED LIFE SUPPORT. ATTEMPT TO RENDEZVOUS EN ROUTE TO THE HOSPITAL.**
J. TRANSFER OF CARE/RENDEZVOUS AND TRANSITION OF PATIENT CARE ALS TO BLS

The ALS clinician-patient relationship is established when the ALS clinician initiates patient assessment and

1. ALS medication(s)* is/are administered or
2. ALS procedure(s)* is/are performed or
3. Upon ALS clinician assessment of the patient there is potential risk of deterioration.

* Based on the medication or procedure as listed in protocol 9.2: Procedures, Medical Devices, and Medications for EMS and Commercial Services.

ALS clinicians may only terminate their EMS clinician-patient relationship when they are assured that the patient will continue to receive care at the same or greater levels, or when they have documented with on-line medical direction that the patient’s condition has improved and that patient care may be transferred safely to an EMS clinician with a lower scope of practice.

BLS clinicians have the right to decline the transition of patient care. When consensus between the clinicians cannot be gained, ALS shall get on-line medical direction.

Clinicians will relay assessment findings and treatment provided to the individual(s) assuming responsibility for the patient(s). Should an ALS clinician perform an EKG (of any type), it shall be imported into the patient care report and a copy shall be sent with the BLS unit to the receiving facility.

K. DOCUMENTATION

A Patient Care Report (PCR) will be completed and delivered to the receiving facility as soon as possible, ideally upon transfer of care. If this is not immediately possible, clinicians must provide documentation of the patient's prehospital care on a template and in a format provided or approved by MIEMSS for inclusion in the patient care record before leaving the receiving facility, then deliver the completed PCR within 24 hours after dispatch, in compliance with COMAR 30.03.04.04.

Only the unit that pronounces death will select the “Dead on Scene” option in the PCR (eMEDS®) and thus all other units will report “Operational Support Only.” If no interventions are performed, the highest level EMS clinician on scene will pronounce death and document “Dead on Scene.” If BLS care was rendered by a BLS unit and then termination of resuscitation and pronouncement of death occurred, the BLS unit will select “Dead at Scene with BLS Intervention” option on the eMEDS® PCR. If ALS care was rendered by an ALS unit and then termination of resuscitation and pronouncement of death occurred, the ALS unit will select “Dead at Scene with ALS Intervention” option on the eMEDS® PCR.

L. CONFIDENTIALITY

Patient confidentiality must be maintained at all times.

M. PROFESSIONAL CONDUCT

All patients should be treated with dignity and respect in a calm and reassuring manner.
2.2 General Patient Care (GPC) – HISTORY AND PHYSICAL EXAMINATION

HISTORY AND PHYSICAL EXAMINATION

TRAUMA PATIENT

Significant MOI
Rapid Trauma Assessment
Head
Crepitation
Chest
Crepitation
Respiration
Paradoxical Motion
Breath Sounds
Abdomen
Rigidity
Distention
Pain on Motion
Blood, Urine, Feces
Extremities
Pulse/Motor/Sensory

D C A P B T L S

Baseline Vital Signs
Obtain SAMPLE History

Non-Significant MOI
Determine Chief Complaint
Perform Focused Examination of the Injured Site and Areas Compatible with Given MOI

D C A P B T L S

Baseline Vital Signs
Obtain SAMPLE History

MEDICAL PATIENT

Unresponsive Patient
Rapid Physical Examination
Head
JVD
Medical Alert Device
Blood, Urine, Feces
Extremities

D C A P B T L S

Baseline Vital Signs
Obtain SAMPLE History

Responsive Patient
Obtain History of Episode
Onset
Provocation
Quality
Radiation
Severity
Time

Baseline Vital Signs
Obtain SAMPLE History
Allergies
Medications
Pertinent History
Last Oral Intake
Events Prior

Focused Physical Exam DCAPBTLS
Check areas suggested by MOI and SAMPLE.

CONSIDER ALS, PERFORM INTERVENTIONS, AND TRANSPORT.
General Patient Care (GPC) – DETAILED AND ONGOING ASSESSMENTS

TRIUMA PATIENT
- Reassess AVPU
- Reassess Airway
- Monitor Breathing
- Reassess Circulation
- Monitor Skin
- Confirm Clinical Priority
- REPEAT INITIAL ASSESSMENT
- REPEAT & RECORD VITAL SIGNS
- REPEAT RAPID TRAUMA ASSESSMENT
- CHECK ALL INTERVENTIONS

MEDICAL PATIENT
- Reassess AVPU
- Reassess Airway
- Monitor Breathing
- Reassess Circulation
- Monitor Skin
- Confirm Clinical Priority
- REPEAT INITIAL ASSESSMENT
- REPEAT & RECORD VITAL SIGNS
- REPEAT FOCUSED ASSESSMENT
- CHECK ALL INTERVENTIONS

GENERAL EXAMINATION
- Scalp & Cranium
- Eyelids
- Discoloration
- Foreign Bodies
- Blood in Anterior Chamber
- Fluid Drainage or Bleeding
- Discoloration
- Mouth
- Teeth & Foreign Bodies
- Breathing Odor
- Discoloration
- Neck
- Jugular Vein Distention
- Trachea Position
- Crepitation
- Chest
- Paradoxical Motion
- Breath Sounds
- Crepitation
- Abdomen
- Pain on Motion
- Pelvis/GU
- Extremities
- Pulse, Motor, Sensory
- Capillary refill
- Posterior

CONSIDER ALS, PERFORM INTERVENTIONS, AND TRANSPORT.
2.3 General Patient Care (GPC) – START TRIAGE ALGORITHM

START Adult Triage

Able to walk?

Yes ➔ MINOR ➔ SECONDARY TRIAGE

No ➔

Spontaneous breathing

No ➔ Position airway ➔ APNEA ➔ EXPECTANT

Yes ➔

Respiratory Rate

>30 ➔ IMMEDIATE

<30 ➔

Perfusion

Radial pulse absent or capillary refill > 2 sec ➔ IMMEDIATE

Radial pulse present or capillary refill < 2 sec ➔

Mental status

Doesn’t obey commands ➔ IMMEDIATE

Obeys commands ➔ DELAYED

Source: U.S. National Library of Medicine
General Patient Care (GPC) – JumpSTART TRIAGE ALGORITHM 2.3

JumpSTART Pediatric Multiple Casualty Incident Triage

Able to walk?
Yes → MINOR → SECONDARY TRIAGE
No →

Spontaneous breathing
No → Position airway
APNEA →

Yes →

Palpable pulse?
Yes →

5 rescue breaths
Spontaneous breathing
APNEA → EXPECTANT

IMMEDIATE

Respiratory Rate
<15 or >45
IMMEDIATE

15-45

Palpable Pulse?
No → IMMEDIATE

Yes →

Neurological Assessment [AVPU]
Inappropriate "P" (e.g., posturing) or "U"
"A," "V," or Appropriate "P" (e.g., withdrawal from painful stimulus)
IMMEDIATE

DELAYED

Neurological Assessment

<p>| | |</p>
<table>
<thead>
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<tbody>
<tr>
<td>A</td>
<td>Alert</td>
</tr>
<tr>
<td>V</td>
<td>Responds to Verbal Stimuli</td>
</tr>
<tr>
<td>P</td>
<td>Responds to Painful Stimuli</td>
</tr>
<tr>
<td>U</td>
<td>Unresponsive to Noxious Stimuli</td>
</tr>
</tbody>
</table>

Use JumpSTART if the Patient appears to be a child.
Use an adult system, such as START, if the patient appears to be a young adult.

Source: U.S. National Library of Medicine
©Lou Romig MD, 2002

www.miemss.org
a) **INDICATIONS**

Adult patients (18 years of age or older) who are identified to be in extremis or are at risk for deterioration to cardiac arrest at any point during their care. These patients can include, but are not limited to, patients with:

1. New onset altered mental status (AVPU – not alert)
2. Airway compromise
3. Acute respiratory distress
4. Signs of poor perfusion
5. Any other patient judged by the clinician to be in extremis or at risk for deterioration to cardiac arrest

b) **BLS**

1. Cease all efforts at patient movement until treatments in this protocol are complete.
2. Obtain a complete patient assessment, including pulse oximetry.
3. Consider the need for more resources, if available, including multiple ALS clinicians.
5. Manage the patient’s airway and ventilation (e.g., BVM with or without OPA/NPA) as indicated and tolerated.
6. Treat hypoxia and respiratory distress aggressively.

c) **ALS**

1. Initiate ETCO₂ monitoring.
2. Obtain 12-lead EKG, if appropriate for patient condition.
3. Obtain vascular access and support perfusion with IV fluids and vasopressors as indicated.
4. Address any other life threats noted on physical exam.
5. **Continue General Patient Care, including transport.**
3.1-A  Adult Emergency Cardiac Care for BLS – Algorithm

UNIVERSAL ALGORITHM FOR ADULT EMERGENCY CARDIAC CARE FOR BLS

Unresponsive Not Breathing

Pulse?

YES
Support ventilation

NO
Begin CPR Attach AED ASAP

Analyze shockable rhythm?

YES
Defibrillate 1 time Resume CPR immediately for 2 minutes

NO
Resume CPR immediately for 2 minutes

ALS & transport

Back to Contents
UNIVERSAL ALGORITHM FOR ADULT EMERGENCY CARDIAC CARE FOR ALS

Assess Responsiveness

Not Responsive: Call for Defibrillator Assess Breathing

Responsive: Observe Treat as Indicated

Breathing

NO

Assess Circulation

If unconscious and no trauma, place in recovery position

YES

Pulse

NO

Begin CPR

VF/VT Present on Monitor

YES

Oxygen as needed VENTILATE as needed Cardiac Monitor Vital Signs IV with LR History & Physical Detailed Assessment

Suspected Cause

Pulmonary Edema/CHF See Protocol

Chest Pain See Protocol

Dysrhythmia

Too Slow

GO TO VT/VF ALGORITHM

GO TO ASYSTOLE ALGORITHM

GO TO PEA ALGORITHM

GO TO BRADYCARDIA ALGORITHM

GO TO TACHYCARDIA ALGORITHM

Too Fast
UNIVERSAL ALGORITHM FOR PEDIATRIC EMERGENCY CARDIAC CARE FOR BLS
Greater than 1 hour old and less than 13 years of age
If less than 1 hour old, refer to Newly Born Protocol

Unresponsive
Not Breathing

Pulse?

YES
Oxygen as needed
VENTILATE as needed
Target ventilation rate = 12–20 bpm
Vital Signs
History & Physical
Detailed Assessment

NO
Begin CPR
Attach AED with pediatric capability
100-120 compressions/minute
100% oxygen

Analyze shockable rhythm?

YES
Defibrillate 1 time
Resume CPR immediately for 2 minutes

NO
Resume CPR immediately for 2 minutes
UNIVERSAL ALGORITHM FOR PEDIATRIC EMERGENCY CARDIAC CARE FOR ALS

Greater than 1 hour old and less than 13 years of age
If less than 1 hour old, refer to Newly Born Protocol

1. Assess Responsiveness
   - Not Responsive:
     - Call for Defibrillator
     - Assess Breathing
   - Responsive:
     - Observe
     - Treat as Indicated

2. Breathing
   - NO: Assess Circulation
   - YES: If unconscious with adequate respiratory rate and effort and no trauma, place in recovery position

3. Pulse
   - NO: Suspected Cause
   - YES: Oxygen as needed
     - VENTILATE as needed
     - Target Ventilation Rate = 12-20 bpm
     - Cardiac monitor
     - Vital signs
     - IV with LR
     - History & Physical
     - Detailed Assessment

   A. Altered Mental Status: See Protocol
   B. Respiratory Distress
      - Allergic Reaction or Anaphylaxis: See Protocol, as appropriate
      - Asthma/COPD: See Protocol
      - Pulmonary Edema/CHF: See Protocol

   C. Dysrhythmia
      - Too Slow: GO TO PEDIATRIC BRADYCARDIA ALGORITHM
      - Too Fast: GO TO PEDIATRIC TACHYCARDIA ALGORITHM
**Indications**
- Slow heart rate, less than 60 bpm
- Bradycardic patients may also present with serious signs and symptoms including:
  - Chest pain or shortness of breath
  - Altered/decreased level of consciousness
  - Hypotension or hypoperfusion
  - Congestive heart failure or pulmonary congestion
  - Acute myocardial infarction

- Assess and treat for shock, if indicated.
- Continuously monitor airway and reassess vital signs every 5 minutes.

**BLS**
- Heart rate less than 60 bpm
- Serious signs or symptoms?

**YES**
- Transcutaneous Pacemaker (a)
- Atropine 0.5–1 mg IVP (b, c, e)
- Epinephrine 1 mL/min using approved epi infusion. (2-10 mcg/min on IV infusion pump) (f)

**NO**
- Second-degree Type II AV Block or Third-degree AV Block (d, e)
- Prepare for Transcutaneous pacing. Start pacing if the patient develops serious signs and symptoms related to the slow heart rate.

(a) Do not delay TCP while awaiting IV or atropine to take effect if the patient is symptomatic.
(b) Denervated transplanted hearts will not respond to atropine. Go directly to TCP.
(c) Atropine should be repeated every 3-5 minutes, if appropriate, to max dose of 0.04 mg/kg.
(d) Do not treat third-degree AV block or ventricular escape beats with amiodarone.

**MC**
- Medical consultation is required to administer atropine for patients with second-degree type II AV block or a third-degree AV block.
- Additional dosing above 1 mL/min (1 drop/second using 60 drop set and approved epinephrine infusion) requires medical consultation. Adults: titrate to systolic BP 90 mmHg.
- If patient develops discomfort with TCP, administer opioid per Pain Management protocol OR administer midazolam 0.1 mg/kg SLOW IVP over 1-2 minutes, in 2 mg increments. Max single dose of 5 mg. Reduce dose by 50% for patients 69 and older.
Indications
- Slow heart rate (refer to Normal Vital Signs Chart)

**BLS**
- Assess and treat for shock, if indicated.
- Continuously monitor airway and reassess vital signs every 5 minutes.

**ALS**

(a) Hemodynamically unstable is defined as a systolic blood pressure less than 60 in neonates (patients less than 28 days old), less than 70 in infants (patients less than 1 year of age), and less than \([70 + (2 \times \text{years}) = \text{systolic BP}]\) for patients greater than 1 year of age.

(b) Neonates (birth to 28 days), epinephrine ET 0.03 mg/kg (0.1 mg/mL) dilute with 1 mL.

(c) Volume infusion for neonates and volume-sensitive children, 10 mL/kg; for infant and child 20 mL/kg.

(d) Calcium chloride, 20 mg/kg (0.2 mL/kg) SLOW IVP/IO (50 mg/min). Max dose 1 gram.

(e) Sodium bicarbonate, 1 mEq/kg with medical consultation.

(f) If patient develops discomfort with TCP, administer opioid per Pain Management protocol OR administer midazolam 0.1 mg/kg SLOW IVP over 1-2 minutes, in 2 mg increments. Max single dose of 5 mg.
3.3-A  
**Adult Tachycardia Algorithm – Irregular Rhythm**

**BLS**
- Place patient in position of comfort.
- Assess and treat for shock, if indicated.
- Continuously monitor airway and reassess vital signs every 5 minutes.

**ALS**

- **Signs and symptoms related to tachycardia:** hypotension, acutely altered mental status, signs of shock, ischemic chest discomfort/AMI, or acute heart failure.
- **Consider sedation** (*midazolam*). However, overall patient status, including BP, may affect ability to administer sedative.
- **Consider calcium chloride** 500 mg IVP for hypotension induced by *diltiazem*.
- **If rate does not slow in 15 minutes,** administer a second dose of *diltiazem* (15-25 mg over 2 minutes).
- **These rhythms include Wolff-Parkinson White (WPW) syndrome, Lown-Ganong-Levine syndrome (LGL), and Mahaim type.**

---

**General Patient Care**

- **Hemodynamically unstable with life-threatening, rate-related signs and symptoms (a) and ventricular rate greater than 150 bpm?**
  - **Yes**
    - **Perfom synchronized cardioversion (b)**
  - **No**

- **Underlying rhythm**
  - **Irregular**
    - **Narrow QRS** (Atrial fibrillation or Atrial flutter)
      - Diltiazem 0.25 mg/kg (10-20 mg) IV/IO over 2 minutes (c,d) BP greater than 100
    - **Wide QRS** (Atrial fibrillation or Atrial flutter with aberrancy)
      - Amiodarone 150 mg IV/IO over 10 minutes (Mixed in 50-100 mL of approved diluent); Repeat if necessary
    - **Polymorphic wide QRS complex tachycardia (Torsades de pointes)**
      - Magnesium sulfate 1-2 grams IV/IO over 2 minutes
    - **History of accessory pathway WPW or LGL irregular wide complex**
      - **(e)**

- **Hemodynamically unstable with life-threatening, rate-related signs and symptoms (a) and ventricular rate greater than 150 bpm?**
  - **Yes**
    - **Perform synchronized cardioversion (b)**
  - **No**

---

**Release Date July 1, 2022**
BLS
- Place patient in position of comfort.
- Assess and treat for shock, if indicated.
- Continuously monitor airway and reassess vital signs every 5 minutes.

ALS

(a) Signs and symptoms related to tachycardia: hypotension, acutely altered mental status, signs of shock, ischemic chest discomfort/AMI, or acute heart failure.
(b) Consider sedation (midazolam). However, overall patient status, including BP, may affect ability to administer sedative.
(c) Be prepared for up to 40 seconds of asystole.
(d) These rhythms include Wolff-Parkinson White (WPW) syndrome, Lown-Ganong-Levine syndrome (LGL), and Mahaim type.
BLS

- Assess and treat for shock, if indicated.
- Continuously monitor airway and reassess vital signs every 5 minutes.

ALS

Identify and treat underlying causes

Evaluate QRS duration

Narrow (less than or equal to 0.09 seconds)

Identify and treat underlying cause

Probable sinus tachycardia

Probable supraventricular tachycardia (a)

Consider vagal maneuvers

Consider adenosine (e)

Consider (c) (d) cardioversion

Wide regular (greater than 0.09 seconds)

Possible VT (g)

Hemodynamically unstable? (b)

YES

Cardiovert 0.5 J/kg (c) (d)

Cardiovert 1 J/kg

Cardiovert 2 J/kg

IV/IO access

Amiodarone (f)

NO

Consider adenosine (e)

Amiodarone (f)

(a) - Ventricular Heart Rates in excess of: Infant 220 bpm or Pediatric 180 bpm
(b) - Hemodynamically unstable is defined as a systolic blood pressure less than 60 in neonates (patients from birth to 28 days old), less than 70 in infants (patients less than 1 year of age), less than [70 + (2 x years) = systolic BP] for patients greater than 1 year of age, altered mental status with hypoperfusion evidenced by delayed capillary refill, pallor, or peripheral cyanosis.
(c) - If calculated joules setting is lower than cardioversion device is able to deliver, use the lowest joules setting possible or obtain medical consultation.
(d) - Consider sedation (midazolam with medical consultation). However, overall patient status, including BP, may affect ability to administer sedative.
(e) - Adenosine: 0.1 mg/kg rapid IV/IO, maximum 6 mg. Second and third doses 0.2 mg/kg rapid IV/IO, maximum single dose 12 mg. Be prepared for up to 40 seconds of asystole. (Contraindicated in polymorphic or irregular wide complex tachycardia)
(f) - Amiodarone: 5 mg/kg IV/IO over 20 minutes (mixed in 50 - 100 mL of approved diluent). Obtain 12-lead EKG prior to administration of amiodarone.
(g) If torsades de pointes, administer magnesium sulfate (25 mg/kg IV/IO to a maximum of 2 grams over 2 minutes).
Cardiac Arrest – Adult

Indications
- Adult patients (medical arrest: 18 years of age and older; trauma arrest: 15 years of age and older) who are unconscious, apneic, and pulseless

**BLS**
- Perform high-quality uninterrupted chest compressions (manual or mechanical) as soon as possible and until defibrillator available.
- Apply AED as soon as available.
- Follow machine prompts regarding rhythm analyses and shocks.
- Limit breaks in compressions to 10 seconds or less for rhythm analysis periods and during shocks; perform compressions while defibrillator is charging.
- **On-scene resuscitation**: Patients who are found in arrest or who arrest prior to transport and are attended to by BLS clinicians must only be resuscitated in place (with minimal movement, no attempts at patient loading, and no attempts at transport) until the following have been accomplished:
  - **Medical etiology**: The patient has received a minimum of five two-minute cycles of chest compressions and rhythm interpretation
  - **Traumatic etiology**: Patient has received treatments for reversible causes per Trauma Protocol: Trauma Arrest protocol
- **Exemptions** from on-scene resuscitation:
  - Physical barriers prevent resuscitation
  - Clinicians are in danger
  - Pregnant patients
  - Patients in cardiac arrest thought to be secondary to hypothermia or submersion
- Following the initial on-scene resuscitation above, clinicians may continue on-scene resuscitation until termination of resuscitation or transport the patient at any time. Clinicians should ensure that a mechanical CPR device is in place (if available) prior to transport.
- Pregnancy: For pregnant patients greater than 20 weeks gestation in cardiac arrest, provide constant left lateral uterine displacement.

**ALS**
- Assess for shockable rhythm at next appropriate interval and treat appropriately.
- **On-scene resuscitation**: Patients who are found in arrest or who arrest prior to transport and are attended to by ALS clinicians must remain in place (with minimal movement, no attempts at patient loading, and no attempts at transport) until the following have been accomplished:
  - **Medical etiology**: The patient has received three doses of epinephrine, regardless of algorithm being followed
  - **Traumatic etiology**: The patient has received treatments for reversible causes per Trauma Arrest protocol
- Following the initial on-scene resuscitation above, clinicians may choose to continue the on-scene resuscitation until termination of resuscitation or to transport the patient at any time. Clinicians should ensure the following prior to transport:
  - Mechanical CPR (mCPR) in place (if available)
  - Placement of an airway that facilitates ventilation during transport by a restrained clinician
- If ROSC, refer to ROSC protocol.
- Consider Termination of Resuscitation when appropriate.

**MC**
- Not applicable.
ALS

Adult Pulseless Electrical Activity (PEA)/Asystole Algorithm

3.4-A

Cardiac: Adult Pulseless Electrical Activity (PEA)/Asystole Algorithm

Release Date July 1, 2022

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3.4-A

(a) Confirm asystole in more than one lead.
(b) Volume infusion is Lactated Ringer’s 20 mL/kg.
(c) Epinephrine is not indicated in adult traumatic cardiac arrest.

All hypothermic patients in cardiac arrest shall be rewarmed; see exclusions to Termination of Resuscitation (3.6-A).
When the patient’s condition changes, indicating transition to a new treatment algorithm, the new treatment shall take into account prior therapy (e.g., previously administered medications).
Ventricular Fibrillation and Pulseless Ventricular Tachycardia Algorithm

Perform CPR and assure adequate ventilation
VF/VT present on monitor

Defibrillate 1 time
Resume CPR immediately for 2 minutes

Confirm Rhythm

Persistent or Recurrent VF/VT

Defibrillate 1 time
Resume CPR immediately for 2 minutes

IV/IO with LR

Epinephrine (0.1 mg/mL)
1 mg IV/IO every 4 minutes up to a max of 4 doses for the initial arrest. If arrest occurs after ROSC, an additional 2 doses may be administered.

Defibrillate 1 time
Resume CPR immediately for 2 minutes

Amiodarone
300 mg IV/IO push
May repeat once 150 mg IV/IO push (a) (b) (c)

Defibrillate 1 time
Resume CPR immediately for 2 minutes

Return of Spontaneous Circulation
GO TO ROSC PROTOCOL

PEA
GO TO PEA ALGORITHM

Asystole
GO TO ASYSTOLE ALGORITHM

(a) - Sodium bicarbonate 1 mEq/kg IV/IO, only in cases for which the suspected cause of cardiac arrest is acidosis, NA channel blocker (tricyclic antidepressant and phenobarbital) overdose.

(b) - If torsades de pointes is present, give magnesium sulfate 1–2 grams IV/IO over 2 minutes before amiodarone.

(c) - For refractory VF/VT after amiodarone, give magnesium sulfate 1–2 grams IV/IO over 2 minutes.

When the patient’s condition changes, indicating the transition to a new treatment algorithm, the new treatment shall take into account prior therapy (e.g., previously administered medications).
**Indications**

- Pediatric patients (medical arrest: less than 18 years of age; trauma arrest: less than 15 years of age) who are unconscious, apneic, and pulseless

**BLS**

- Perform high-quality uninterrupted chest compressions (manual or mechanical) as soon as possible and until defibrillator available.
- Apply AED as soon as available.
- Follow machine prompts regarding rhythm analyses and shocks.
- Limit breaks in compressions to 10 seconds or less for rhythm analysis periods and during shocks; perform compressions while defibrillator is charging.
- **On-scene resuscitation**: Patients who are found in arrest or who arrest prior to transport and are attended to by BLS clinicians must only be resuscitated in place (with minimal movement, no attempts at patient loading, and no attempts at transport) until the following have been accomplished:
  - **Medical etiology**: the patient has received a minimum of five two-minute cycles of chest compressions and rhythm interpretation
  - **Traumatic etiology**: patient has received treatments for reversible causes per Trauma Protocol: Trauma Arrest protocol
- **Exemptions** from on-scene resuscitation:
  - Physical barriers prevent resuscitation
  - Clinicians are in danger
  - Pregnant patients
  - Patients in cardiac arrest thought to be secondary to hypothermia or submersion
- Following the initial on-scene resuscitation above, clinicians may continue on-scene resuscitation until termination of resuscitation or transport the patient at any time. Clinicians should ensure that a mechanical CPR device is in place (if available) for patients 13 years of age and older prior to transport.
- Pregnancy: For pregnant patients greater than 20 weeks gestation in cardiac arrest, provide constant left lateral uterine displacement.

**ALS**

- Assess for shockable rhythm at next appropriate interval and treat appropriately.
- Only in a pediatric or neonatal arrest situation, naloxone, atropine sulfate, and epinephrine, can be administered via the ET route. Medications administered for pediatric patients via the endotracheal tube route shall be 2–2.5 times the IV dose for naloxone and atropine sulfate, and 10 times the IV dose for epinephrine (1 mg/mL). All ET medications shall be diluted in 5 mL of Lactated Ringer's for pediatric patients.
- **On-scene resuscitation**: See BLS section above.
- Following initial on-scene resuscitation, clinicians may choose to continue the on-scene resuscitation until termination of resuscitation or to transport the patient at any time. Clinicians should ensure the following prior to transport:
  - Mechanical CPR (mCPR) in place for patients 13 years of age and older (if available)
  - Placement of an airway that facilitates ventilation during transport by a restrained clinician
- If ROSC, perform 12-lead EKG and transport the patient to Children’s National Medical Center or Johns Hopkins Children’s Center by ground or medevac. If arrival time is greater than 30 minutes to either of these destinations, transport to the closest appropriate facility.
- If no ROSC, transport to the closest appropriate facility or consider Termination of Resuscitation protocol, as appropriate.

**MC**

- Not applicable.
(a) - Continue cycle of epinephrine, defibrillation at 8 J/kg then 10 J/kg.

(b) - Neonates (0–28 days), epinephrine ET 0.03 mg/kg (0.1 mg/mL) dilute with 1 mL.

(c) - Sodium bicarbonate, 1 mEq/kg, with medical consultation. See sodium bicarbonate.

(d) - Calcium chloride, 20 mg/kg (0.2 mL/kg) SLOW IVP/IO (50 mg/min). Max dose 1 gram.

(e) - Volume infusion for neonates and volume-sensitive children, 10 mL/kg; for infant and child 20 mL/kg.

(f) - If torsades de pointes, administer magnesium sulfate (25 mg/kg IV/IO to a maximum of 2 grams over 2 minutes before amiodarone).

When the patient’s condition changes, indicating the transition to a new treatment algorithm, the new treatment shall take into account prior therapy (e.g., previously administered medications).
Return of Spontaneous Circulation (ROSC) – Adult

Indications
- Patients 18 years of age and older who have been revived from cardiac arrest (return of pulses) due to a medical etiology
- For patients resuscitated from traumatic arrest, refer to Multiple/Severe Trauma protocol.

BLS
- Verify presence of a carotid pulse. If any doubt exists as to whether a carotid pulse is present, initiate CPR and refer to appropriate Cardiac Arrest protocol.
- If apneic or inadequate respirations, continue to support ventilations.
- Frequently reassess vital signs. Treat any abnormalities in accordance with appropriate shock, respiratory, or cardiac protocols.
- Rendezvous with ALS or transport to the closest ED.
- If available and not already in place, apply mechanical CPR (mCPR) device in standby mode.

ALS
- Obtain 12-lead EKG; if STEMI, treat according to STEMI protocol.
- Establish IV/IO access, if not already obtained.
- Identify cardiac rhythm and treat according to appropriate algorithm.
  - If VF or VT was present during arrest and amiodarone not yet given, consider amiodarone 150 mg IV/IO over 10 minutes. (Presence of a perfusing sinus rhythm is necessary for the administration of amiodarone.)
- Treat hypotension with Lactated Ringer’s fluid bolus, titrate to systolic blood pressure of 90 mmHg, or if ineffective, refer to the Epinephrine Infusion protocol. Refer to Shock: Hypoperfusion protocol.
- Reassess need for airway management or intubation, if not already addressed.
- Identify and treat underlying causes that contributed to the cardiac arrest.
- Initiate transport to a Cardiac Intervention Center, unless exceptions below apply.
  - Exceptions:
    - Obvious non-cardiac cause for arrest (e.g., drowning, asphyxiation, opiate overdose)
    - Transport time to Cardiac Interventional Center is more than 45 minutes greater than transport time to nearest ED

MC
- Obtain medical consultation if patient’s clinical instability will not allow for safe transport to Cardiac Interventional Center due to extended transport time.
- All post-cardiac arrest patients are priority 1, and require medical consultation.

Clinical Pearls
- Consider use of helicopter transport if patient has sustained ROSC and it would provide a time-appropriate arrival at a cardiac intervention center.
**Indications**
- Pediatric patients less than 18 years of age who have been revived from cardiac arrest (return of pulses) due to a medical etiology
- For patients resuscitated from traumatic arrest, refer to *Multiple/Severe Trauma* protocol.

**BLS**
- Verify presence of a carotid pulse. If any doubt exists as to whether a carotid pulse is present, initiate CPR and refer to appropriate *Cardiac Arrest* protocol.
- If apneic or inadequate respirations, continue to support ventilations.
- Frequently reassess vital signs. Treat any abnormalities in accordance with appropriate shock, respiratory, or cardiac protocols.
- Rendezvous with ALS or transport to the closest ED.
- For patients 13 years of age and older, apply mechanical CPR (mCPR) device in standby mode, if available and not already in place.

**ALS**
- Establish IV/IO access, if not already obtained.
- Identify cardiac rhythm and treat according to appropriate algorithm.
- Reassess need for airway management or intubation, if not already addressed.
- If lungs are clear, treat hypotension with *Lactated Ringer’s* 20 mL/kg IV fluid bolus per *Shock: Hypoperfusion* protocol, with the following blood pressure goals:
  - For patients 10 years and older (including adults), systolic blood pressure greater than 90 mmHg
  - For patients under 10 years of age, systolic blood pressure greater than 70 + 2x age in years mmHg; or
  - Systolic blood pressure ordered by the Pediatric Base Station. May repeat *Lactated Ringer’s* 20 mL/kg fluid bolus one time.
- Transport to Children’s National Medical Center or Johns Hopkins Children’s Center with the following exceptions:
  - Transport time is 30 minutes greater than transport time to nearest ED, or
  - Patient’s clinical instability will not allow for safe transport to one of the above centers due to transport time.

**MC**
- If patient’s clinical instability will not allow for safe transport to a pediatric center, obtain a medical consult.
- All post-cardiac arrest patients are priority 1, and require medical consultation with a Pediatric Base Station, which may assist with destination determination.
- Third and subsequent fluid boluses, *Lactated Ringer’s* 20 mL/kg IV/IO require medical consultation
- Pediatric *epinephrine* infusion dosage
  - The following dosing chart should be used for pediatric patients less than 50 kg, using approved *epinephrine* infusion and 60-drop set:

<table>
<thead>
<tr>
<th>Weight range (kg)</th>
<th>Initial epinephrine dose</th>
<th>If goal blood pressure not achieved at 5 min, increase to</th>
</tr>
</thead>
<tbody>
<tr>
<td>LESS than 10 kg</td>
<td>6 drops/min (0.1 mL/min)</td>
<td>12 drops/min (0.2 mL/min)</td>
</tr>
<tr>
<td>10-19 kg</td>
<td>12 drops/min (0.2 mL/min)</td>
<td>24 drops/min (0.4 mL/min)</td>
</tr>
<tr>
<td>20-29 kg</td>
<td>18 drops/min (0.3 mL/min)</td>
<td>36 drops/min (0.6 mL/min)</td>
</tr>
<tr>
<td>30-39 kg</td>
<td>24 drops/min (0.4 mL/min)</td>
<td>48 drops/min (0.8 mL/min)</td>
</tr>
<tr>
<td>40-49 kg</td>
<td>30 drops/min (0.5 mL/min)</td>
<td>60 drops/min (1.0 mL/min)</td>
</tr>
</tbody>
</table>

- If blood pressure goal in ALS section has not been met after 10 minutes, obtain medical consultation.
Termination of Resuscitation – Adult

3.6-A

Indications

- Patients who are in cardiac arrest due to medical or traumatic etiology

Exclusions

- The following patients should receive care according to appropriate protocol, without TOR, and transport to the closest appropriate facility:
  - Pregnant patients
  - Patients in cardiac arrest that is suspected to be due to hypothermia or submersion

<table>
<thead>
<tr>
<th>BLS</th>
<th>If the patient meets the criteria listed in the <em>Pronouncement of Death in the Field</em> protocol, EMS clinicians should terminate resuscitation efforts.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• BLS clinicians may terminate resuscitation for adult patients (age 18 or older) if:</td>
</tr>
<tr>
<td></td>
<td>- ALS resources are genuinely unavailable, and</td>
</tr>
<tr>
<td></td>
<td>- The patient has received a minimum of 15 two-minute cycles of HPCPR, and</td>
</tr>
<tr>
<td></td>
<td>- During the five AED analyses immediately prior to TOR there was “no shock advised.”</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ALS</th>
<th>Medical etiology: may terminate resuscitation for adult patients (age 18 years or older) if:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Patient has received 15 two minute cycles of HPCPR, and the patient is:</td>
</tr>
<tr>
<td></td>
<td>- in asystole, or</td>
</tr>
<tr>
<td></td>
<td>- in VF, pulseless VT, or PEA with an ETCO₂ of less than 15 mmHg</td>
</tr>
<tr>
<td></td>
<td>• Traumatic etiology: may terminate resuscitation regardless of total resuscitation time for adult patients (15 years or older) if:</td>
</tr>
<tr>
<td></td>
<td>- Patient presents in asystole, or</td>
</tr>
<tr>
<td></td>
<td>- Patient’s cardiac rhythm changes to asystole during the resuscitation, or</td>
</tr>
<tr>
<td></td>
<td>- Blunt trauma patient remains in PEA or VF after 5 two-minute cycles of HPCPR according to the <em>Trauma Protocol: Trauma Arrest</em> protocol</td>
</tr>
</tbody>
</table>

| MC  | Not applicable. |

Clinical Pearls

- If the patient does not meet TOR criteria, continue resuscitation and re-evaluate at the next rhythm check.
- For traumatic arrest patients, asystole and resuscitations lasting longer than 10 minutes are independent predictors of mortality. Treatment of the trauma arrest patient should focus on identifying and treating reversible causes during that narrow resuscitative window. TOR and transport decisions should only be made after administering time-sensitive therapies.
Adult Termination of Resuscitation: Medical Arrest
(Age 18 Years and Older)

Cardiac Arrest (Considering Termination of Resuscitation)

Exclusions:
- Pregnant
- Hypothermia or submersion

NO

Meets Pronouncement of Death criteria

YES

Should terminate resuscitation

NO

Minimum of 15 2-minute cycles of HPCR. Identify and treat reversible causes

ROSC

YES

Transport

NO

Asystole OR VF/Pulseless VT/PEA with ETCO₂ less than 15 mmHg

YES

May terminate resuscitation

NO

Continue resuscitation. Reevaluate at next rhythm check.
Adult Termination of Resuscitation: Trauma Arrest (Age 15 Years and Older)

Cardiac Arrest (Considering Termination of Resuscitation)

Exclusions:
- Pregnant
- Hypothermia or submersion

NO

Meets Pronouncement of Death criteria

YES

Should terminate resuscitation

Rhythm

VF, VT, or PEA

Penetrating

Asystole

Less than 15 min. to trauma center

Minimum of 5 two-minute cycles of HPCPR. Identify and treat reversible causes.

HPCPR/Treat reversible causes, transport to trauma center

May terminate resuscitation

More than 15 min. to trauma center

HPCPR/Treat reversible causes, transport to local ER

ROSC

YES

Transport

NO
Indications
• Patients who are in cardiac arrest due to medical or traumatic etiology

Exclusions
• The following patients should receive care according to appropriate protocol, without TOR, and transport to the closest appropriate facility:
  ■ Pregnant patients
  ■ Patients in cardiac arrest that is suspected to be due to hypothermia or submersion

**BLS**
- If the patient meets the criteria listed in the *Pronouncement of Death in the Field* protocol, EMS clinicians should terminate resuscitation efforts.
- May not terminate resuscitation for pediatric **medical** arrest patients (under age 18 years).
- May terminate resuscitation for pediatric **traumatic** arrest patients (under age 15 years) if:
  ■ ALS resources are genuinely unavailable, **and**
  ■ The patient has received a minimum of 15 two-minute cycles of HPCPR, **and**
  ■ During the five AED analyses immediately prior to TOR there was “no shock advised.”

**ALS**
- **Medical etiology:** may terminate resuscitation of pediatric patients (less than 18 years of age) if:
  ■ Patient has received 15 two-minute cycles of HPCPR, **and** at least 1 dose of **epinephrine** **and**:
    ♦ Patient is in asystole, **and**
    ♦ Patient has a sustained ETCO₂ of less than 15 mmHg, **and**
    ♦ In the judgment of EMS and law enforcement on scene, there is adequate social/ emotional support and safety for civilians and professionals on scene, **and**
    ♦ In the judgment of EMS and law enforcement, scene is amenable to leaving patient on scene.
- **Traumatic etiology:** may terminate resuscitation for pediatric patients (less than 15 years of age) if:
  ■ Patient has received 5 two-minute cycles of HPCPR without ROSC according to the *Trauma Protocol: Trauma Arrest* protocol **and**
  ■ Patient is in asystole, **and**
  ■ Patient has a sustained ETCO₂ of less than 15 mmHg, **and**
  ■ In the judgment of EMS and law enforcement on scene, there is adequate social/ emotional support and safety for civilians and professionals on scene, **and**
  ■ In the judgment of EMS and law enforcement, scene is amenable to leaving patient on scene.

**MC**
- Not applicable.

Clinical Pearls
- If patient does not meet TOR criteria, continue resuscitation and reevaluate at the next rhythm check.
Pediatric Termination of Resuscitation: Medical Arrest
(Patients Under 18 Years of Age)

Cardiac Arrest (Considering Termination of Resuscitation)

Exclusions:
- Pregnant
- Hypothermia or submerison

NO

Meets Pronouncement of Death criteria?

YES
- Should terminate resuscitation

NO
- Minimum of 15 two-minute cycles of HPCPR—treat reversible causes of arrest

ROSC

YES
- Transport

NO

Asystole with ETCO2, less than 15 mmHg and administration of at least 1 dose of epinephrine (0.1 mg/mL)

YES
- May terminate resuscitation

NO
- Continue resuscitation. Reevaluate at next rhythm check/consider transport.
Pediatric Termination of Resuscitation: Trauma Arrest
(Patients Under 15 Years of Age)

Cardiac Arrest (Considering Termination of Resuscitation)

Exclusions:
- Pregnant
- Hypothermia or submersion

NO

Meets Pronouncement of Death criteria?

YES

Should terminate resuscitation

NO

Minimum of 5 two-minute cycles of HPCPR—treat reversible causes of arrest

Rhythm

Asystole AND ETCO$_2$ less than 15 mmHg?

May terminate resuscitation

VF, VT, PEA OR Asystole with ETCO$_2$ greater than 15 mmHg?

Less than 15 min. to Pediatric Trauma Center

HPCPR/Treat reversible causes, transport to Pediatric Trauma Center

More than 15 min. to Pediatric Trauma Center

HPCPR/Treat reversible causes, transport to local ED or Adult Trauma Center
Indications

- EMS clinicians may use this protocol to pronounce the death of a patient when one or more of the following criteria have been met:
  - Decapitation
  - Rigor mortis
  - Decomposition
  - Dependent lividity
  - Pulseless, apneic patient in a multi-casualty incident where system resources are required for the stabilization of living patients
    - Patient may be “black tagged” by BLS or ALS, but asystole must be confirmed by ALS prior to formal pronouncement of death.
  - Pulseless, apneic patient with an injury not compatible with life
    - Exception: Obviously pregnant female patient should have resuscitation initiated and be transported to the closest appropriate facility.
  - EMS clinician has terminated resuscitation per the Termination of Resuscitation protocol

BLS

- Confirm that the patient is unresponsive, pulseless, and apneic.
- Document the exact time and location of the pronouncement of death.
- Notify law enforcement and follow local jurisdictional policies.
- Organ donor hotline: If deceased patient is organ donor and law enforcement has released the body to the family, please assist the family in calling either:
  - For Charles, Montgomery and Prince George's Counties: 703-641-0100
  - For all other areas of Maryland: 800-923-1133
- If death is pronounced during transport, deliver the patient to the hospital and follow hospital policies. Law enforcement must be notified, as they may need to notify the medical examiner's office.

ALS

- Refer to BLS protocol.

MC

- Not applicable.

Clinical Pearls

- Health General Article §5-202 provides that: an individual is dead if, based on ordinary standards of medical practice, the individual has sustained either:
  - Irreversible cessation of circulatory and respiratory functions; or
  - Irreversible cessation of all functions of the entire brain, including the brain stem
Indications
- A MOLST Form or Acceptable EMS DNR Order is presented to EMS by family/caregivers or found on scene, and
  - Patient is in cardiac or respiratory arrest, or
  - Patient is non-verbal or lacks medical decision-making capacity

- Resuscitation status:
  - Attempt CPR – if cardiac or respiratory arrest occurs: perform CPR, artificial ventilation, and all medical efforts that are indicated during arrest in order to restore or stabilize cardiopulmonary function
  - MOLST A-1 – if cardiac or respiratory arrest occurs: do not attempt resuscitation (no CPR)
    ♦ Prior to arrest: maximal restorative efforts including intubation
  - MOLST A-2 – if cardiac or respiratory arrest occurs: do not attempt resuscitation (no CPR)
    ♦ Prior to arrest: comprehensive efforts to prevent arrest excluding intubation
  - MOLST B – if cardiac or respiratory arrest occurs: do not attempt resuscitation (no CPR)
    ♦ Prior to arrest: limited, palliative care only

- Acceptable DNR Orders
  - Maryland MOLST Form or Bracelet
    ♦ May be an original, copy, or electronic format for patient care decisions, however, sending facility must provide paper copy to EMS prior to patient transport
  - Maryland EMS/DNR Form or Bracelet
    ♦ There is no expiration on older versions of DNR forms.
  - Medic Alert DNR Bracelet or Necklace
  - Out-of-state EMS/DNR Form
  - Oral DNR Order from EMS System Medical Consultation
  - Oral DNR Order from other on-site physician, physician assistant, or nurse practitioner

- Unacceptable DNR Orders
  - Advanced directives (without a MOLST or DNR Order) or other oral or written requests shall not be honored by EMS without EMS System Medical Consultation

- Revocation of DNR Orders
  - An EMS/DNR Order may be revoked at any time by:
    ♦ Physical cancellation or destruction of all EMS/DNR Order devices; or
    ♦ A verbal statement by the patient made directly to EMS clinicians requesting resuscitation or palliative care only. In this case, EMS/DNR devices do not need to be destroyed. EMS clinicians must thoroughly document the revocation. A verbal revocation by the patient is only good for the current response for which it was issued.
  - An authorized decision-maker, other than the patient, cannot revoke an EMS/DNR Order verbally.
    ♦ Decision-makers with the authority to revoke an EMS/DNR Order must either void or withhold all EMS/DNR Order devices if they wish resuscitation for the patient. If there is any confusion, the EMS clinician should consult a Base Station.
EMS DNR/MOLST Medical Protocols

- Perform limited patient assessment.
  - Check for a palpatable pulse.
  - Check for respirations in an unresponsive patient.
  - Check for MOLST form or other acceptable EMS/DNR Order.

- Resuscitate/Do Not Resuscitate Criteria
  - If MOLST form or other acceptable EMS/DNR Order is present and the patient is in cardiac or respiratory arrest, no resuscitative measures shall be initiated.
  - If MOLST form or other acceptable EMS/DNR Order is not present, revoked, or otherwise void, EMS clinician shall treat and transport the patient, as appropriate.
    - If EMS clinicians believe that resuscitation or further resuscitative efforts are futile, they may initiate the Termination of Resuscitation protocol.
  - If the patient is conscious and able to communicate directly to EMS clinicians that they revoke the MOLST or other EMS/DNR Order verbally, then EMS clinicians shall treat and transport the patient, as appropriate.
  - If the EMS/DNR patient (Option A-1, A-2, B) experiences respiratory or cardiac arrest, EMS shall withhold or withdraw further resuscitation and provide support to the family and caregivers.

- MOLST A-1 – Maximal Restorative Care, including intubation
  - Prior to respiratory or cardiac arrest: the Option A-1 patient shall receive the full scope of interventions permissible under The Maryland Medical Protocols for Emergency Medical Services, including: intubation, CPAP/BiPAP, cardiac monitoring, cardioversion, cardiac pacing, IVs, and medications in attempt to forestall cardiac or respiratory arrest.
  - If respiratory or cardiac arrest occurs: do not initiate CPR or any resuscitative efforts. Withhold or withdraw resuscitative efforts if they were already in progress prior to discovery of the MOLST or EMS/DNR Order.

- MOLST A-2 – Comprehensive Efforts, excluding intubation
  - Prior to respiratory or cardiac arrest: same as option A-1, except no intubation is permitted
  - If respiratory or cardiac arrest occurs: no CPR, same as option A-1

- MOLST B – Palliative Care
  - Prior to respiratory or cardiac arrest, provide supportive treatment:
    - Respiratory
      - Open and maintain airway using chin lift, jaw thrust, finger sweep, nasopharyngeal or oropharyngeal airway, Heimlich maneuver, or laryngoscopy with Magill forceps for suspected airway obstruction, but no intubation, cricothyroidotomy, or tracheostomy
      - Oxygen: may provide passive oxygen via nasal cannula or non-rebreather mask, but no positive pressure oxygen via BVM, demand valve or ventilator. Pulse oximetry and capnography may be used.
      - Ventilator patients: if the patient is found on an outpatient ventilator and is not in cardiac arrest, maintain ventilator support during transport to the hospital
        - If the patient on an outpatient ventilator is found in cardiac arrest, contact online medical direction before disconnecting the ventilator.
    - Suction as necessary
    - Position for comfort
External bleeding
- Standard treatment; direct pressure, tourniquet
- No IVs

Immobilize fractures with devices to minimize pain

Uncontrolled pain or other symptoms (e.g., severe nausea)
- Allow patient, family or other health care clinicians to administer patient-prescribed medications. Document this on the PCR.
- Patient controlled analgesia (PCA) systems shall be maintained and monitored.
- For the patient with significant pain or pain with prolonged transport, initiate the Pain Management protocol.

Existing IV lines shall be maintained in place.

Transport: upon request of the patient, family or caregivers, EMS clinicians may transport Option B EMS/DNR patients to a specified inpatient hospice facility for pain control, symptom management or respite care (in lieu of transport to a hospital-based emergency department). EMS clinicians must notify the hospice facility prior to transport.

Documentation
- A copy of the MOLST or other acceptable EMS/DNR Order must be transported with the patient to the emergency department or inpatient hospice facility.
- MOLST or EMS/DNR order status must be documented in the patient care report.

Non-transported EMS/DNR Patients
- Follow local operational procedures for handling deceased patients.
- Do not remove DNR or Medical Alert Bracelets or Necklaces from the patient; leave the original MOLST or EMS/DNR Order with the patient.
- Law enforcement or medical examiner's office need to be notified only in the case of sudden or unanticipated death that occurs:
  - By violence
  - By suicide
  - As the result of an accident
  - Suddenly, if the deceased was in apparent good health, or
  - In any suspicious or unusual manner

Refer to BLS protocol

An oral DNR Order from EMS System Medical Consultation is acceptable if a MOLST or DNR form is not present.

Obtain medical consultation if the MOLST or DNR form instructions are unclear or the form is unreadable.
EMS DNR/MOLST Flowchart

EMS/DNR Order Presented:
1. Maryland EMS/DNR Order Form
2. Other State EMS/DNR Order Form
3. Maryland EMS/DNR Bracelet Insert
4. Medic Alert DNR Bracelet or Necklace
5. Oral DNR Order from medical consultation
6. Oral DNR Order from other on-site physician, physician assistant, or nurse practitioner
7. Maryland MOLST form
8. Maryland MOLST Bracelet Insert

If spontaneous respirations are ABSENT, OR palpable pulse is ABSENT, OR patient meets “Pronouncement of Death” criteria:
DO NOT ATTEMPT RESUSCITATION

If spontaneous respirations AND palpable pulse are PRESENT:
DETERMINE DNR CARE OPTION “A” OR “B”

If OPTION “A” or “A (DNI)”: Treat in accordance with all Maryland Protocols

If OPTION “B”: Treat in accordance with Maryland Palliative Care Protocol

If patient loses spontaneous respirations or palpable pulse, withdraw resuscitative efforts.
Indications

- Angina or anginal equivalents
- Chest pain, pressure or discomfort
- Pain or discomfort in the upper abdomen, arm, or jaw
- Shortness of breath
- Unexplained diaphoresis

**BLS**

- Place patient in position of comfort.
- Administer aspirin 324 mg or 325 mg chewed, if not given prior to EMS arrival.
- Assist with administration of patient-prescribed nitroglycerin (BLS) 0.4 mg SL.
  - May be repeated in 3-5 minutes if chest pain persists, blood pressure is greater than 90 mmHg, and pulse is between 60-150 bpm. Maximum 3 doses total (patient and EMT-assisted)
- Assess and treat for shock if indicated.

**ALS**

- Perform a 12-lead EKG as soon as possible.
  - EKG should occur within 10 minutes of contact with an EKG-capable clinician.
  - Document the patient’s last name, first initial, age, and gender on the EKG. These identifiers should be on the transmission copy (if able to transmit) and shall be on the delivered printed copy.
- Establish IV access.
- Nitroglycerin 0.4 mg SL. May be repeated if symptoms persist, blood pressure is greater than 90 mmHg and pulse is between 60-150 bpm, to a maximum dose of 1.2 mg SL.
  - If patient does not have a prescription or previous history of nitroglycerin use, an IV must be established prior to administration.
- Administer opioid per Pain Management protocol.

**MC**

- Medical consultation must be obtained for nitroglycerin in the following scenarios:
  - Additional doses of nitroglycerin, beyond 3 doses or 1.2 mg SL
  - No IV access
  - If patient’s systolic blood pressure drops more than 20 mmHg per dose of nitroglycerin given, consult for further doses.
- Pediatrics: Consult a Pediatric Base Station for patients who have not reached their 18th birthday with chest pain with associated dysrhythmias, cardiac disease, or blunt trauma.

**Clinical Pearls**

- Nitroglycerin is contraindicated for any patient having taken medication for pulmonary artery hypertension (e.g., Adcirca® or Revatio®) or erectile dysfunction (e.g., Viagra®, Levitra®, or Cialis®) within the past 48 hours.
**Indications**
- Patient must meet both criteria:
  - Three or more distinct ICD shocks **and**
  - Obvious device malfunction with at least one EMS clinician-witnessed inappropriate shock (e.g., alert patient in atrial fibrillation with rapid ventricular rate or SVT)

**BLS**
- Place patient in position of comfort.
- Assess and treat for shock, if indicated.

**ALS**
- Establish IV/IO access.
- Monitor cardiac rhythm and treat according to appropriate algorithm.
- **Donut magnet:** Place a donut magnet directly over device. Magnet placed directly over ICD will deactivate device and shocks will not be delivered. After defibrillator is deactivated, tape the magnet firmly in place and treat cardiac rhythm according to the appropriate algorithm.
- If ICD deactivation indications are questionable or deactivation is unsuccessful (or a donut magnet is not available) and undesired shocks continue, medications may be administered for patient comfort:
  - Administer opioid per *Pain Management* protocol or midazolam with consultation.
- If the patient becomes unstable or in the event of a rhythm change where a shock is desired, remove the magnet to reactivate the ICD. If reactivation does not occur, use manual defibrillator in accordance with *Tachycardia* protocol.

**MC**
- **Midazolam** 0.1 mg/kg SLOW IVP/IN/IM/IO. Maximum single dose is 5 mg. (Paramedic may perform without consultation.) IN administration max 1 mL per nare. IM administration requires all clinicians to obtain consultation.
- **Midazolam** 0.1 mg/kg SLOW IV/IO over 1–2 minutes. Maximum single IV/IN/IO dose 2 mg. Maximum total dose 5 mg. IN administration max 1 mL per nare. If IV cannot be established, administer 0.2 mg/kg IM. Max single IM dose is 5 mg. (IM requires all clinicians to obtain medical consultation.) Maximum total dose 5 mg.
- Consult a Pediatric Base Station for children (who have not reached their 18th birthday) with an ICD device delivering shock therapy or malfunctioning.

**Clinical Pearls**
- If the patient is in cardiac arrest, perform CPR and use the AED as appropriate despite the patient's ICD, which may or may not be delivering shocks.
- If the patient has a combination ICD and pacemaker, deactivating the ICD may or may not deactivate the pacemaker.
Indications

- Patient with acute coronary syndrome (ACS) symptoms, including angina or angina equivalents such as shortness of breath, chest, epigastric, arm or jaw pain or discomfort, diaphoresis, and/or nausea and meets one of the following criteria on diagnostic quality EKG:
  - New ST elevation of 1 mm (or greater) in two or more contiguous leads
  - Posterior MI: ST depression greater than 1 mm in V1-V3

BLS

- Not applicable; ALS protocol only

ALS

- Aspirin, nitroglycerin, Pain Management protocol.
- STEMI patients are Priority 1 and require transmission of an EKG and medical consultation with clear communication of an incoming “STEMI Alert” patient.
- STEMI patients shall be transported to the closest cardiac interventional center by air or ground as long as the delivery time is not more than 45 minutes greater than transport to the nearest ED.
- STEMI patients may bypass the ED and go directly to the cardiac catheterization lab, as directed by the receiving ED physician.
- If the patient cannot be delivered to a cardiac interventional center within the allotted time, complete the Fibrinolytic Therapy Checklist for STEMI.
  - If patient meets all of the criteria for fibrinolytic therapy, transport to the closest ED.
  - If the patient does not meet all of the criteria for fibrinolytic therapy, consult with the nearest cardiac interventional center and the closest ED to determine the most appropriate receiving facility.
- If inferior MI, obtain a right-sided EKG (V4R) to evaluate for right ventricular involvement. If ST elevation is noted in V4R, do not give nitrates due to risk for hypotension.
- If hypotensive with clear lung sounds, administer Lactated Ringer’s 250 mL IV.
- If the patient does not have ST elevations greater than 1 mm in two contiguous leads on 12-lead EKG, the patient shall be transported to the closest appropriate ED.
For STEMI patients with hypotension, obtain medical consultation for Lactated Ringer’s doses greater than 500 mL.

Left Bundle Branch Block (LBBB) / Paced Rhythm plus any of the following requires consultation with the closest appropriate EMS base station or cardiac interventional center:
- Patient presents in cardiogenic shock
- EKG shows excessive ST segment elevation greater than 5mm
- EKG shows ST segment deviation (elevation or depression) in the same direction as the QRS complex.

Other high-risk EKG findings that require consultation with EMS base station or cardiac interventional center:
- Wellens’ wave: biphasic T waves or deeply inverted T waves in the precordial leads (V2-V3 +/- V4).
- ST segment elevation in aVR: with coexisting multi-lead ST segment depression
- Hyperacute T waves: peaked, broad-based T waves

For STEMI patients who have not reached the 18th birthday, consult a pediatric base station for management and destination.

### Clinical Pearls
- Approximately 40% of inferior STEMIs have right ventricular involvement, which predisposes these patients to hypotension and increased mortality. Avoid nitrates if there is suspected right ventricular involvement. Clinically, patients with RV involvement often have clear lung sounds, hypotension, and JVD.
- ST depressions in V1-V3 may indicate anterior ischemia or a posterior STEMI. Obtain a posterior EKG to differentiate between these two conditions.

### Fibrinolytic Therapy Checklist for STEMI

Use this checklist if a STEMI patient cannot be delivered to a Cardiac Interventional Center within 45 minutes greater than transport to the nearest ED. All of the “YES” boxes and all of the “NO” boxes must be checked before a patient should be transported to the nearest emergency department.

**INCLUSION CRITERIA**
(All of the “YES” boxes must be checked)

YES
- 18 years of age or older
- Signs and symptoms of STEMI
- Patient cannot be delivered to a Cardiac Interventional Center within 45 minutes greater than transport to the nearest ED

**EXCLUSION CRITERIA**
(If any of the “NO” are unchecked, clinician must consult with a Cardiac Interventional Center and nearest ED to determine most appropriate receiving facility.)

PATIENT HAS NO:
- Active internal bleeding (e.g., GI or urinary bleeding within the last 21 days)
- Known bleeding disorder
- Within 3 months of intracranial surgery, serious head trauma, or stroke
- Within 14 days of major surgery or serious trauma
- History of intracranial hemorrhage
- Witnessed seizure at onset
- History of cancer of the brain
Indications

- Adult patients who have an implantable ventricular assist device (VAD), including left ventricular assist device (LVAD), right ventricular assist device (RVAD), or biventricular assist device (BiVAD) and have symptoms of cardiovascular compromise or cardiac arrest

<table>
<thead>
<tr>
<th>BLS</th>
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<tbody>
<tr>
<td></td>
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<tr>
<td><strong>Assess level of consciousness and vitals</strong></td>
</tr>
<tr>
<td>- Note: most VAD patients will not have a palpable pulse or detectable systolic and diastolic blood pressures due to the nature of the pump</td>
</tr>
<tr>
<td>- An automated blood pressure cuff may be used which may obtain a mean arterial pressure (MAP). The normal range for MAP is between 60 and 90 mmHg.</td>
</tr>
<tr>
<td>- Check for breathing and assist ventilation if necessary.</td>
</tr>
<tr>
<td>- Assess for perfusion: check skin color, skin temperature, capillary refill, MAP, and mental status.</td>
</tr>
<tr>
<td><strong>Altered mental status/adequate perfusion.</strong> If the patient has altered mental status, but has other signs of adequate perfusion, assess for causes of altered mental status (4.4).</td>
</tr>
<tr>
<td>- Check blood glucose and refer to Hypo/Hyperglycemia (4.7).</td>
</tr>
<tr>
<td>- If concern for overdose, refer to Overdose/Poisoning-Adult (7.7-A).</td>
</tr>
<tr>
<td><strong>Unresponsive/abnormal perfusion.</strong> If the patient is unconscious/unresponsive, not breathing, has delayed capillary refill, and unable to obtain a MAP, initiate manual chest compressions and ventilations per Cardiac Arrest-Adult protocol (3.4-A).</td>
</tr>
<tr>
<td>- Listen for pump sound “hum” or “whirling sound” over the chest.</td>
</tr>
<tr>
<td>- Check power and connections from the controller to the batteries and driveline.</td>
</tr>
<tr>
<td>- Contact patient’s VAD coordinator, using phone number on the device, and/or VAD-trained companion, who will likely be the best source of information for need to return to tertiary care center.</td>
</tr>
<tr>
<td>- Johns Hopkins (cell phone): 410-382-6885</td>
</tr>
<tr>
<td>- MedStar (pager): 202-801-9796</td>
</tr>
<tr>
<td>- University of Maryland (phone): 410-328-4903</td>
</tr>
<tr>
<td>- Change VAD batteries and/or controller, if indicated.</td>
</tr>
<tr>
<td>- If VAD batteries require changing, only change ONE at a time.</td>
</tr>
<tr>
<td>- Transport the “backup bag” with batteries and a second controller with the patient.</td>
</tr>
<tr>
<td>- For VAD-related complications or suspected cardiac/respiratory conditions: transport to the medical facility where the VAD was placed, if patient’s clinical condition and time allows.</td>
</tr>
<tr>
<td>- For all other conditions: transport to closest appropriate emergency department without manipulating the device.</td>
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</tbody>
</table>
3.12 Ventricular Assist Device (VAD) Protocol (continued)

**Clinical Pearls**
- LVAD patients are preload dependent and afterload sensitive.
- VAD patients require both anticoagulation and antiplatelet medication to prevent pump thrombosis, both of which make them high risk for life-threatening bleeding.
- VAD patients are at increased risk of sepsis due to driveline infection.
- Chest compressions must be manual (using hands only); automated chest compression devices may increase the risk of VAD dislodgement.
- VAD patients can be cardioverted and defibrillated without any changes or adjustments to the VAD.
- Always ask VAD patients if they have had any alarms and if their parameters are within normal range.

**ALS**
- Initiate cardiac monitoring and end-tidal CO$_2$ monitoring.
- Establish IV/IO access
- Obtain 12-lead EKG.
- For hypoperfusion (MAP less than 60 mmHg) due to hypovolemia with a functioning VAD OR ventricular arrhythmias associated with low-flow alarms:
  - Administer *Lactated Ringer’s* fluid boluses (250 mL) at a time, to achieve a target MAP of 60 mmHg, if there are no signs of pulmonary edema.
  - A maximum of 1 liter *Lactated Ringer’s* may be given over 15 minutes, using a push-pull method of drawing up the fluid and pushing it through the IV.
- Defibrillate or cardiovert, as appropriate for the patient’s rhythm. The controller or batteries should not be disconnected to deliver electrical therapy.

**MC**
- Additional IV fluid beyond 1 liter of *Lactated Ringer’s* requires medical consultation.
Indications
- Injuries or burns in a pattern suggesting intentional infliction
- Injuries in various stages of healing or injuries scattered over multiple areas of the body
- Patient, parent, or caregiver responding in an inappropriate manner to the situation
- Malnutrition or extreme lack of cleanliness of the patient or environment
- Bulging of fontanels and altered mental status in infants

Stabilize and treat injuries according to the appropriate protocol
- Discourage patient from washing if sexual abuse is suspected
- Document the following in the patient care report:
  - All statements made by the patient, parent, or caregiver; include verbatim statements in quotation marks
  - Any abnormal behavior on the part of the patient, parent, or caregiver
  - The condition of the environment and other residents present
  - Document the time the police or social service agency was notified along with the name and identifier, if possible
  - Document the name of the receiving health care clinician (RN, PA, or MD)
- Report all cases of suspected child or vulnerable adult abuse or neglect directly to either the local police or social service agency, as required by law. Do not initiate the report in front of the patient, parent, or caregiver.

Clinical Pearls
- Maryland EMS clinicians are protected from liability if they make a report of child or vulnerable adult abuse and neglect in good faith.
4.2-A Agitation – Adult

Indications

- **Mild symptoms** – Patient is agitated but cooperative and making rational decisions. No immediate concern for patient or clinician safety.
- **Moderate symptoms** – Patient is irrational and exhibiting behavior that puts themselves or clinicians at risk.
- **Severe symptoms** – Patient is physically violent and presents an immediate and imminent threat to themselves or others.

**Maintain scene safety and have a low threshold for requesting law enforcement.**
- Assess patient’s capacity and risk for self-harm
- Place the patient in supine position (face up) as soon as practical.
- Consider causes of agitation (medical, head trauma, psychiatric, drug/alcohol ingestion)

**Mild Agitation**
- Attempt verbal de-escalation and provide emotional support by using SAFER Model:
  - Stabilize the situation by containing and lowering the stimuli.
  - Assess and acknowledge the crisis.
  - Facilitate the identification and activation of resources (chaplain, family, friends, or police).
  - Encourage patient to use resources and take actions in their best interest.
  - Recovery or referral – leave patient in care of responsible person/professional or transport.

**Moderate Agitation**
- Evaluate for source of agitation and treat as follows:
  - Medical delirium (e.g., infection)
    - *Droperidol* 2.5 mg IM (1.25 mg for patients 69 years of age or older)
  - Psychiatric emergency (e.g., schizophrenia, patient off medications)
    - *Droperidol* 2.5 mg IM (1.25 mg for patients 69 years of age or older)
  - Drug or alcohol ingestion
    - *Midazolam* 5 mg IM/IV (2.5 mg for patients 69 years of age or older)
  - Head injury
    - *Midazolam* 5 mg IM/IV (2.5 mg for patients 69 years of age or older)
  - Unknown or other
    - *Midazolam* 5 mg IM/IV (2.5 mg for patients 69 years of age or older)

**Severe Agitation**
- *Midazolam* 5 mg IM/IV (2.5 mg for patients 69 years of age or older)
  OR
- *Ketamine* 1 mg/kg IV/IO (max 100 mg) or *ketamine* 4 mg/kg IM (max 400 mg) if there is immediate and imminent danger to patient or EMS

**Following sedation, perform the following interventions:**
- Initiate cardiac monitoring, continuous ETCO₂, pulse oximetry.
- Obtain 12-lead EKG to evaluate for prolonged QTc.
- Evaluate for trauma.
- Check blood glucose.
- Check temperature and initiate passive cooling measures (e.g., cold packs), as appropriate.
- If tachycardic or hyperthermic, initiate *Lactated Ringer’s* 20 mL/kg fluid bolus.
- Apply physical restraints as indicated in Physical Restraint protocol, only when imminent and immediate danger to self or others.
**Agitation – Adult (Continued)**

**MC**

- *Ketamine* 1 mg/kg IV/IO (max 100 mg) OR *ketamine* 4 mg/kg IM (max 400 mg) requires medical consultation unless immediate and imminent danger to patient or clinicians
- Additional doses of medication (beyond first dose): *droperidol*, *ketamine*, or *midazolam* require medical consultation
- *Diphenhydramine* 25-50 mg IM/IV may be administered if a dystonic reaction occurs (associated with *droperidol*)

**Clinical Pearls**

**Ketamine**:
- *Ketamine* should be avoided if possible in agitated elderly patient due to the risk for over-sedation and apnea.
- Advanced airway equipment, BVM, oxygen, and suction must be immediately available at all times for patients receiving *ketamine*.
- All patients that receive *ketamine* must be transported with at least two EMS clinicians, one of which must be an ALS clinician.

**Severe agitation**:
- Patients with severe agitation should not receive *droperidol* or *diphenhydramine* (Benadryl®) for sedation. These medications may worsen anticholinergic crisis.

**Droperidol**:
- *Droperidol* may prolong the QTc interval, which increases the risk of cardiac dysrhythmia. Do not administer *droperidol* if QTc is known to be 440 ms or greater.
- Dystonic reactions (extrapyramidal symptoms) may occur after administration of *droperidol*.
- *Droperidol* is contraindicated for pregnant patients.
Indications

- **Mild symptoms** – Patient is agitated but cooperative and making rational decisions. No immediate concern for patient or clinician safety.
- **Moderate symptoms** – Patient is irrational and exhibiting behavior that puts themselves or clinicians at risk.
- **Severe symptoms** – Patient is physically violent and presents an immediate and imminent threat to themselves or others.

**BLS**

- Maintain scene safety and have a low threshold for requesting law enforcement.
- Assess patient's capacity and risk for self-harm.
- Place the patient in a supine position (face up) as soon as practical.
- Consider causes of agitation (medical, head trauma, psychiatric, drug/alcohol ingestion).

**Mild Agitation**

- Attempt verbal de-escalation and provide emotional support by using SAFER Model:
  - **S**tabilize the situation by containing and lowering the stimuli.
  - **A**ssess and acknowledge the crisis.
  - **F**acilitate the identification and activation of resources (chaplain, family, friends, or police).
  - **E**ncourage patient to use resources and take actions in their best interest.
  - **R**ecovery or referral – leave patient in care of responsible person/professional or transport.

**ALS**

- **Moderate Agitation**
  - Less than 5 years of age: no medication indicated
  - 5-12 years of age: medical consultation required for midazolam
  - 13-18 years of age: droperidol 2.5 mg IM or consult for midazolam 0.1 mg/kg IV or 0.2 mg/kg IM/IN (max 5 mg). IM route preferred.

- **Severe Agitation**
  - Less than 5 years of age: no medication indicated
  - 5-12 years of age: medical consultation required for midazolam or ketamine
  - 13-18 years of age:
    - **Ketamine** 1 mg/kg IV/IO (max 100 mg) or ketamine 4 mg/kg IM (max 400 mg) if there is immediate and imminent danger to patient or EMS OR
    - **Midazolam** 0.1 mg/kg IV or 0.2 mg/kg IM/IN (max 5 mg). IM route preferred.

- Following sedation, perform the following interventions:
  - Initiate cardiac monitoring, continuous ETCO₂, pulse oximetry.
  - Obtain 12-lead EKG to evaluate for prolonged QTc.
  - Evaluate for trauma.
  - Check blood glucose.
  - Check temperature and initiate passive cooling measures (e.g., cold packs), as appropriate.
  - If tachycardic or hyperthermic, initiate Lactated Ringer’s 20 mL/kg fluid bolus.
  - Apply physical restraints as indicated in Physical Restraint protocol, only when imminent and immediate danger to self or others.
Less than 13 years of age:
- **Ketamine** 1 mg/kg IV/IO (max 100 mg) or **ketamine** 4 mg/kg IM (max 400 mg) OR
- **Midazolam** 0.1 mg/kg IV or 0.2 mg/kg IM/IN (max 5 mg). IM route preferred.
13-18 years of age: **Ketamine** 1 mg/kg IV/IO (max 100 mg) OR **Ketamine** 4 mg/kg IM (max 400 mg) requires medical consultation unless immediate and imminent danger to patient or clinicians

Additional doses of droperidol, ketamine, or midazolam require medical consultation.

**Diphenhydramine** 1 mg/kg IM/IV may be administered if a dystonic reaction occurs (associated with droperidol).

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**Clinical Pearls**

**Ketamine:**
- Advanced airway equipment, BVM, oxygen, and suction must be immediately available at all times for patients receiving ketamine.
- All patients that receive ketamine must be transported with at least two EMS clinicians, one of which must be an ALS clinician.

**Severe agitation:**
- Patients with severe agitation should not receive droperidol or diphenhydramine (Benadryl®) for sedation. These medications may worsen anticholinergic crisis.

**Droperidol:**
- **Droperidol** may prolong the QTc interval, which increases the risk of cardiac dysrhythmia. Do not administer droperidol if QTc is known to be 440 ms or greater.
- Dystonic reactions (extrapyramidal symptoms) may occur after administration of droperidol.
- **Droperidol** is contraindicated for pregnant patients.
Allergic Reaction – Adult

Indications

- **Mild symptoms**: localized swelling and itching at the site
- **Moderate symptoms**: hives and/or mild wheezing
- **Severe symptoms**: diffuse wheezing, pharyngeal swelling, dyspnea, hypoperfusion, abnormal skin color, stridor, and/or loss of peripheral pulses (Refer to Anaphylaxis protocol)

BLS

- **Mild symptoms** (if history of life-threatening allergic reaction to same allergen)
  - Epinephrine auto-injector (BLS) 0.3 mg IM OR
  - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.5 mg IM.

- **Moderate symptoms**
  - Epinephrine auto-injector (BLS) 0.3 mg IM OR
  - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.5 mg IM.
  - Albuterol (BLS) inhaler (2 puffs inhaled) or nebulized albuterol (BLS). May repeat dose one time, as needed, within 30 minutes.

ALS

- **Mild symptoms**
  - Diphenhydramine 25 mg SLOW IV or IM.
  - Epinephrine (1 mg/mL) 0.5 mg IM if patient has a history of life-threatening allergic reaction to the same allergen.

- **Moderate symptoms**
  - Epinephrine (1 mg/mL) 0.5 mg IM. May repeat every 5 minutes, for a total of 3 doses, for recurrent or worsening symptoms.
  - Establish IV access.
  - Diphenhydramine 50 mg SLOW IVP or IM.
  - Albuterol 2.5 mg and ipratropium 500 mcg nebulizer. May repeat albuterol one time for recurrent or worsening symptoms.

MC

- Additional doses of epinephrine auto-injector, epinephrine, albuterol, ipratropium, diphenhydramine beyond those listed above require medical consultation.

Clinical Pearls

- Re-check dosing and concentration of epinephrine prior to administration.
- Epinephrine 1 mg/mL (previously known as 1:1,000) is appropriate for the IM route only.
- Epinephrine should never be given by IV route, except for an epinephrine infusion for patients in anaphylaxis or for patients in cardiac arrest.
**Indications**

- **Mild symptoms:** localized swelling and itching at the site
- **Moderate symptoms:** hives and/or mild wheezing
- **Severe symptoms:** diffuse wheezing, pharyngeal swelling, dyspnea, hypoperfusion, abnormal skin color, stridor, and/or loss of peripheral pulses (Refer to Anaphylaxis protocol)

**BLS**

- **Mild symptoms** (if history of life-threatening allergic reaction to same allergen)
  - Less than 5 years of age: pediatric epinephrine auto-injector (BLS) 0.15 mg IM **OR**
    - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.15 mg IM
  - 5 years of age or greater: epinephrine auto-injector (BLS) 0.3 mg IM **OR**
    - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.5 mg IM

- **Moderate symptoms**:
  - Less than 5 years of age:
    - Pediatric epinephrine auto-injector (BLS) 0.15 mg IM **OR**
    - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.15 mg IM
  - 5 years of age or greater:
    - Epinephrine auto-injector (BLS) 0.3 mg IM **OR**
    - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.5 mg IM
  - **Albuterol (BLS)** inhaler (2 puffs inhaled) or albuterol (BLS) nebulizer. May repeat dose one time, as needed, within 30 minutes.
    - For infants and children less than 2 years of age, administer nebulized albuterol (BLS) 1.25 mg.
    - For patients 2 years of age or greater, administer nebulized albuterol (BLS) 2.5 mg.

**ALS**

- **Mild symptoms**
  - **Diphenhydramine** 1 mg/kg SLOW IV or IM. Maximum single dose 25 mg **OR**
  - **Epinephrine** if history of life-threatening allergic reaction to the same allergen.
    - Less than 5 years of age: epinephrine (1 mg/mL) 0.15 mg IM
    - 5 years of age or greater: epinephrine (1 mg/mL) 0.5 mg IM

- **Moderate symptoms**
  - **Epinephrine**
    - Less than 5 years of age: epinephrine (1 mg/mL) 0.15 mg IM
    - 5 years of age or greater: epinephrine (1 mg/mL) 0.5 mg IM
  - Establish IV access.
  - If age-related vital signs and patient’s condition indicate hypoperfusion, administer initial fluid bolus of 20 mL/kg LR IV. If patient’s condition does not improve, administer the second bolus of fluid at 20 mL/kg LR IV.
  - **Diphenhydramine** 1 mg/kg SLOW IVP or IM. Maximum single dose 50 mg.
  - **Albuterol / ipratropium** nebulized.
    - For an infant less than 1 year of age: albuterol 1.25 mg via nebulizer; ipratropium is contraindicated.
    - For a child 1 year of age or greater, but less than 2 years of age: albuterol 1.25 mg and ipratropium 250 mcg.
    - For a patient 2 years of age or greater: albuterol 2.5 mg and ipratropium 500 mcg.
    - May repeat albuterol one time for recurrent or worsening symptoms.

**MC**

- Additional doses of epinephrine auto-injector, epinephrine, albuterol, ipratropium, diphenhydramine beyond those listed above require medical consultation.
Altered Mental Status

4.4

Check Pulse

Pulse Present

Assess vital signs
Physical exam
Assess for signs of trauma
Check blood glucose

Suspected OD/Tox Environmental

Blood glucose < 70 or > 300

Recent seizure activity, history of seizures

Fever/ tachycardia/ low BP/ suspected sepsis

Numbness/ weakness/ speech abnormality/ suspected stroke

Tox/ Environmental (7.1-7.10)

Hypo/ Hyperglycemia (4.7)

Seizures (4.11)

Sepsis (4.12)

Stroke (4.14)

No Pulse

Cardiac Arrest (3.4)
**Indications**

- Acute onset of severe illness after exposure to a known allergen with **two or more** of the following:
  - Urticaria (hives) or acute swelling of the mucosa (e.g., tongue, airway, stridor, lips)
  - Respiratory compromise
  - Hypotension
  - GI symptoms, such as persistent nausea/vomiting, abdominal pain, or diarrhea
- Acute onset of severe illness after exposure to a known allergen with hypotension

**BLS**

- **Epinephrine auto-injector (BLS)** 0.3 mg **OR**
- If BLS epinephrine OSP approved, **epinephrine (BLS)** (1 mg/mL) 0.5 mg IM in the lateral thigh.
- **Albuterol (BLS)** inhaler (2 puffs inhaled) or **albuterol (BLS)** 2.5 mg nebulized for wheezing/bronchospasm/shortness of breath. May repeat dose one time, as needed, within 30 minutes.

**ALS**

- Administer **epinephrine** (1 mg/mL) 0.5 mg IM. May repeat epinephrine IM every 5 minutes to a maximum of 3 doses for persistent severe reactions.
- Establish IV/IO access.
- For patients who are in extremis with severe hypotension or impending respiratory failure, initiate an epinephrine infusion (after having administered 3 doses of IM epinephrine) as follows:
  - Add 1 mg of epinephrine (either 1 mg/mL or 0.1 mg/mL) in a 100 mL bag of LR or NS
  - Use a Microdrip set (60 drops/mL) for infusion administration
  - Adult epinephrine infusion dosage:
    - Administer infusion through a free-flowing IV, ideally 20 gauge or larger, or by IO
    - Start infusion at 1 mL/min (60 drops/min) IV/IO
    - Check blood pressure every 5 minutes. If MAP is less than 65 mmHg or systolic blood pressure is less than 90 mmHg, increase to a maximum rate of 2 mL/min (120 drops/min).
- Additional treatments **after** administration of the initial dose of epinephrine:
  - **Albuterol** 2.5 mg nebulized and **ipratropium** (Atrovent®) 500 mcg nebulized; may repeat albuterol nebulized 2.5 mg one time.
  - **Diphenhydramine** 50 mg SLOW IVP or IM
  - Administer 20 mL/kg bolus IV/IO for hypotension (MAP less than 65 mmHg or systolic blood pressure less than 90 mmHg)
  - **Dexamethasone** 10 mg IV/IO

**MC**

- Additional doses of epinephrine auto-injector, epinephrine, albuterol, ipratropium, diphenhydramine beyond those listed above require medical consultation.
- If blood pressure goals are not met upon reaching epinephrine infusion rate of 2 mL/min (120 drops/min), obtain medical consultation.

**Clinical Pearls**

- Re-check dosing and concentration of epinephrine prior to administration.
- **Epinephrine** 1 mg/mL (previously known as 1:1,000) is appropriate for the IM route only.
- Epinephrine should never be given by IV route, except for an epinephrine infusion for patients in anaphylaxis or for patients in cardiac arrest.
Anaphylaxis – Pediatric

**Indications**
- Acute onset of severe illness after exposure to a known allergen with **two or more** of the following:
  - Urticaria (hives) or acute swelling of the mucosa (e.g., tongue, airway, stridor, lips)
  - Respiratory compromise
  - Hypotension
  - GI symptoms, such as persistent nausea/vomiting, abdominal pain, or diarrhea
- Acute onset of severe illness after exposure to a known allergen with hypotension

**BLS**
- **Epinephrine (BLS)**
  - Less than 5 years of age:
    - Pediatric epinephrine auto-injector (BLS) 0.15 mg IM in the lateral thigh OR
    - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.15 mg IM
  - 5 years of age or greater:
    - Epinephrine auto-injector (BLS) 0.3 mg IM in the lateral thigh OR
    - If BLS epinephrine OSP approved, epinephrine (BLS) (1 mg/mL) 0.5 mg IM
- **Albuterol (BLS)** – for wheezing/bronchospasm/shortness of breath.
  - Less than 2 years of age: albuterol (BLS) inhaler (2 puffs) inhaled or albuterol (BLS) 1.25 mg nebulized. May repeat dose once, as needed, within 30 minutes.
  - 2 years of age or greater: albuterol (BLS) inhaler (2 puffs) inhaled or albuterol (BLS) 2.5 mg nebulized. May repeat dose once, as needed, within 30 minutes.

**ALS**
- **Administer epinephrine IM.** May repeat every 5 minutes for a total of 3 doses.
  - Less than 5 years of age: epinephrine (1 mg/mL) 0.15 mg IM in the lateral thigh.
  - 5 years of age or greater: epinephrine (1 mg/mL) 0.5 mg IM in the lateral thigh.
- Establish IV/IO access.
- Additional treatments to consider after administration of the initial dose of epinephrine:
  - Albuterol and ipratropium (Atrovent®) via nebulizer:
    - Less than 1 year of age: albuterol 1.25 mg; ipratropium is contraindicated.
    - Greater than 1 year of age but less than 2 years of age: albuterol 1.25 mg and ipratropium 250 mcg.
    - For a patient 2 years of age or greater: albuterol 2.5 mg and ipratropium 500 mcg.
    - For all age groups, one additional dose of albuterol only may be given via nebulizer.
  - Diphenhydramine 1 mg/kg SLOW IVP or IM to a maximum of 50 mg
  - Administer 20 mL/kg bolus for hypotension; systolic blood pressure less than 70 x 2 (age in years)
  - Dexamethasone 0.5 mg/kg to a maximum of 10 mg IV/IO

**MC**
- Additional doses of pediatric epinephrine auto-injector, epinephrine, albuterol, ipratropium, diphenhydramine beyond those listed above require medical consultation.
- Consider pediatric epinephrine infusion for refractory anaphylactic shock.

**Clinical Pearls**
- Re-check dosing and concentration of epinephrine prior to administration.
- Epinephrine 1 mg/mL (previously known as 1:1,000) is appropriate for the IM route only.
- Epinephrine should never be given by IV route, except for an epinephrine infusion for patients in anaphylaxis or for patients in cardiac arrest.

Release Date July 1, 2022

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**Indications**
- Infant or child less than 2 years of age
- Episode that is frightening to the observer that includes some combination of the following:
  - Apnea
  - Skin color change (cyanosis, pallor, erythema)
  - Marked change in muscle tone
  - Choking or gagging not associated with feeding or witnessing foreign body aspiration

**BLS**
- Perform assessment using the Pediatric Assessment Triangle
- Obtain a description of the event including nature, duration, and severity
- Assess the environment for possible causes
- When obtaining the medical history, include questions to identify any: current medications, chronic diseases, current or recent infections, evidence of seizure activity, gastroesophageal reflux, or recent trauma
- Apply oxygen and be prepared to support ventilation during transport

**ALS**
- Place patient on cardiac monitor
- Establish IV/IO access only if required by patient’s clinical condition

**MC**
- If the parent or guardian refuses medical care or transport, clinician SHALL contact a Pediatric Base Station physician

**Clinical Pearls**
- Most patients will appear stable upon assessment. However, this episode may be a sign of serious underlying illness or injury. All suspected ALTE/BRUE patients should be transported for further medical evaluation.
**Indications**
- Blood glucose less than 70 mg/dL or greater than 300 mg/dL
- Patient-reported low or high blood glucose
- Diabetic patients with other medical symptoms (e.g., vomiting)
- Altered mental status
- Alcohol intoxication, suspected
- Seizure
- Stroke symptoms
- Unresponsive patients
- Cardiac arrest

**BLS**
- Check blood glucose level
- If blood glucose is less than 70 mg/dL, administer 10-15 grams of oral glucose between the patient’s gum and cheek.
- Administer additional dose of 10-15 grams of oral glucose if not improved after 10 minutes.

**ALS**
- **HYPOglycemia**: If blood glucose is less than 70 mg/dL, administer 10% dextrose in 50 mL (5 gram) boluses, 1 minute apart, to a maximum of 250 mL OR 25 grams of 50% dextrose IVP, until:
  - the patient has a return to normal mental status, and
  - the patient’s blood glucose is at least 90 mg/dL
- If patient has persistently altered mental status and blood glucose less than 90 mg/dL despite treatment, repeat dosing regimen above.
- If unable to initiate an IV and blood glucose is less than 70 mg/dL, administer glucagon 1 mg IM/IN.
  - If the patient has persistently altered mental status and blood glucose less than 90 mg/dL at 15 minutes, transport to the hospital should not be delayed.
- **HYPERglycemia**: If blood glucose is greater than 300 mg/dL, administer 10 mL/kg Lactated Ringer’s bolus unless rales, wheezing, pedal edema, or history of renal failure or CHF is present.

**MC**
- Not applicable
HYPOglycemia/HYPERglycemia – Pediatric

**Indications**
- Blood glucose less than 70 mg/dL or greater than 300 mg/dL
- Patient-reported low or high blood glucose
- Diabetic patients with other medical symptoms (e.g., vomiting)
- Altered mental status
- Alcohol intoxication, suspected
- Seizure
- Stroke symptoms
- Unresponsive patients
- Cardiac arrest
- Pediatric bradycardia

**BLS**
- Check blood glucose level
- If blood glucose is less than 70 mg/dL, administer 10-15 grams of *oral glucose* between the patient’s gum and cheek.
- Administer additional dose of 10-15 grams of *oral glucose* if not improved after 10 minutes.

**ALS**
- **Patient less than 28 days:** If blood glucose is less than 40 mg/dL, administer 2 mL/kg of *10% dextrose IV/IO*. Recheck glucose after first dose.
- **Patient 28 days or greater until the 18th birthday:** If blood glucose is less than 70 mg/dL, administer 2–4 mL/kg of *10% dextrose IV/IO* to a maximum of 25 grams. Recheck glucose after first dose.
  - If unable to start IV and blood glucose is less than 70 mg/dL:
    - **28 days–4 years of age:** Glucagon 0.5 mg IM/IN.
    - **5 years of age until 18th birthday:** Glucagon 1 mg IM/IN. Recheck blood glucose. If the patient has persistently altered mental status and blood glucose less than 90 mg/dL at 15 minutes, transport to the hospital should not be delayed.

**HYPOglycemia**
- **Patient less than 28 days:** If blood glucose is less than 40 mg/dL, administer 2 mL/kg of *10% dextrose IV/IO*. Recheck glucose after first dose.

**HYPERglycemia**
- If blood glucose is greater than 300 mg/dL, no fluid bolus is indicated for pediatric patients. Consult with the receiving hospital.

**MC**
- **Patient less than 28 days:** If blood glucose is less than 40 mg/dL, obtain medical consultation to administer second dose of *10% dextrose IV/IO*.
- **Patient 28 days – 18 years of age:** If blood glucose is less than 70 mg/dL, obtain medical consultation to administer second dose of *10% dextrose IV/IO*.
**Indications**
- Renal failure or chronic kidney disease patients or history of poorly-functioning kidneys
- Renal dialysis patients who are hemodynamically unstable or patients suspected of having elevated potassium with EKG changes (peaked T waves, wide QRS complexes, or bradycardia)
- Crush syndrome (entrapped patients with prolonged extrication time)

**BLS**
- Place patient in position of comfort.
- Assess and treat for shock, if indicated.

**ALS**
- Establish IV/IO access.
- *Lactated Ringer’s per Shock: Hypoperfusion* protocol
- If the patient is bradycardic, refer to *Bradycardia* protocol.
- *Calcium chloride* 0.5–1 gram SLOW IVP over 3–5 minutes. Maximum dose 1 gram or 10 mL.
- *Sodium bicarbonate* 50 mEq IV over 5 minutes
- **Crush syndrome or patients with functional kidneys by history**
  - *Sodium bicarbonate* 50 mEq SLOW IV over 5 minutes and then initiate drip of *sodium bicarbonate* 100 mEq in 1,000 mL to run over 30–60 minutes

**MC**
- Albuterol 20 mg nebulized (high dose)

**Clinical Pearls**
- Flush IV with 5 mL of *Lactated Ringer’s* between *calcium chloride* and *sodium bicarbonate* administration to avoid precipitation in the IV.
Indications
- Renal failure or chronic kidney disease patients or history of poorly-functioning kidneys
- Renal dialysis patients who are hemodynamically unstable or patients suspected of having elevated potassium with EKG changes (peaked T waves, wide QRS complexes, or bradycardia)
- Crush syndrome (entrapped patients with prolonged extrication time)

BLS
- Place patient in position of comfort.
- Assess and treat for shock, if indicated.

ALS
- Establish IV/IO access.
- *Lactated Ringer’s per Shock: Hypoperfusion protocol*
- If the patient is bradycardic, refer to *Bradycardia protocol*.
- *Calcium chloride* 20 mg/kg (0.2 mL/kg) slow IVP/IO (50 mg/min). Maximum dose 1 gram or 10 mL.

MC
- *Albuterol* nebulized
  - For patients less than 2 years of age, administer *albuterol* 1.25 mg.
  - For patients 2 years of age or greater, administer *albuterol* 2.5 mg.
- **Crush syndrome or patients with functional kidneys by history**
  - *Sodium bicarbonate* 1 mEq/kg IV over 5 minutes. Maximum dose 50 mEq. For patients less than 1 year of age, must be diluted (1:1) with *Lactated Ringer’s*.

Clinical Pearls
- Flush IV with 5 mL of *Lactated Ringer’s* between *calcium chloride* and *sodium bicarbonate* administration to avoid precipitation in the IV.
### Indications
- Nausea
- Vomiting
- Active motion sickness
- Medication side effect/complication
- Prevention of nausea/vomiting (e.g., penetrating eye injury, high risk for aspiration, opioid administration)

### BLS
- Place patient in position of comfort or in left lateral position, with consideration for *spinal motion restriction* if required.
- Allow patient to inhale vapor from an isopropyl alcohol wipe 3 times every 15 minutes, as needed and tolerated.

### ALS
- Establish IV access, if appropriate.
- *Lactated Ringer’s* fluid bolus, 20 mL/kg, if appropriate. Titrate to systolic blood pressure of 90 mmHg.
- *Ondansetron* 8 mg slow IV over 2-5 minutes OR 4-8 mg IM OR 8 mg orally disintegrating tablet (ODT). May repeat dose one time if needed.

### MC
- A third dose of *ondansetron* may be administered, with medical consultation, to a maximum of 24 mg.

### Clinical Pearls
- Higher doses of *ondansetron* may prolong the patient’s QTc interval and lead to cardiac dysrhythmias. Initiate cardiac monitoring when repeat doses are administered.
**Indications**
- Nausea
- Vomiting
- Active motion sickness
- Medication side effect/complication
- Prevention of nausea/vomiting (e.g., penetrating eye injury, high risk for aspiration, opioid administration)

**BLS**
- Place patient in position of comfort or in left lateral position, with consideration for *spinal motion restriction* if required.
- Allow patient to inhale vapor from an isopropyl alcohol wipe 3 times every 15 minutes, as needed and tolerated.

**ALS**
- Establish IV access, if appropriate.
- *Lactated Ringer’s* fluid bolus, 20 mL/kg, if age-related vital signs and patient’s condition indicate hypoperfusion. Titrate to systolic blood pressure of (2 x patient’s age in years) + 70 mmHg.
- **Ondansetron**
  - 28 days to 12-years-old: *Ondansetron* 0.1 mg/kg SLOW IV over 2–5 minutes
  - 13-18th birthday: *Ondansetron* 8 mg ODT OR 8 mg SLOW IV over 2–5 minutes
  - OR *ondansetron* 0.1 mg/kg IM, if IV access is not available (with max single dose of 8 mg)
  - May repeat dose one time if needed.

**MC**
- A third dose of *ondansetron* may be administered, with medical consultation, to a maximum total dose of 0.3 mg/kg or 24 mg, whichever is lower.

**Clinical Pearls**
- Higher doses of *ondansetron* may prolong the patient’s QTc interval and lead to cardiac dysrhythmias. Initiate cardiac monitoring when repeat doses are administered.
Indications

- Patient presents with a painful condition that would benefit from treatment with an analgesic. This includes DNR/MOLST patients and patients being pre-medicated for a painful procedure.
  - **Mild to moderate pain:** Pain rated in the 1-5 range on a scale of 1-10. Isolated musculoskeletal injuries such as sprains and strains.
  - **Moderate to severe pain:** Pain rated in the 5-10 range on a scale of 1-10.

- Measure level of pain. Ask patient to rate their pain on a scale from 0 (no pain) to 10 (worst pain imaginable). Patients who have a difficult time communicating their condition can be asked to rate their pain using the FACES scale.

## Pain Rating Scale

- **10 - Worst Pain Possible**
  - Unbearable
  - (Unable to do any activities because of pain)
- **9**
- **8 - Intense/Dreadful/Horrible**
  - (Unable to do most activities because of pain)
- **7 - Severe Pain**
- **6 - Miserable/Distressing**
  - (Unable to do some activities because of pain)
- **5 - Moderate Pain**
- **4 - Nagging/Uncomfortable**
  - (Can do most activities with rest periods)
- **3**
- **2 - Mild Pain**
  - Annoying
  - (Pain is present but does not limit activity)
- **1**
- **0 - No Pain**

- Allow patient to remain in position of comfort unless contraindicated by patient’s condition.
- **Mild to Moderate Pain (1-5 on FACES scale):**
  - *Acetaminophen* for mild to moderate pain:
    - FOUR unit doses of 160 mg/5 mL each for a total of 640 mg/20 mL OR
    - 325 mg pill or tablet X 2 for a total of 650 mg with sips of water as tolerated by the patient. No repeat doses.
Clinical Pearls

- Administration of pain medication does not eliminate the need for transport of the patient to the hospital to receive a comprehensive evaluation of the cause of their pain and appropriate definitive treatment.
- *Ketamine* is indicated only for musculoskeletal and back pain. Do not administer for chest pain, abdominal/flank pain, or for headaches.
- Chest pain that is thought to be due to acute coronary syndrome should initially be managed with nitroglycerin. If pain remains refractory to nitroglycerin, consider the use of opioid analgesia. Avoid opioids for patients with suspected exacerbation of congestive heart failure.
- Use opioid analgesia with caution in the management of the multiple trauma patient. Observe for evidence of hypotension and correct with fluid boluses. Reassess vital signs after administration of the medicine.
- Use analgesia with caution in the management of patients with altered mental status. Observe for respiratory depression and take steps to ensure a stable airway.
- Patients who have received a parenteral (IV/IO/IM/IN) dose of opioid, benzodiazepine, or ketamine from sending facility or ALS must be transported by ALS:
  - If any of the above medications were given within the past 1 hour OR
  - If the patient has an altered mental status without return to their baseline after receiving any of the above medications OR
  - If the patient has potential for respiratory compromise (RR less than 14, oxygen saturation less than 94%, clinician judgment) after receiving any of the above medications.
Indications
- Patient presents with a painful condition that would benefit from treatment with an analgesic. This includes DNR/MOLST patients and patients being pre-medicated for a painful procedure.
  - **Mild to moderate pain:** Pain rated in the 1-5 range on a scale of 1-10. Isolated musculoskeletal injuries such as sprains and strains. Pain related to childhood illnesses such as headache, ear infection, and pharyngitis.
  - **Moderate to severe pain:** Pain rated in the 5-10 range on a scale of 1-10.

- Measure level of pain. Ask older children to rate their pain on a scale from 0 (no pain) to 10 (worst pain imaginable). Young children can be asked to rate their pain using the FACES scale, which provides 5 levels of pain perception.

### Pain Rating Scale

- **Hurts Worse**
  - 10 - Worst Pain Possible
  - 9

- **Hurts Whole Lot**
  - 8 - Intense/Dreadful/Horrible
  - 7 - Severe Pain

- **Hurts Even More**
  - 6 - Miserable/Distressing
  - 5 - Moderate Pain

- **Hurts Little Worse**
  - 4 - Nagging/Uncomfortable
  - 3

- **Hurts Little Bit**
  - 2 - Mild Pain
  - 1

- **No Hurt**
  - 0 - No Pain

- Allow patient to remain in position of comfort unless contraindicated.

**Mild to Moderate Pain (1-5 on FACES scale):**
- **Acetaminophen** for mild to moderate pain per child or parent (1-5 on FACES scale)
  - Less than 3 months of age: Not indicated
  - 3 months to 2 years of age:

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<th>3 months</th>
<th>4-11 months</th>
<th>12-23 months</th>
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<td>1.25 mL</td>
<td>2.5 mL</td>
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</table>
### Clinical Pearls
- **Administration of pain medication** does not eliminate the need for transport of the patient to the hospital to receive a comprehensive evaluation of the cause of their pain and appropriate definitive treatment.
- **Ketamine** is indicated only for musculoskeletal and back pain. Do not administer for chest pain, abdominal/flank pain or for headaches.

### 4.10-P

#### 2–4 years: Unit dose 160 mg/5 mL
- 5–12 years: TWO unit doses of 160 mg/5 mL each for a total of 320 mg/10 mL
- 13 years and above: FOUR unit doses of 160 mg/5 mL each for a total of 640 mg/20 mL OR in a form of 325 mg pill or tablet x2 for a total of 650 mg with sips of water as tolerated by the patient.

### 4.10-P

#### Moderate to Severe Pain (5-10 on FACES scale):
- **Fentanyl** IN. If IN route not accessible, then use IV/IO/IM route.
  - Administer 1 mcg/kg to a maximum initial dose of 200 mcg (For IN route, dosing may be limited due to volume limitations - administration of max 1mL per nare).
  - Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of fentanyl 1 mcg/kg to a maximum dose of 200 mcg.
- OR
  - **Morphine** IV/IM 0.1 mg/kg maximum single dose of 20 mg.
  - Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of morphine 0.05 mg/kg to a maximum additional dose of 10 mg.
- OR
  - **Ketamine** IV/IO/IN/IM 0.2 mg/kg IV/IO over 1–2 minutes. Maximum single dose 20 mg.
  - Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of ketamine 0.2 mg/kg IV/IO over 1–2 minutes. Maximum single dose 20 mg.
  - If IV unavailable, administer 0.5 mg/kg IN/IM (If delivery device is available, divide administration of the dose equally between the nares to a maximum of 1 mL per nare). Reassess in 15 minutes. If pain remains moderate to severe, then administer a second dose of ketamine 0.5 mg/kg IN/IM.
- OR
  - **Ketorolac** IV/IM:
    - Patients who have not yet reached their 2nd birthday: Contraindicated.
    - Age 2 to patients who have not yet reached their 18th birthday: Administer 0.5 mg/kg IV to a maximum total dose of 15 mg. No repeat doses.
      - If IV is unavailable, administer 1 mg/kg IM to a maximum total dose of 30 mg. No repeat doses.

### 4.10-P

#### Obtain on-line medical direction for additional doses of **fentanyl**, **morphine**, or **ketamine**, based on the patient’s level of pain following two doses of medication. **Ketorolac** and **acetaminophen** are single dose-only medications.
Seizures – Adult

**Indications**
- Involuntary, repetitive muscle movements, usually abrupt in onset
- Observed seizure activity
- Decreased mental status
- Unconscious
- Incontinence
- Head injury

**Seizure has stopped**
- Identify and treat injuries
- Check blood glucose and treat per *Hypoglycemia* protocol
  - If blood glucose is less than 70 mg/dL, administer *oral glucose* paste (10-15 grams) between the gum and cheek. Administer single additional dose of *oral glucose* if not improved after 10 minutes.

**Active seizure**
- Do not restrain the patient or place any device into the patient’s mouth.
- Protect the patient from injury.
- Identify and treat potential underlying cause of seizure: epilepsy, head injury, hypoxia, hypoglycemia, hypoperfusion, infection, stroke, alcohol or drug abuse or withdrawal, head injury

**BLS**

**Active seizure**
- Check blood glucose and treat per *Hypoglycemia* protocol
- Establish IV/IO access. This should not delay *midazolam* IM/IN for patients with active seizures.
- **Midazolam:***
  - IM/IN: 5 mg IN or IM. IM route preferred.
  - IV/IO: 0.1 mg/kg in 2 mg increments SLOW IVP/IO over 1–2 minutes with a maximum single dose of 5 mg
  - Reduce *midazolam* dose by 50% for patients 69 and older
- **Diazepam:** if *midazolam* is not available, administer *diazepam* in 2.5 mg increments SLOW IVP/IO with a maximum single dose of 10 mg
- Pregnant patients: administer *midazolam* followed by *magnesium sulfate* 4 grams IV/IO over 10 minutes (mixed in 50–100 mL of approved diluent).

**MC**

- **Midazolam:** consult for additional doses of *midazolam* up to a maximum total dose of 10 mg
- **Diazepam:** IM doses of *diazepam* require consultation, except if suspected nerve agent exposure (CANA and nerve agent antidote kits may be used without consultation)
- **Magnesium sulfate:** consult for additional dose of *magnesium sulfate* for pregnant patients if seizures persist.

**Clinical Pearls**
- *Midazolam* by the intramuscular (IM) route affords the fastest route to terminating a seizure
**Indications**
- Involuntary, repetitive muscle movements, usually abrupt in onset
- Observed seizure activity
- Decreased mental status
- Unconscious
- Incontinence
- Head injury

**BLS**

**Seizure has stopped**
- Identify and treat injuries
- Check blood glucose and treat per Hypoglycemia protocol
  - If blood glucose is less than 70 mg/dL, administer oral glucose paste (10-15 grams) between the gum and cheek. Administer single additional dose of oral glucose if not improved after 10 minutes.

**Active seizure**
- Do not restrain the patient or place any device into the patient’s mouth.
- Protect the patient from injury.
- Identify and treat potential underlying cause of seizure: epilepsy, head injury, hypoxia, hypoglycemia, hypoperfusion, infection (fever/stiff neck), stroke, alcohol or drug abuse or withdrawal, CVA, head injury

**ALS**

**Active seizure (or seizure lasting more than 10 minutes)**
- Check blood glucose and treat per Hypoglycemia protocol
- Assist patients with the administration of their prescribed benzodiazepine, if available
- Establish IV/IO access. This should not delay midazolam IM/IN for patients with active seizures.
  - **Midazolam:**
    - IM/IN: 0.2 mg/kg IN or IM. Maximum total dose 5 mg. IM route preferred.
    - IV/IO: 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes. Maximum total dose 5 mg.
  - **Diazepam:** if midazolam is not available.
    - Rectal: up to 0.2 mg/kg rectally. Maximum total dose 10 mg
    - SLOW IVP/IO/IM: 0.1 mg/kg in 2.5 mg increments. Maximum total dose 5 mg.
- Pregnant patients: administer midazolam followed by Magnesium sulfate 4 grams IV/IO over 10 minutes (mixed in 50–100 mL of approved diluent).

**MC**

- **Midazolam:** consult for additional doses of midazolam up to a maximum total dose of 5 mg
- **Diazepam:** IM doses of diazepam require consultation, except if suspected nerve agent exposure (CANA and nerve agent antidote kits may be used without consultation)
- **Magnesium sulfate:** consult for additional dose of magnesium sulfate for pregnant patients if seizures persist
**Indications**

- Adult patients (18 years of age and older) with a suspected source of infection **and** present with at least **two** of the following criteria:
  - Temp greater than 100.4°F (38°C) or less than 95.9°F (35.5°C)
  - HR greater than 100 bpm
  - RR greater than 25 (or ETCO2 less than or equal to 32 mmHg)
  - Hypotension (systolic BP less than 90 mmHg)

**BLS**

- Place patient in position of comfort, or supine if hypotension is present.
- Monitor airway and respiratory status, manage as required using the appropriate respiratory distress protocol.
- Rendezvous with ALS or transport to the closest appropriate facility. Use the term “sepsis alert” in consultation with the receiving facility. Patients with suspected sepsis and either altered mental status or hypotension are Priority 1 patients.

**ALS**

- Establish large-bore IV access.
- Obtain a second IV, if it will not delay transport.
- If IV access is unsuccessful and transport time will exceed 20 minutes, obtain IO access for Priority 1 or hypotensive (septic shock) patients.
- Obtain peripheral blood samples if time permits.
- If lungs are clear, and patient does not have a history of CHF or end-stage renal failure, infuse *Lactated Ringer’s* 2 L **IV wide-open.** Reassess every 500 mL for shortness of breath, blood pressure, and SpO₂ saturation changes. Accurately document IV fluid start time and amount of *Lactated Ringer’s* infused.

**OR**

- If patient is fluid-sensitive (i.e., has a history CHF, pulmonary edema, or end-stage renal disease), infuse 250 mL and carefully monitor and reassess. Repeat 250 mL once if no worsening of respiratory status is noted, to a max of 500 mL (consultation may be obtained to provide more fluid).
- Monitor cardiac rhythm and perform 12-lead EKG. Do not delay IV therapy or fluid bolus for these interventions.
- If hypotension persists after 2 L of *Lactated Ringer’s*, provide additional *Lactated Ringer’s* up to a maximum of 30 mL/kg IV, as long as the patient is not volume-sensitive.
- Administer *epinephrine* infusion for patients with persistent hypotension after 30 mL/kg fluid bolus.
  - Add 1 mg of *epinephrine* (either 1 mg/mL or 0.1 mg/mL concentration) in a 100 mL bag of *Lactated Ringer’s* or NS.
  - Use a Microdrip set (60 drops/mL) for infusion administration.
  - Adult *epinephrine* infusion dosage
    - Administer infusion through a free-flowing IV, ideally 20 gauge or larger, or by IO.
    - Start infusion at 1 mL/min (60 drops/min) IV/IO.
    - Check blood pressure every 5 minutes. If MAP is less than 65 mmHg or systolic blood pressure is less than 90 mmHg, increase to a maximum rate of 2 mL/min (120 drops/min).
Clinical Pearls

- Infection can cause a systemic response resulting in fever, altered mental status, shock (including or excluding hypotension), and death. Early recognition and treatment with aggressive fluids, when not contraindicated, and early hospital notification may improve survival rates and patient outcomes.

- The following patient populations are considered especially high-risk for sepsis and should have their temperature measured:
  - Altered mental status
  - Patients in long-term care facilities (nursing home)
  - Indwelling catheters
  - Oncology patients
  - Solid organ transplant
  - Bed-bound
  - Post-operative
  - Currently on antibiotics
  - Asplenic
  - Left ventricular assist device

- Additional fluid beyond 30 mL/kg, or a modification to fluid dose may be approved by online medical consultation.

- If above blood pressure goals are not met upon reaching maximum rate (2 mL/min) of epinephrine infusion, obtain online medical consultation.
**Indications**

- Pediatric patients (less than 18 years of age) with a suspected source of infection and present with at least three of the Pediatric Sepsis Rule-In Criteria by Age.
- Patients who do not meet at least three criteria may be treated using this protocol with approval from a pediatric base station if sepsis is suspected by EMS clinician.

**Pediatric Sepsis Rule-In Criteria by Age**

<table>
<thead>
<tr>
<th>Suspected or known infection plus three criteria</th>
<th>Less than 28 days</th>
<th>1-12 months</th>
<th>1 year but less than 2 years</th>
<th>2-4 years</th>
<th>5-12 years</th>
<th>13-17 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart Rate (sustained)</td>
<td>greater than 205 bpm</td>
<td>greater than 205 bpm</td>
<td>greater than 190 bpm</td>
<td>greater than 140 bpm</td>
<td>greater than 140 bpm</td>
<td>greater than 100 bpm</td>
</tr>
<tr>
<td>Respiratory Rate</td>
<td>greater than 60 rpm</td>
<td>greater than 60 rpm</td>
<td>greater than 40 rpm</td>
<td>greater than 40 rpm</td>
<td>greater than 34 rpm</td>
<td>greater than 25 rpm</td>
</tr>
<tr>
<td>Temp</td>
<td>greater than 38.0 Cº or greater than 100.4 Fº</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cap Refill/Skin</td>
<td>Delayed (greater than 3 seconds), mottled</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic BP (mmHg)</td>
<td>less than 60</td>
<td>less than 70</td>
<td>(less than 70+ (age x2))</td>
<td>(less than 70+ (age x2))</td>
<td>(less than 70+ (age x2))</td>
<td>less than 90</td>
</tr>
<tr>
<td>Mental Status</td>
<td>Unresponsive, confused, inappropriate, lethargic</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High Risk Condition</td>
<td>Cancer, Asplenia, Sickle Cell Disease, bone marrow or solid organ transplant, central or indwelling line/catheter, immunodeficiency or immunosuppression</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Meeting any of these criteria indicates standing order initiation of a fluid bolus.

- Patients who meet the sepsis rule-in criteria and have at least one of the High-Risk Sepsis Rule-In Criteria by Age (shaded) should receive aggressive standing order fluid therapy. Other patients meeting the pediatric sepsis rule-in criteria, but not having one of the high-risk signs, may be treated only after contacting a pediatric base station for medical consultation.

**BLS**

- Place patient in position of comfort, or supine if hypotension is present.
- Monitor airway and respiratory status, manage as required using the appropriate respiratory distress protocol.
- If altered mental status, blood glucose check is required.
- Transport to the closest appropriate facility. Use the term “sepsis alert” in consultation with the receiving facility. Patients with suspected sepsis and either altered mental status or hypotension are Priority 1 patients.
If patient meets the pediatric sepsis rule-in criteria and meets one of the high-risk criteria (shaded), initiate IV/IO access and provide Lactated Ringer’s 20 mL/kg bolus IV/IO over 5–20 min. Maximum single dose of 2 L. Accurately document IV fluid start time and amount of Lactated Ringer’s infused.

Monitor closely for signs of respiratory distress, rales, or delayed capillary refill (greater than 2 seconds). If respiratory status deteriorates rapidly, stop bolus and obtain medical consultation.

**OR**

- For volume-sensitive children, administer initial fluid bolus of 10 mL/kg Lactated Ringer’s IV/IO (max of 250 mL). Volume-sensitive children include: neonates (birth to 28 days), congenital heart diseases, chronic lung disease, or chronic renal failure.

- If patient’s vital signs do not improve after 20 mL/kg fluid, consider additional 20 mL/kg Lactated Ringer’s boluses (up to a max of 60 mL/kg total, including first bolus, in one hour).

- Monitor cardiac rhythm.

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**Pediatric epinephrine infusion dosage**

- The following dosing chart should be used for pediatric patients less than 50 kg who do not achieve an age-appropriate BP (70 + 2 x Age) after 60 mL/kg fluid bolus (using approved epinephrine infusion and 60 drop set):

<table>
<thead>
<tr>
<th>Weight range (kg)</th>
<th>Initial epinephrine dose</th>
<th>If goal blood pressure not achieved at 5 min, increase to</th>
</tr>
</thead>
<tbody>
<tr>
<td>LESS than 10 kg</td>
<td>6 drops/min (0.1 mL/min)</td>
<td>12 drops/min (0.2 mL/min)</td>
</tr>
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<td>10-19 kg</td>
<td>12 drops/min (0.2 mL/min)</td>
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<td>40-49 kg</td>
<td>30 drops/min (0.5 mL/min)</td>
<td>60 drops/min (1.0 mL/min)</td>
</tr>
</tbody>
</table>

- Blood pressure goal:
  - For patients 10 years and older (including adults), systolic blood pressure greater than 90 mmHg;
  - For patients under 10 years of age, systolic blood pressure greater than 70 + 2x age in years mmHg; or
  - Systolic blood pressure ordered by the pediatric base station.
  - If above blood pressure goal not met after 10 minutes, obtain online medical consultation.

- If a pediatric patient meets any of the sepsis rule-in plus one or more shaded areas in the chart, consultation with pediatric base station is required. This should be combined with local base station consultation.

- Additional fluid beyond 60 mL/kg or a modification to fluid dose may be approved by online medical consultation.
Clinical Pearls

- Infection can cause a systemic response resulting in fever, altered mental status, shock (including or excluding hypotension), and death. Early recognition and treatment with aggressive fluids, when not contraindicated, and early hospital notification may improve survival rates and patient outcomes.
- The pediatric septic patient may be difficult to identify due to a poor history or clinicians may have difficulty identifying an obvious source of infection, as many pediatric sepsis patients are very young children or infants.
- The following pediatric patients are at greater risk for sepsis and should have their temperature measured:
  - Altered mental status
  - Asplenia (spleen removed from treatment of trauma or illness)
  - Bone marrow or solid organ transplant
  - Cancer patients
  - Cerebral Palsy
  - Sickle Cell Disease
  - Central or indwelling catheters
  - Immunodeficiency or immunosuppression
  - Bed-bound
  - Severe mental delay
Indications

- Shock is a state of inadequate blood flow to meet the oxygen demands of the cells.
- Patient may exhibit any of the following:
  - Altered mental status; cool, clammy skin; diaphoresis; hypotension, tachycardia or weak pulses; shallow, labored respirations; general weakness; and/or a decreasing pulse pressure.

BLS

- Place the patient in supine position with legs elevated.
- Closely monitor airway and vital signs every 5 minutes
- Pregnancy: For pregnant patients greater than 20 weeks gestation with hypotension, provide constant left lateral uterine displacement.

ALS

- Establish IV/IO access.
- Initiate treatment based on the suspected cause of shock.

- **Cardiogenic Shock**
  - If rales are present, administer small fluid bolus (maximum bolus of 250 mL Lactated Ringer’s). Titrate to a systolic blood pressure of 90 mmHg or greater.
  - Additional fluid requires medical consultation.
  - Initiate epinephrine infusion.
  - Add 1 mg of epinephrine (either 1 mg/mL or 0.1 mg/mL concentration) in a 100 mL bag of Lactated Ringer’s or NS
  - Use a microdrip set (60 drops/mL) for infusion administration.
  - Adult epinephrine infusion dosage:
    - Administer infusion through a free-flowing IV, ideally 20 gauge or larger, or by IO.
    - Start infusion at 1 mL/min (60 drops/min) IV/IO.
    - Check blood pressure every 5 minutes. If MAP is less than 65 mmHg or systolic blood pressure is less than 90 mmHg, increase to a maximum rate of 2 mL/min (120 drops/min).
    - If above blood pressure goals are not met upon reaching maximum rate, obtain online medical consultation.

- **Hemorrhagic Shock**
  - Administer small boluses of Lactated Ringer’s (maximum single bolus of 250 mL prior to additional blood pressure check) to achieve and maintain a systolic blood pressure of 90 mmHg or greater (or mean arterial pressure of 65 mmHg).
  - If head injury is suspected, administer small boluses of Lactated Ringer’s to maintain a systolic blood pressure of 110 mmHg or greater.
  - For patients 15 years of age and older with suspected hemorrhagic shock (SBP less than 90) due to trauma, administer TXA 1 gram in 100 mL of approved diluent (normal saline/Lactated Ringer’s/D5W) IV/IO over 10 minutes. Injury must have occurred within the past one (1) hour. Do not delay transport to initiate TXA.

- **Hypovolemic or Septic Shock**
  - If lungs are clear, administer fluid bolus of 20 mL/kg of Lactated Ringer’s IV. Titrate to a systolic blood pressure of 90 mmHg (or mean arterial pressure of 65 mmHg). Maximum patient dose of 2,000 mL of Lactated Ringer’s.
  - If hypotension persists after 2 L of Lactated Ringer’s are provided, consider additional Lactated Ringer’s up to a maximum of 30 mL/kg total, as long as the patient is not volume-sensitive.
  - Initiate epinephrine infusion if systolic blood pressure remains less than 90 mmHg (or mean arterial pressure less than 65 mmHg) after IV fluid bolus of 30 mL/kg Lactated Ringer’s.
**Anaphylactic shock:** Initiate *epinephrine* infusion for patients who are in extremis with severe hypotension or impending respiratory failure, after having administered 3 doses of IM *epinephrine*. (Refer to *Anaphylaxis* protocol.)

**Neurogenic shock** (suspected spinal cord injury which typically presents with hypotension and bradycardia)

- If lungs are clear, administer fluid bolus of 20 mL/kg of *Lactated Ringer’s IV*. Titrate to a systolic blood pressure of 110 mmHg (or mean arterial pressure of 85 mmHg). Maximum patient dose of 2,000 mL of *Lactated Ringer’s*.
- Initiate *epinephrine* infusion if systolic blood pressure remains less than 110 mmHg (or mean arterial pressure less than 85 mmHg).

**ALS**

**MC**

- Not applicable.
**Indications**

- Shock is a state of inadequate blood flow to meet the oxygen demands of the cells.
- Patient may exhibit any of the following:
  - Altered mental status; cool, clammy skin; diaphoresis; tachycardia or weak pulses; shallow, labored respirations; delayed capillary refill greater than 2 seconds, pallor or peripheral cyanosis; general weakness; and/or a decreasing pulse pressure.
  - Hypotension
    - Neonates (birth to 28 days): a systolic blood pressure less than 60 mmHg
    - Infants (less than 1 year of age): a systolic blood pressure less than 70 mmHg
    - Children (1 to 9 years of age): a systolic blood pressure of less than \[70 + (2 \times \text{years})\]
    - Children (10 to 17 years of age): systolic blood pressure of less than 90 mmHg

**BLS**

- Place the patient in supine position with legs elevated.
- Closely monitor airway and vital signs every 5 minutes
- Pregnancy: For pregnant patients greater than 20 weeks gestation with hypotension, provide constant left lateral uterine displacement.

**ALS**

- Establish IV/IO access.
- If age-related vital signs and patient’s condition indicate hypoperfusion, administer initial fluid bolus of Lactated Ringer’s 20 mL/kg IV/IO.
  - If patient's condition does not improve, administer a second bolus of Lactated Ringer’s 20 mL/kg IV/IO.
  - **OR**
  - For volume-sensitive children administer initial fluid bolus of Lactated Ringer’s 10 mL/kg IV/IO. If patient’s condition does not improve, administer a second bolus of Lactated Ringer’s 10 mL/kg Lactated Ringer’s IV/IO.
    - Volume-sensitive children include: neonates (birth to 28 days), children with congenital heart disease, chronic lung disease, or chronic renal failure.
  - **For patients 15 years of age and older** with suspected hemorrhagic shock (SBP less than 90) due to trauma, administer TXA 1 gram in 100 mL of approved diluent (normal saline/Lactated Ringer’s/D5W) IV/IO over 10 minutes. Injury must have occurred within the past one (1) hour. Do not delay transport to initiate TXA.
Third and subsequent fluid boluses, *Lactated Ringer’s 20 mL/kg IV/IO* require medical consultation.

**Pediatric epinephrine** infusion dosage

- The following dosing chart should be used for pediatric patients less than 50 kg, using approved *epinephrine* infusion and 60-drop set:

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</tr>
</tbody>
</table>

**Blood pressure goal:**

- For patients 10 to 17 years of age, systolic blood pressure greater than 90 mmHg;
- For patients less than 10 years of age, systolic blood pressure greater than 70 + 2x age in years mmHg; OR
- Systolic blood pressure ordered by the pediatric base station.
- If above blood pressure goal not met after 10 minutes, obtain online medical consultation.
Indications
- Blurred vision (including intermittent loss of vision in one or both eyes, which may have resolved upon arrival of EMS)
- Difficulty speaking
- Numbness or weakness (often one side only)
- Sudden onset of dizziness or loss of balance
- Severe, unexplained headache

**Position patient with head elevated at 30 degrees.**
**Check blood glucose level; if less than 70 mg/dL, treat per Hypoglycemia protocol.**
**Pediatrics: for patients who have not reached their 18th birthday, administer oxygen 2-6 lpm via nasal cannula, unless the patient is hypoxic or in respiratory distress.**
**Perform Cincinnati Prehospital Stroke Scale (any abnormality is positive for stroke):**
  - **Facial droop** – have patient smile or show teeth
    - Normal – both sides of face move equally
    - Abnormal – one side of the face does not move as well as the other side
  - **Arm drift** – patient closes eyes and holds both arms straight out for 10 seconds
    - Normal – both arms move the same or both arms do not move at all
    - Abnormal – one arm does not move or one arm drifts down compared with other
  - **Speech abnormal** – have the patient say “you can’t teach an old dog new tricks”
    - Normal – patient uses correct words with no slurring
    - Abnormal – patient slurs words, uses wrong words, or is unable to speak

**Perform Posterior Cerebellar Assessment (any abnormality is positive for stroke):**
  - **Balance:** patient complains of sudden onset of loss of balance or dizziness
  - **Eyes:** patient has sudden vision loss (including intermittent loss of or blurred vision)

If either the Cincinnati Prehospital Stroke Scale or Posterior Cerebellar Assessment is positive, then calculate the suspected stroke patient’s Los Angeles Motor Scale (LAMS) score:

- **Facial droop**
  - Absent 0
  - Present 1

- **Arm drift**
  - Absent 0
  - Drifts down 1
  - Falls rapidly 2

- **Grip strength**
  - Normal 0
  - Weak grip 1
  - No grip 2

**Obtain and document a telephone number for one or more individuals who have knowledge of the patient’s presenting symptoms, last known well time, and medical history. Communicate this information to receiving hospital staff.**
• Destination determination for a suspected stroke patient who can be delivered to the appropriate stroke center within 22 hours from when patient was last known well:
  ▪ **LAMS score 0-3**: transport the patient to the closest Designated Acute Stroke Ready, Primary, or Comprehensive Stroke Center.
  ▪ **LAMS score of 4 or greater**: transport the patient to the closest Comprehensive Stroke Center or thrombectomy-capable Primary Stroke Center. If the patient cannot be delivered to an appropriate center within 30 minutes, go to the closest Designated Acute Stroke Ready or Primary Stroke Center.
  ▪ **For suspected stroke patients greater than 30 minutes from any stroke center**: transport patient to the closest hospital or request aviation if there would be a time savings.
  ▪ **For pediatric suspected stroke patients (have not reached their 18th birthday)**: consult with a local base station and pediatric base station to arrange transport to a Pediatric Trauma Center.

• Establish IV access, preferably on the unaffected side of the body
• Obtain blood sample using a closed system
• If blood glucose is less than 70 mg/dL, treat per **Hypoglycemia** protocol

• For all suspected stroke patients within 22 hours of last known well time, notify the receiving stroke center or hospital as soon as possible. During the consultation, the clinician shall use the verbiage, “Priority 1, Stroke Alert patient with a last known well time of XX:XX” as the universal method of notifying the facility that the patient meets the stroke inclusion criteria.
• If the patient is hypotensive, obtain medical consultation.
• Do not treat hypertension in the field.

**Clinical Pearls**
• While strokes during pregnancy or shortly after giving birth are rare, there has been a significant rise reported in the literature. Mothers-to-be and postpartum mothers have an increased risk.
• Strokes are less common in children than in adult patients. However, children with the following conditions have a higher risk of stroke: congenital heart defects, brain injury, sickle cell disease (blood disorders), and certain types of infections (meningitis, encephalitis).
Support ABCs and provide any needed BLS/ALS Interventions Check Glucose

Determine presence of stroke severity using Posterior Cerebellar assessment and Cincinnati Prehospital Stroke Scale

Either test positive for stroke

Treat and transport per pt presentation

YES

Determine time patient last known well
LAMS Assessment

Signs and symptoms consistent with stroke AND onset less than 22 hrs.

NO

Transport to closest Stroke Center (a) (b)

YES

LAMS 4 or greater?

TRANSPORT TO CLOSEST STROKE CENTER AS PRIORITY 1 AND STROKE ALERT

NO

(a) - Designated Acute Stroke Ready, Primary, or Comprehensive Stroke Center
(b) - Patients under 18 years of age should be transported to a Pediatric Trauma Center

Transport to closest Comprehensive Stroke Center or thrombectomy-capable Primary Stroke Center if within 30 minutes as Priority 1 and Stroke Alert

NO

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Indications

- Transient loss of consciousness with an inability to maintain postural tone
- Symptoms may resolve without intervention or prior to EMS arrival.
- Patients who “feel like they are going to pass out” (near-syncope)
- For children less than 24 months of age, refer to ALTE/BRUE protocol.

BLS

- Place patient in the supine position, with feet elevated.
- Check blood glucose. If less than 70 mg/dL, refer to Hypoglycemia protocol.
- Perform Cincinnati Stroke Scale. If any abnormal findings are present, refer to Stroke protocol.
- If the patient sustained a fall, assess for trauma and treat injuries per trauma protocols.

ALS

- Monitor cardiac rhythm and treat according to appropriate algorithm.
- Obtain 12-lead EKG for patients 13 years of age and older.
- Establish IV access.
- Administer LR as appropriate to maintain a systolic blood pressure of at least 90 mmHg (or MAP of 65 mmHg) for adults.
  - For patients under the age of 18, administer 20 mL/kg LR bolus titrated to age-appropriate blood pressure.

MC

- Not applicable.

Clinical Pearls

- History, physical examination, and 12-lead EKG should all be used to determine the patient’s risk of an adverse outcome. Patients with history or evidence of heart failure, structural cardiac anomaly, and/or abnormal finding on EKG are at higher risk for adverse outcomes.
- Syncope in children may be associated with serious medical conditions such as heart failure or structural cardiac anomalies.
1. Indications
Patient presents pregnant, with contractions and/or pain, accompanied by bleeding or discharge, crowning during contraction, the feeling of an impending bowel movement, and/or a rock-hard abdomen.

2. Treatment
- Pre-Arrival Information
  - Excessive Bleeding? YES → Absorb Bleeding Treat for Shock
  - NO → Seizures YES → Transport Left Lateral Position Maintain Body Temp. Have Suction Ready (d)
  - NO → Baby’s Head Presents?
    - YES → Hand/Foot/ Butt Presents?
      - YES → Left Lateral Position Deliver Body Support Baby’s Wt. Form V to Open Airway
      - NO → Cord Presents?
        - YES → Position Mother Face Down & Butt Up Wrap Cord Keep Moist Insert Gloved Hand to Lift Baby (a,b)
        - NO → Amniotic Sac Broken? NO → Puncture Sac
          - YES → Suction mouth then nose only if non-vigorous or obvious airway obstruction
    - NO → Support Head

(Continued on next page)
5.1 BLS

Shoulders Delivered? 

- NO 
  - Assist Upper then Lower Shoulder 
  - NO 

- YES 
  - Deliver Remainder of Baby 
    - YES 
      - Breathing? 
        - NO 
          - Clamp and Cut Cord 
          - Deliber Placenta 
            - NO 
              - Multiple Birth? 
                - YES 
                  - GO TO NEWLY BORN PROTOCOL 
                - NO 
                  - Excessive Bleeding? 
                    - YES 
                      - Consider Uterine Massage (c) 
                      - NO 
                        - Transport to Medical Facility 

- GO TO BEGINNING 

Reconsider ALS, Priority Transport & Notify Receiving Facility

(a) - Keep presenting part of baby off the cord. Monitor and attempt to maintain the pulse in the cord.
(b) - Position of mother: 📆
(c) - Uterine massage is performed with the heel of the hand applying firm pressure from the pubis toward the umbilicus only. This massage is continued until bleeding diminishes. Transport rapidly.
(d) - Go to Seizure protocol: Consider midazolam.
1. Indications
This protocol applies to the infant within the first hour after delivery.

**UNIVERSAL ALGORITHM FOR THE NEWLY BORN FOR BLS**

- **Dry, Warm, Position, Stimulate**
- Suction if non-vigorous or obvious airway obstruction

**If Apnea/Gasping, HR is less than 100 or central cyanosis**
Ventilate with BVM @ 40–60 breaths/min using room air for the first minute (40-60 breaths) before connecting to 100% oxygen

**HR less than 60 after 30 seconds of BVM**
120 compressions/minute with 3:1 compressions: ventilations

**AED NOT INDICATED FOR NEWLY BORN**

- ALS Care for Rhythm Management & Treatment Medications (ALS Only)
**Universal Algorithm for the Newly Born for ALS**

**Dry, Warm, Position, Stimulate**

**Assess respirations**

- **Respirations Spontaneous with Good Effort**
- **Respiratory Rate Slow/Gasping, Absent**
  - Position airway
  - Ventilate with BVM @ 40-60 breaths/min using room air for first minute (a)

**Evaluate Heart Rate**

- **Heart Rate less than 60**
  - Perform CPR
  - 120 compressions/minute with 3:1 compressions: ventilations on 100% oxygen
  - Consider intubation (a)

- **Heart Rate 60–100**
  - Support ventilations with BVM at a rate of 40-60 breaths/min. Use room air for an additional 30 seconds before connecting to 100% oxygen

- **Heart Rate greater than 100**
  - Reassess respiratory rate and effort
  - Remain on room air Monitor $\text{SpO}_2$ (a)
  - Evaluate skin color
  - APGAR at 1 min, repeat at 5 mins

**Reassess**

- IV/IO with LR (c)
  - Epinephrine IV/IO 0.01 mg/kg (0.1 mg/mL)
  - Neonates (0–28 days), Epinephrine ET 0.03 mg/kg (0.1 mg/mL) dilute with 1 mL
  - Repeat every 4 minutes
  - Consider intubation (a)
  - Consider causes (b)

- Administer supplemental $\text{O}_2$
  - Monitor IV/IO with LR if poor perfusion (c)

- Monitor and maintain body temperature
  - Transport

**Medical consult**

**Monitor and maintain body temperature**

**Transport**

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**Release Date July 1, 2022**

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**APGAR Chart**

(a) - Acceptable Target SpO₂ after Birth
   - 1 min – 60-65%
   - 2 min – 65-70%
   - 3 min – 70-75%
   - 4 min – 75-80%
   - 5 min – 80-85%
   - 10 min – 85-95%

(b) - Consider possible causes of depressed newborn.
   (Parenthesis = possible therapies and treatments)
   Respiratory depression (Premature infants less than 32 weeks gestation will likely require ongoing BVM ventilations due to immature lungs.)
   Hypoglycemia (Threshold for treatment = 30 mg/dL) (D10W 2–4 mL/kg IV/IO (D10W is prepared by mixing one part of D50W with four parts LR.))
   Hypothermia (Warming)
   Hypovolemia (Volume infusion – see “c”, below)

(c) - Volume infusion is 10 mL/kg.

### APGAR Chart

<table>
<thead>
<tr>
<th>SIGN</th>
<th>0</th>
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<th>2</th>
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<tr>
<td>MUSCLE TONE (ACTIVITY)</td>
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<td>ACTIVE, GOOD FLEXION</td>
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<tr>
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<td>LESS THAN 100/MIN</td>
<td>GREATER THAN 100/MIN</td>
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<tr>
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<td>NO RESPONSE</td>
<td>SOME GRIMACE OR AVOIDANCE</td>
<td>COUGH, CRY OR SNEEZE</td>
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<tr>
<td>COLOR (APPEARANCE)</td>
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<td>PINK BODY, BLUE HANDS/FEET</td>
<td>PINK</td>
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<tr>
<td>RESPIRATIONS</td>
<td>ABSENT</td>
<td>SLOW/IRREGULAR, INEFFECTIVE</td>
<td>CRYING, RHYTHMIC EFFECTIVE</td>
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</table>

*Nasal or Oral Suction Catheter Stimulus*
Indications
- Heavy vaginal bleeding as a result of pregnancy, miscarriage, postpartum bleeding, or sexual assault
- Patient may exhibit signs or symptoms of shock

BLS
- Place absorbent pads under the patient.
- If signs of hypotension, place the patient in the supine position with the patient’s feet elevated at a 30-degree angle.
- If postpartum bleeding, perform uterine massage from the pubis toward the umbilicus only.
- Request ALS for patients with signs or symptoms of shock.

ALS
- Establish IV/IO access, as appropriate.
- Administer LR fluid bolus 20 mL/kg IV; titrate to systolic blood pressure of 90 mmHg and refer to the Shock: Hypoperfusion protocol.

MC
- Fluid administration beyond 2,000 mL requires medical consultation.

Clinical Pearls
- Bring all products of conception to the hospital.
- Do not pull products of conception from vaginal opening without medical consultation.
**Indications**
- Shortness of breath with wheezing or decreased air entry, presumed to be due to bronchospasm from reactive airway disease, asthma, or COPD
- Signs of respiratory distress which may include:
  - Accessory muscle use and/or tripod positioning
  - Cyanosis, mottled skin
  - Nasal flaring, retractions

**BLS**
- **Albuterol (BLS) inhaler (2 puffs inhaled)** or **albuterol (BLS) 2.5 mg nebulized.** May repeat dose one time, as needed, within 30 minutes.
- For severe respiratory symptoms, administer **epinephrine auto-injector (BLS) 0.3 mg IM** or administer **epinephrine (BLS) (1 mg/mL) 0.5 mg in the lateral thigh.**
- Medical consult required if patient has cardiac history or is pregnant.

**ALS**
- **Albuterol 2.5 mg nebulized** and **ipratropium (Atrovent®) 500 mcg nebulized;** may repeat **albuterol nebulized 2.5 mg one time.**
- For severe respiratory symptoms, administer **epinephrine auto-injector 0.3 mg IM via or epinephrine (1 mg/mL) 0.5 mg IM.** May repeat every 5 minutes for a total of 3 doses, as needed.
- Establish IV/IO access for patients with moderate to severe symptoms.
- For moderate to severe exacerbations, administer **dexamethasone 10 mg IV/PO.**
- Consider **terbutaline 0.25 mg IM** for moderate exacerbations or severe exacerbations with a cardiac history. May repeat dose one time after 15 minutes if there is not improvement. Maximum total dose 0.5 mg IM.
- For severe exacerbations or patients whose condition deteriorates despite treatments above, administer nebulized treatments, along with high-flow oxygen, continuous positive airway pressure (CPAP), or bag-valve mask (BVM).

**MC**
- (BLS) For patients with a cardiac history or pregnancy, obtain medical consultation for **epinephrine (1 mg/mL) or epinephrine auto-injector.**
- (ALS/BLS) Additional doses of **albuterol, ipratropium, epinephrine (1 mg/mL) or epinephrine auto-injector** beyond those listed above require medical consultation.
- (ALS) For moderate to severe exacerbations, administer **magnesium sulfate 1–2 grams,** mixed in 50–100 mL of approved diluent, IV/IO over 10–20 minutes.

**Clinical Pearls**
- If respiratory distress is due to a suspected allergic reaction or anaphylaxis, refer to Allergic Reaction and Anaphylaxis protocols.
Indications

- Shortness of breath with wheezing or decreased air entry, presumed to be due to bronchospasm from reactive airway disease or asthma
- Signs of respiratory distress which may include:
  - Accessory muscle use and/or tripod positioning
  - Cyanosis, mottled skin
  - Nasal flaring, retractions

**BLS**

- **Albuterol (BLS) inhailer (2 puffs inhaled) or albuterol (BLS) nebulizer.** May repeat dose one time, as needed, within 30 minutes.
  - For infants and children less than 2 years of age, administer nebulized albuterol (BLS) 1.25 mg.
  - For patients 2 years of age or greater, administer nebulized albuterol (BLS) 2.5 mg.
- **For severe respiratory symptoms:**
  - Less than 5 years of age: pediatric epinephrine auto-injector (BLS) 0.15 mg IM or epinephrine (BLS) (1 mg/mL) 0.15 mg IM
  - 5 years of age or greater: epinephrine auto-injector (BLS) 0.3 mg IM or epinephrine (BLS) (1 mg/mL) 0.5 mg IM

**ALS**

- **Albuterol and Ipratropium (Atrovent®) nebulized.**
  - For an infant less than 1 year of age: albuterol 1.25 mg nebulized; ipratropium is contraindicated.
  - For a child 1 year of age or greater, but less than 2 years of age: albuterol 1.25 mg and ipratropium 250 mcg nebulized.
  - For a patient 2 years of age or greater: albuterol 2.5 mg and ipratropium 500 mcg nebulized.
- **May repeat albuterol one time for recurrent or worsening symptoms**
- **Epinephrine (1 mg/mL)**
  - Less than 5 years of age: epinephrine (1 mg/mL) 0.15 mg IM
  - 5 years of age or greater: epinephrine (1 mg/mL) 0.5 mg IM
- **May repeat epinephrine (1 mg/mL) every 5 minutes for a total of 3 doses, as needed.**
- For moderate to severe exacerbations, consider the administration of dexamethasone 0.5 mg/kg PO/IV, up to a maximum dose of 10 mg.
- For patients 12 years of age and older, consider terbutaline 0.25 mg IM for moderate to severe exacerbations. May repeat dose one time after 15 minutes if no improvement. Maximum total dose 0.5 mg IM.

**MC**

- (BLS) For patients with congenital heart or lung disease, obtain medical consultation for epinephrine (1 mg/mL), pediatric epinephrine auto-injector, or epinephrine auto-injector.
- (ALS/BLS) Additional doses of albuterol, ipratropium, epinephrine (1 mg/mL) or epinephrine auto-injector beyond those listed above require medical consultation.
- (ALS) For moderate to severe exacerbations, administer magnesium sulfate 50 mg/kg IV/IO to a max of 2 grams given over 10–20 minutes (mixed in 50 - 100 mL of approved diluent). For children, administer 20 ml/kg fluid bolus of LR with magnesium to reduce risk of hypotension.

Clinical Pearls

- If respiratory distress is due to a suspected allergic reaction or anaphylaxis, refer to Allergic Reaction and Anaphylaxis protocols.
Indications
- Exposure to chlorine or phosgene with respiratory symptoms
- Signs of respiratory distress may include:
  - Wheezing and/or crackles, abnormal respiratory rate, dyspnea, diminished or absent breathing sounds, and/or tripod positioning.
  - Cyanosis or mottled skin
  - Altered mental status
  - Rapid heart rate

BLS
- For patients with dyspnea with wheezing or symptoms related to bronchospasm, treat with albuterol per Asthma/COPD protocol.
- For patients with dyspnea and rales or suspected pulmonary edema, treat supportively with oxygen, as indicated per CHF protocol.

ALS
- For patients with dyspnea with wheezing or symptoms related to bronchospasm, treat with bronchodilators and dexamethasone per Asthma/COPD protocol.
- For patients with dyspnea and rales or suspected pulmonary edema, treat supportively with oxygen and CPAP, as indicated per CHF protocol.

MC
- As noted in Asthma/COPD and CHF protocols.

Clinical Pearls
- Ensure BSI and decontamination of the patient with suspected chlorine or phosgene exposure.
- Symptoms of chlorine or phosgene exposure may be delayed for up to 48 hours. Patients can rapidly deteriorate once symptoms present.
Indications

- Shortness of breath (dyspnea) due to suspected pulmonary edema, which may present with some of the following associated signs or symptoms:
  - Pulmonary: rales, wheezing, hypoxia, tachypnea
  - Systemic: peripheral edema, hypertension, jugular venous distention (JVD), orthopnea, paroxysmal nocturnal dyspnea
  - History of CHF or cardiac disease (MI)
- **Mild:** Mild dyspnea at rest, despite oxygen administration and able to speak in full sentences, or dyspnea with minimal exertion
- **Moderate:** Moderate dyspnea, pulse oximetry less than 94% on supplemental oxygen. SBP usually greater than 150 mmHg. Unable to speak in full sentences with a normal mental status.
- **Severe:** Severe dyspnea, respiratory failure, hypoxia with pulse oximetry less than 90% on supplemental oxygen, diaphoresis, SBP usually greater than 180 mmHg. Speaking one word at a time, altered level of consciousness.

**BLS**
- Place patient in High Fowler's position.
- Assess the severity of patient’s dyspnea on a scale where 0 is “no trouble breathing” and 10 is “the worst trouble breathing.”
- For patients with pulse oximetry less than 94%, administer oxygen per GPC.

**ALS**
- Monitor cardiac rhythm and treat according to appropriate algorithm.
- Perform 12-lead EKG.
- Establish IV access.
- **Mild symptoms:**
  - For patients with pulse oximetry less than 94%, administer oxygen per GPC.
  - Administer low-dose nitroglycerin 0.4 mg SL every 3-5 minutes, to a maximum dose to a maximum dose of 1.2 mg.
- **Moderate and severe symptoms:**
  - Initiate CPAP. Until CPAP is applied, administer high-dose nitroglycerin as noted below.
  - Assess BP before each nitroglycerin dose.
  - For patients with SBP greater than 150 mmHg, administer high-dose nitroglycerin every 3-5 minutes until CPAP is applied or if CPAP is not tolerated:
    - Administer nitroglycerin 0.4 mg SL and apply 1 inch of nitroglycerin paste.
    - Administer nitroglycerin 0.8 mg SL.
    - Continue nitroglycerin 0.8 mg SL, every 3-5 minutes, to achieve a 20% reduction in SBP.
  - An IV is not required to start nitroglycerin for patients with moderate and severe symptoms, but should be established after acute interventions to stabilize respiratory status.
  - If SBP drops below 90 mmHg, treat with Lactated Ringer’s fluid bolus of 250 mL; may repeat once, if the patient remains hypotensive.
- **Hypotensive CHF patients:**
  - Administer a small Lactated Ringer’s fluid bolus of 250 mL.
  - If the patient remains hypotensive, administer epinephrine infusion (1 mg epinephrine in 100 mL LR), starting at 1 mL/min (60 drops/min). Titrate to systolic BP of 90 mmHg or MAP of 65 mmHg. Maximum rate of 2 mL/min (120 drops/min). IV infusion pump preferred.
Clinical Pearls

- Geriatric patients demonstrating marked hypertension in association with shortness of breath/respiratory distress and wheezing (in the absence of asthma or infection) strongly suggests congestive heart failure/pulmonary edema.
- For CHF patients with hypertension, the goals of treatment are to reduce the pressure of blood returning to the heart (preload) and the resistance that the left ventricle must pump against (afterload). The most effective and safe medication for these goals is nitroglycerin.
- CPAP is the preferred therapy for moderate to severe CHF patients with hypertension. Do not remove CPAP to continue administering NTG.
- Some CHF patients will present with hypotension and pulmonary edema. These patients may benefit from small fluid boluses (250 mL) and epinephrine infusion, if SBP remains below 90 mmHg or MAP below 65 mmHg. Nitroglycerin is contraindicated in these hypotensive patients.

- If an inferior wall MI is present on EKG, consult prior to administering nitroglycerin.
- Additional Lactated Ringer’s fluid boluses beyond 250 mL for CHF patients require medical consultation.
- If a hypotensive CHF patient continues to have SBP less than 90 mmHg (MAP less than 65 mmHg), despite maximum epinephrine infusion dose, obtain medical consultation.
### Indications
- Shortness of breath (dyspnea) with any of the following associated signs or symptoms:
  - Rales
  - Peripheral edema
  - Jugular venous distention (JVD)
  - History of congenital heart or chronic lung disease

### BLS
- Place patient in semi-Fowler’s position.
- For patients with pulse oximetry less than 94%, administer oxygen per GPC.
- Transport to the pediatric specialty center that follows the patient, if clinical condition permits.

### ALS
- Monitor cardiac rhythm and treat according to appropriate algorithm.
- Establish IV/IO access.
- Patients with moderate to severe respiratory distress may require high flow oxygen via non-rebreather mask, CPAP, or BVM while receiving medication via nebulizer.

### MC
- Medical consultation is required for patients with congenital heart or chronic lung disease.
- (ALS/BLS) Consider *albuterol* nebulized.
  - For children less than 2 years, *albuterol* 1.25 mg nebulized.
  - For children greater than or equal to 2 years, *albuterol* 2.5 mg nebulized.
- (ALS) Consider *morphine* 0.1 mg/kg SLOW IVP/IO/IM (1–2 mg/min). Maximum dose 5 mg.
- (ALS) If hypotensive, consider pediatric *epinephrine* infusion.
Indications

- **Mild**: Barky cough without stridor at rest (Priority 2)
- **Moderate**: Barky cough with stridor at rest, without agitation; may exhibit mild respiratory distress (Priority 2)
- **Severe**: Stridor at rest, signs of severe respiratory distress that is associated with agitation or decreased level of consciousness (Priority 1)

**BLS**

- Ensure that the patient has a patent airway and adequate respiratory effort.
- Assess respiratory status, looking specifically for signs and/or symptoms of respiratory distress (nasal flaring, retractions, increased/decreased respirations, skin color, change in level of consciousness).

**ALS**

- Monitor cardiac rhythm.
- **Mild**: Administer dexamethasone 0.5 mg/kg PO up to a maximum dose of 10 mg.
- **Moderate**: Administer dexamethasone 0.5 mg/kg PO up to a maximum dose of 10 mg.
  If no change in patient’s condition, then administer 2.5 mL of epinephrine (1 mg/mL) nebulized.
- **Severe**: for severe respiratory distress and imminent respiratory arrest:
  - Administer 0.01 mg/kg of epinephrine (1 mg/mL) IM. Max single dose of 0.5 mg IM.
  - Administer dexamethasone 0.5 mg/kg IV up to a maximum dose of 10 mg. If IV is not available, administer IM.
  - Administer 2.5 mL of epinephrine (1 mg/mL) nebulized.

**MC**

- If epiglottitis is suspected (i.e., drooling with above signs and symptoms), obtain medical consultation prior to initiating this protocol.
- Obtain medical consultation for:
  - Patients less than 1 year of age
  - Patients requiring additional nebulized epinephrine due to level of respiratory distress
  - Additional interventions needed to stabilize the patient

**Clinical Pearls**

- All patients who receive nebulized epinephrine must be transported by an advanced life support unit to the closest appropriate medical facility.
Pediatric Respiratory Distress for BLS – Universal Algorithm

Assess Responsiveness

Not Responsive
Assess ABCs

Responsive
Assess Breathing

Go to Universal Algorithm for Pediatric Emergency Cardiac Care for BLS

If respiratory with adequate rate and effort (b):
Oxygen 90–100% via non-rebreather mask

If respiratory with inadequate rate and effort: (a)
BVM with 100% oxygen at 12–20 breaths/min

Suspected Cause

Acute onset of upper airway symptoms:
Stridor, head bobbing, drooling
Assess/treat for foreign body obstruction
See GPC D. 4. Airway See Croup Protocol

History of life-threatening allergic reaction or severe symptoms
See Allergic Reaction or Anaphylaxis Protocol, as appropriate

History of asthma/chronic lung disease
See Asthma/COPD Protocol

History of congenital or acquired heart disease
See Pulmonary Edema/Congestive Heart Failure Protocol

Transport to nearest appropriate medical facility

Consider ALS Rendezvous

(a) Inadequate RR: Infant less than 20 breaths per minute, Child less than 16 breaths per minute, Adolescents less than 12 breaths per minute. Inadequate effort: Poor chest rise, shallow respirations/poor air movement, cyanosis, severe retractions, paradoxical breathing.

(b) For children with chronic lung disease or congenital heart disease: Maintain or increase home oxygen to maintain patient’s target saturations.
Pediatric Respiratory Distress for ALS – Universal Algorithm

Assess Responsiveness

Not Responsive
Assess ABCs

Responsive
Assess Breathing

Suspected Cause

If respiratory with adequate rate and effort (b):
Oxygen 90–100% via non-rebreather mask

If respiratory with inadequate rate and effort: (a)
BVM with 100% oxygen at 12–20 breaths/min

Go to Universal Algorithm for Pediatric Emergency Cardiac Care for ALS

Acute onset of upper airway symptoms:
Stridor, head bobbing, drooling
Assess/treat for foreign body obstruction:
See GPC D. 4. Airway See Croup Protocol

History of life-threatening allergic reaction or severe symptoms
See Allergic Reaction or Anaphylaxis Protocol, as appropriate

History of asthma/chronic lung disease or acute onset of lower airway symptoms:
Wheezeing, retractions, nasal flaring
See Asthma/COPD Protocol

History of congenital or acquired heart disease or acute onset of heart failure:
Wheezing/crackles, edema, poor perfusion
See Pulmonary Edema/Congestive Heart Failure Protocol

Transport to nearest appropriate medical facility

(a) Inadequate RR: Infant less than 20 breaths per minute, Child less than 16 breaths per minute, Adolescent less than 12 breaths per minute. Inadequate effort: Poor chest rise, shallow respirations/poor air movement, cyanosis, severe retractions, paradoxical breathing.

(b) For children with chronic lung disease or congenital heart disease: Maintain or increase home oxygen to maintain patient’s target saturations.
**Indications**

- Suspected exposure to carbon monoxide (CO), e.g., house fires, malfunctioning furnaces, suicide attempts
- Symptoms are variable, but may include: headache, dizziness, and nausea and vomiting, most frequently. Symptoms can also include: chest pain, altered mental status, dyspnea, and/or seizures.

**Remove patient from toxic environment by appropriately trained personnel using proper level PPE.**

**Decontaminate as appropriate.**

**Administer high-flow oxygen.**

**Treat respiratory and/or cardiac symptoms per appropriate protocol.**

**Consider transport to the Hyperbaric Medicine Specialty Center, if indications apply:**

- Patients with exposure to products of combustion (smoke) or carbon monoxide who have a carboxyhemoglobin value of greater than 25%, with or without symptoms OR
- Patients with proven exposure to products of combustion (smoke) or carbon monoxide who have:
  - **Any** of the following diagnostic indicators:
    - Carboxyhemoglobin value (co-oximeter or blood) of greater than 15%
    - Alarm of EMS or fire agency-maintained passive carbon monoxide monitor
    - Targeted atmospheric carbon monoxide value 100 ppm or greater in the patient environment
  - **And** one or more of the following:
    - History of loss of consciousness during exposure (may have since resolved)
    - GCS persistently less than or equal to 13
    - Rapid decline of neurological symptoms, including actively seizing patients with appropriate airway stabilization
    - Pregnancy
    - Chest pain
    - Extremes of age
    - Per clinician discretion

**Contraindications** for transport to Hyperbaric Medicine Specialty Center:

- Transport time to Hyperbaric Medicine Specialty Center is greater than 1 hour
- Patients in cardiac arrest or ROSC post-cardiac arrest
- Patients with burns and trauma should be transported to the nearest appropriate trauma center.

**Obtain a blood sample using closed system, particularly if carboxyhemoglobin co-oximeter measurement is not available.**

**Establish IV access.**

**If shock is present, administer 20 mL/kg bolus of Lactated Ringer’s and treat per Shock/ Hypoperfusion protocol.**

**For smoke inhalation patients, refer to OSP: Cyanide protocol, if jurisdiction is participating.**
Overdose/Poisoning: Carbon Monoxide/Smoke Inhalation (continued)

Clinical Pearls
- Pulse oximetry may not be accurate for carbon monoxide victims. Patients may have normal SpO\textsubscript{2} levels with carbon monoxide toxicity.
- **Pregnant patients and infants:** Fetal hemoglobin has a very high affinity for carbon monoxide. A pregnant mother may be asymptomatic, yet fetal levels may be dangerously high.
- Patients who do not meet Hyperbaric Medicine Specialty Center referral criteria should be transported to the nearest hospital-based emergency department.
# Cold-Related Emergencies

**Indications**
- **Frostbite**: reddened, mottled, bluish or white-gray appearing skin in a cold-exposed area; pain in the area followed by numbness in later stages
- **Hypothermia**: exposure to cold environment, with symptoms including: shivering, cool, mottled or pale skin, stiffness of the muscles, altered gait, shallow respirations, altered level of consciousness, weak pulses

## BLS
- Remove patient from cold environment and remove any wet clothing.
- Protect from further heat loss.
- For frostbite:
  - Handle potential frostbitten areas gently; do not rub affected areas.
  - Cover affected areas with lightweight gauze.
- For hypothermia:
  - Passively rewarm the patient within a warm environment.
  - Use a thermal blanket to cover the patient and a separate blanket to cover the patient’s head, except for the face.
  - Administer warmed oxygen, if available.
  - Handle all hypothermic patients gently; rough handling may precipitate cardiac arrest.
- For hypothermic patients in cardiac arrest:
  - Attempt resuscitation on all hypothermic cardiac arrest patients, unless there is an injury incompatible with life.
  - Administer AED shocks, if indicated.
  - Transport hypothermic patients in cardiac arrest to the closest appropriate facility, unless there is an injury incompatible with life.

## ALS
- Establish IV/IO access, if appropriate.
- Provide analgesia per *Pain Management* protocol.
- For patients with hypothermia and hypothermic cardiac arrest:
  - Monitor cardiac rhythm and treat according to the *Cardiac Arrest* protocol.

## MC
- Obtain early medical consultation for hypothermic cardiac arrest patients. Drug and defibrillation intervals may change depending on degree of hypothermia.

## Clinical Pearls
- For patients with severe hypothermia, pulse and respiratory rates may be very slow. Initiate CPR if there is any doubt as to whether the patient is in cardiac arrest.
- Hypothermic patients in cardiac arrest may survive after a prolonged downtime. Hypothermia is a contraindication to the *Termination of Resuscitation* protocol.
Environmental Emergencies: Depressurization/Decompression

Indications
- History of SCUBA, breathing in a pressurized environment (flight) or altitude chamber with sudden depressurization
- Confusion, seizure, vertigo, focal weakness, pain (especially in the back or joints), visual disturbances, speech difficulty, marbled rash, numbness, tingling, cardiac arrest

BLS
- Remove patient from the water or pressurized environment.
- Treat for hypothermia, as appropriate.
- Apply high-flow oxygen, 15 lpm, regardless of initial oxygen saturation.

ALS
- Establish IV access and administer Lactated Ringer’s, as indicated.
- Pain Management protocol

MC
- Transport to a Hyperbaric Medicine Specialty Center if within a 30-minute transport time. If transport time is greater than 30 minutes, obtain medical consultation with a Hyperbaric Medicine Specialty Center and the closest appropriate facility for a transport destination decision.

Clinical Pearls
- Aeromedical transport may be appropriate for patients with barotrauma.
7.4 Environmental Emergencies: Hazardous Materials Exposure

**Indications**
- Exposure to a known or unknown hazardous material
- Signs and symptoms will vary based on the type of substance, route, and duration of exposure.

**BLS**
- Personnel must be trained and appropriately equipped with proper PPE prior to entering a hazardous materials scene.
- Decontaminate patient(s) as indicated.
- If multiple patients are on-scene, triage using START / JumpSTART Algorithm.
- Protect patient from hypothermia and treat for any signs or symptoms of hypothermia.
- Notify EMRC and receiving hospital(s) with the following information:
  - Number of patients and ETA
  - Type of hazardous material(s) involved
  - Decontamination performed on-scene

**ALS**
- Establish IV/IO access in a clean (decontaminated) area if medication administration is anticipated.
- Administer antidote to specific agent, if available.

**MC**
- Antidote or medication that is outside of the protocols
- Antibiotic specific to agent in mass casualty incident, if available

**Clinical Pearls**
- Ensure medical follow-up within 48 hours for all public safety personnel who come into close contact with hazardous materials. Personnel should be advised of possible delayed symptoms post-exposure.
Indications
- Exposure to a warm temperature environment with any of the following symptoms:
  - **Heat Cramps**: Moist, cool skin, cramps, normal to slightly elevated body temperature
  - **Heat Exhaustion**: Moist, cool skin, cramps, weakness, dizziness, normal to elevated body temperature, nausea
  - **Heat Stroke**: Hot, dry skin (25% of patients will still be moist), seizures, altered mental status, dilated pupils, rapid heart rate, or arrhythmia

**BLS**
- Remove patient from the warm environment.
- Move patient to air-conditioned ambulance and remove excess clothing, as practical.
- If patient is conscious and alert, without nausea, give electrolyte-rich fluid by mouth, if available.
- If heat stroke, aggressively cool patient and place patient in semi-Fowler’s position
  - Apply truncal cool packs and wet towels or sheets
  - Misting exposed skin with water from a spray bottle, if available, and air-conditioner running
- If active cooling is in progress by a team/event physician or athletic trainer, allow cooling to continue prior to making an attempt to transport until the patient has an improved mental status or the temperature has decreased to 102 degrees or less.

**ALS**
- Establish IV access.
- Administer *Lactated Ringer’s*; titrate to a minimum systolic blood pressure of 90 mmHg, or age-appropriate blood pressure.

**MC**
- If any disagreement in the patient care plan with a team/event physician or an athletic trainer occurs, contact an appropriate Base Station for on-line medical consultation.
Environmental Emergencies: Near-Drowning – Adult & Pediatric

**Indications**
- Confirmed or suspected near-drowning
- Patient may also have any of the following symptoms: altered level of consciousness, dyspnea, cyanosis, vomiting, seizures, cardiac arrest.

**BLS**
- Remove patient from water.
- Protect patient from and treat for hypothermia.
- Provide supplemental oxygen, as appropriate, to maintain oxygen sat of 94% or greater.

**ALS**
- Establish IV/IO access.
- Administer Lactated Ringer’s; titrate to systolic blood pressure of 90 mmHg, or age-appropriate blood pressure.
- Identify cardiac rhythm and refer to appropriate cardiac protocol.

**MC**
- If the parent or guardian refuses medical care or transport, clinician shall contact a Pediatric Base Station physician while completing the patient-initiated refusal.

**Clinical Pearls**
- Abdominal thrusts are contraindicated unless the patient has a foreign-body airway obstruction.
- Enter the water only if trained and as a last resort (Reach, Throw, Row, Go with Assistance).
- All near-drowning victims should be transported even if they appear uninjured or think they have recovered from the incident due to risk of delayed pulmonary edema.
Indications
- **Beta-blockers** – bradycardia, hypotension, altered mental status, seizure, hypoglycemia
- **Calcium channel blockers** – bradycardia, hypotension
- **Dystonic reaction** – involuntary, slow, sustained muscular contractions; typical in patients taking an antipsychotic, anti-nausea, or antidepressant medication
- **Opioids** – respiratory depression, apnea, altered mental status/decreased level of consciousness, and/or constricted pupils
- **Organophosphates** – SLUDGE symptoms – salivation, lacrimation, urination, defecation, GI pain/upset, emesis (vomiting); seizures; bradycardia, bronchospasm, bronchorrhea
- **NA channel blockers (including tricyclic antidepressants and phenobarbital)** – altered mental status/decreased level of consciousness, hallucinations, hypotension, dysrhythmias, anticholinergic symptoms (hyperthermia, flushed skin, dilated pupils)
- **Other/Unknown substances** – symptoms are variable; may include nausea, vomiting, diarrhea, altered mental status

**BLS**
- Identify the substance and amount ingested
- **Opioids**: if signs of respiratory depression or hypoventilation are present:
  - Assist ventilations with BVM and provide supplemental oxygen
  - Naloxone (BLS) 2 mg IN, dividing the dose equally between the nares to a maximum of 1 mL per nare OR
  - Naloxone (BLS) 4 mg/0.1 mL IN in one nare.
  - Administer additional doses of naloxone (BLS) to maintain a normal respiratory rate and not necessarily until return of consciousness
  - Monitor patient for recurrent respiratory depression and decreased mental status
- **Other/Unknown substances** – obtain medical consultation and Poison Center recommendations. Medication orders can only be accepted from an approved base station.

**ALS**
- Establish IV access, if appropriate
- **Opioids**: if signs of respiratory depression or hypoventilation are present:
  - Naloxone (ALS) 0.4-2 mg IVP/IO (titrated)/IM/IN. For the intranasal route, divide the 2 mg dose equally between the nares to a maximum of 1 mL per nare OR
  - Naloxone (ALS) 4 mg/0.1 mL IN in one nare.
  - Administer additional doses of naloxone (ALS) to maintain a normal respiratory rate and not necessarily until return of consciousness
  - Monitor patient for recurrent respiratory depression and decreased mental status
- **Calcium channel blocker overdose**: calcium chloride 0.5-1 gram slow IVP over 10 minutes; maximum dose of 1 gram. Note: calcium chloride is contraindicated in calcium channel blocker overdose for a patient taking digoxin.
- **Dystonic reaction**: diphenhydramine 25 mg IV or IM
- **NA channel blockers (including tricyclic antidepressants and phenobarbital)**: sodium bicarbonate 1 mEq/kg IVP bolus initial; 0.5 mEq/kg every 10 minutes
Clinical Pearls
- Do not give anything by mouth to a poisoned patient without medical consultation.
- Call the Poison Center early (preferably via EMRC with a base station also online or 800-222-1222) as the poison specialists may have valuable treatment recommendations.
- For opioid overdoses, patients may become agitated or violent following naloxone administration due to opioid withdrawal, hypoxia, or other the effects of other ingested substances.
- The opioid reversal effect of naloxone is limited to approximately 1 hour. Extended release opioids (methadone, oxycodone extended-release) may have a longer duration of action. These patients are at a higher risk of recurrent respiratory depression.
Indications

- **Beta-blockers** – bradycardia, hypotension, altered mental status, seizure, hypoglycemia
- **Calcium channel blockers** – bradycardia, hypotension
- **Dystonic reaction** – involuntary, slow, sustained muscular contractions; typical in patients taking an antipsychotic, anti-nausea, or antidepressant medication
- **Opioids** – respiratory depression, apnea, altered mental status/decreased level of consciousness, and/or constricted pupils
- **Organophosphates** – SLUDGE symptoms – salivation, lacrimation, urination, defecation, GI pain/upset, emesis (vomiting); seizures; bradycardia, bronchospasm, bronchorrhea
- **NA channel blockers (including tricyclic antidepressants and phenobarbital)** – altered mental status/decreased level of consciousness, hallucinations, hypotension, dysrhythmias, anticholinergic symptoms (hyperthermia, flushed skin, dilated pupils)
- **Other/Unknown substances** – symptoms are variable; may include nausea, vomiting, diarrhea, altered mental status

**BLS**

- Identify the substance and amount ingested
- **Opioids**: if signs of respiratory depression or hypoventilation are present:
  - Assist ventilations with BVM and provide supplemental oxygen
  - Age 28 days to adult: *Naloxone (BLS)* 2 mg IN, dividing the dose equally between the nares to a maximum of 1 mL per nare or
  - *Naloxone (BLS)* 4 mg/0.1 mL IN in one nare.
  - Administer additional doses of *naloxone (BLS)* to maintain a normal respiratory rate and not necessarily until return of consciousness
  - Monitor patient for recurrent respiratory depression and decreased mental status
- **Other/Unknown substances** – obtain medical consultation and Poison Center recommendations. Medication orders can only be accepted from an approved base station.

**ALS**

- Establish IV access, if appropriate
- **Opioids**: if signs of respiratory depression or hypoventilation are present:
  - Age 28 days to adult: *Naloxone (ALS)* 0.1 mg/kg IVP/IO (titrated)/IM/IN. For the intranasal route, divide the 2 mg dose equally between the nares to a maximum of 1 mL per nare or
  - *Naloxone (ALS)* 4 mg/0.1 mL IN in one nare
  - *Naloxone (ALS)* ET dose: 0.2-0.25 mg/kg via endotracheal tube
  - Administer additional doses of *naloxone (ALS)* to maintain a normal respiratory rate and not necessarily until return of consciousness
  - Monitor patient for recurrent respiratory depression and decreased mental status
- **Calcium channel blocker overdose**: calcium chloride 20 mg/kg (0.2 mL/kg) slow IVP (50 mg/min); maximum dose of 1 gram. Note: *calcium chloride* is contraindicated in calcium channel blocker overdose for a patient taking digoxin.
- **Dystonic reaction**: diphenhydramine 1 mg/kg IVP or IM, Maximum single dose 25 mg.
Clinical Pearls

- Do not give anything by mouth to a poisoned patient without medical consultation.
- Call the Poison Center early (preferably via EMRC with a base station also online or 800-222-1222) as the poison specialists may have valuable treatment recommendations.
- For opioid overdoses, patients may become agitated or violent following naloxone administration due to opioid withdrawal, hypoxia, or other the effects of other ingested substances.
- The opioid reversal effect of naloxone is limited to approximately 1 hour. Extended release opioids (methadone, oxycodone extended-release) may have a longer duration of action. These patients are at a higher risk of recurrent respiratory depression.
- EMS clinicians should emphasize the need for transport to the closest appropriate facility for all suspected overdose cases.

For ingested substances, activated charcoal without sorbitol 1 gram/kg PO

**Beta-blocker overdose:**
- 5 years of age and older: glucagon 1 mg IVP every 5 minutes, as necessary
- 28 days to 5th birthday: glucagon 0.5 mg IVP every 5 minutes, as necessary

**Organophosphates:** atropine 0.02 mg/kg IVP/IO or IM. May be repeated every 5-10 minutes until the patient has dry secretions and adequate oxygenation. Maximum dose 2 mg.

**NA channel blockers (including tricyclic antidepressants and phenobarbital):** sodium bicarbonate 1 mEq/kg IVP/IO. For less than 1 year, dilute 1:1 with LR.

**Beta-blocker overdose:**
- 5 years of age and older: glucagon 1 mg IVP every 5 minutes, as necessary
- 28 days to 5th birthday: glucagon 0.5 mg IVP every 5 minutes, as necessary

**Organophosphates:** atropine 0.02 mg/kg IVP/IO or IM. May be repeated every 5-10 minutes until the patient has dry secretions and adequate oxygenation. Maximum dose 2 mg.

**NA channel blockers (including tricyclic antidepressants and phenobarbital):** sodium bicarbonate 1 mEq/kg IVP/IO. For less than 1 year, dilute 1:1 with LR.
Environmental Emergencies: Overpressurization

Indications
- History of SCUBA, breathing in a pressurized environment or altitude chamber or exposure to blast concussion waves can lead to arterial gas embolism.
- Confusion, seizure, vertigo, focal weakness, visual disturbances, speech difficulty, marbled rash, numbness, tingling, pain (particularly pain in the joints and back pain), bleeding from body orifices, cardiac arrest

BLS
- Remove patient from the pressurized environment.
- Treat associated traumatic injuries.

ALS
- Establish IV access and administer Lactated Ringer’s, as indicated.
- Pain Management protocol

MC
- Transport to a Hyperbaric Medicine Specialty Center if within a 30-minute transport time.
  If transport time is greater than 30 minutes, obtain medical consultation with a Hyperbaric Medicine Specialty Center and the closest appropriate facility for a transport destination decision.

Clinical Pearls
- Associated injuries (ruptured eardrum) may make assessment and communication difficult. Symptoms may be slow to present.
- Aeromedical transport may be appropriate for patients with barotrauma.
### Indications
- Localized pain
- Puncture wounds
- Erythema local edema
- Numbness or tingling
- Nausea, vomiting, or diarrhea
- Altered mental status
- Seizures or muscle twitching
- Hypoperfusion
- Metallic or rubbery taste

- For patients with moderate to severe allergic reaction symptoms, or mild symptoms with a history of life-threatening allergic reaction to prior snakebite:
  - Age 5 and older: Epinephrine (BLS) (1 mg/mL) 0.5 mg IM or adult epinephrine auto-injector (BLS)
  - Less than 5 years of age: Epinephrine (BLS) (1 mg/mL) 0.15 mg IM or pediatric epinephrine auto-injector (BLS)
  - Albuterol (BLS) 2.5 mg nebulized or albuterol (BLS) MDI 2 puffs inhaled. For patients less than 2 years of age: albuterol (BLS) 1.25 mg nebulized or albuterol (BLS) 2 puffs inhaled.

- Remove all jewelry on affected extremity
- Immobilize extremity as soon as possible
- Take a picture of the snake, if possible. Do not attempt to capture the snake or transport it to the hospital due to risk of additional envenomation events.

- For a poisonous snakebite, do not: apply distal or proximal constricting bands such as tourniquets, apply ice packs, locally incise the bite, copiously wash the wound, or attempt to remove venom by “sucking” or suctioning

- Establish IV access
  - Lactated Ringer’s 20 mL/kg bolus in uninjured extremity. Titrate to a systolic blood pressure of 90 mmHg for adults or age-appropriate blood pressure for pediatric patients.

- Not applicable
Indications
- Patient with known or suspected stimulant drug use
- Signs and symptoms may include: anxiety, chest pain, hallucinations, hypertension, hyperthermia, respiratory distress, seizures, tachycardia

BLS
- Ensure scene is secure and safe from paraphernalia.
- Identify stimulant name, amount, route, and time introduced into the body, if possible.

ALS
- Establish IV access with Lactated Ringer's.
- For stimulant use associated with severe symptoms or tachycardia (greater than 150 bpm):
  - Administer midazolam 0.1 mg/kg in 2 mg increments slow IVP over 1-2 minutes per increment with maximum single dose of 5 mg.
    - Reduce dose by 50% for patients 69 years of age or older.
    - If IV unavailable, 5 mg IN or IM may be administered. IN administration max 1 mL per nare.
  - Administer midazolam 0.1 mg/kg in 2 mg increments slow IVP over 1-2 minutes per increment with maximum single dose of 5 mg.
    - If IV unavailable, administer 0.2 mg/kg IN to a maximum single dose of 2 mg or 0.2 mg/kg IM to a maximum single dose of 5mg. IN administration max 1 mL per nare.
- Initiate Chest Pain or STEMI protocol, as appropriate, for patients with suspected acute coronary syndrome (ACS) and stimulant toxicity.
- For patients with active seizures, refer to Seizure protocol.

MC
- Additional doses of midazolam may be required to treat patients with stimulant toxicity.

Clinical Pearls
- Supraventricular tachycardia (SVT) associated with stimulant toxicity may resolve with midazolam. Many drugs have longer half-lives than adenosine, which renders it ineffective at terminating SVT associated with stimulant toxicity.
8.1 Indications

- Burns (electrical, thermal, chemical) as evidenced by any of the following: reddening of the skin, deep and intense pain, blisters, mottled appearance, or charred black or brown areas with severe or no pain.

- Extricate the patient from burning vehicles or buildings when safe.
- Stop the burning process; remove wet clothing and dry the patient to prevent hypothermia.
- Administer high concentration of oxygen (if smoke inhalation, refer to Carbon Monoxide protocol).
- Treat associated trauma.
- Cover wounds appropriately with a clean sheet or Mylar® blanket.
- Remove all rings, bracelets, and other jewelry.
- Determine percentage of body surface area (BSA) burned and depth; use of the Palmar method is recommended for pediatrics.
- For burns greater than 10% BSA, follow Cold Emergencies protocol.
- For chemical burns, brush off dry chemical, remove clothing, and flush with water.
- Do not give anything by mouth.
- Do not place ice or ice packs on any patient with burns.

**Destination Determination for Burn Patients:**

- Transport patients who meet any of the following criteria to a burn center:
  - All third-degree burns (full thickness)
  - Second-degree burns (partial thickness) greater than 10% BSA
  - Burns of the face, hands, feet, major joints, genitalia, or perineum
  - Electrical burns, including lightning or contact with high voltage (greater than 120 volts)
  - Suspected smoke inhalation
  - Circumferential burns involving the extremities or torso

- Consider aeromedical transport if the patient is more than 30 minutes by ground from a burn center or hyperbaric medicine specialty center.
- Chemical burns should be transported to the closest appropriate hospital for decontamination prior to referral to a burn center.
- Patients with burns and trauma should be transported to the nearest appropriate trauma center for initial care.
- Children who have not reached 15th birthday who meet burn center criteria should be transported to a pediatric burn center.
For adults, Lactated Ringer’s doses greater than 2,000 mL require medical consultation. If transport time exceeds 30 minutes, obtain medical consultation for maintenance fluid recommendation.

### Clinical Pearls
- Pulse oximetry is not reliable in presence of carbon monoxide or cyanide exposure.
- If suspected smoke inhalation, closely monitor the patient’s airway for delayed airway obstruction, respiratory distress, or oxygen desaturation. The patient may need emergent airway management.
Note: The surface of the patient’s palm equals 1% of their body surface area.
Indications

- Traumatic injury to the eye
- May include the following signs and symptoms in either eye or surrounding facial area:
  - Profuse bleeding
  - Foreign objects
  - Impaled objects and/or soft-tissue damage
  - Avulsions or lacerations

**BLS**

- **Foreign objects NOT embedded in the eye(s):** Flush with copious amounts of water (preferably sterile), normal saline, or *Lactated Ringer’s* from the bridge of the nose outward.
- **Injury to orbits (area around the eye):** Evaluate for need for head stabilization and *Spinal Motion Restriction* protocol.
- **Lacerations/injuries to the eye or globe:** Shield affected eye and dress other eye to reduce movement and protect loss of fluids; consider head stabilization and spinal protection and elevate the head to decrease intraocular pressure.
- **Impaled objects:** Stabilize object, shield affected eye, and dress other eye to reduce movement.
- For isolated eye trauma or injuries, transport to the nearest eye trauma center. If the patient has other associated trauma or burns, transport the patient to the closest appropriate trauma or burn center.

**ALS**

- Establish IV access with *Lactated Ringer’s*, if appropriate.
- Administer opioid per *Pain Management* protocol.

**MC**

- Not applicable.

**Clinical Pearls**

- Never apply pressure to the eye or globe.
- Do not use chemical cold packs on the face.
Indications

**UPPER EXTREMITY:** Patients should have stable vital signs with an isolated upper extremity injury, and no other major or multiple system traumatic injuries.

- Stable patients with an isolated upper extremity injury at or below the mid-humerus
- Complete or incomplete hand or finger amputation (except distal fingertip)
- Degloving, high pressure injection, or crush injury
- Compartment syndrome, suspected (excessive swelling and significant pain to extremity)
- Complicated nerve or vascular injury of the forearm and hand
- High-pressure injection injuries to hand or upper extremity
- Complicated nerve, vessel, or compartment syndrome (excessive swelling and pain of extremity with possible evolving nerve deficit) injury of the forearm and hand

**LOWER EXTREMITY:**

- Complete or incomplete amputation of the lower extremity, ankle, foot
- Degloving, high-pressure injection, or crush injury
- Compartment syndrome, suspected (excessive swelling and significant pain to extremity)
- Complicated nerve or vascular injury of the lower extremity

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**BLS**

- Control bleeding.
  - Apply direct pressure to the area of bleeding.
  - Apply tourniquet early if hypovolemic shock is present and/or bleeding is difficult to control. If bleeding source is unclear, place tourniquet as proximal as possible on the limb.
  - If bleeding from a non-compressible injury (i.e., not able to place a tourniquet to stop bleeding), consider wound packing and/or hemostatic gauze.
- Splint suspected fracture or dislocated extremity or joint. If suspected fracture appears to have compromised perfusion or neurological function, apply gentle traction and splint in anatomic position.
- Package amputated extremity in sealed plastic bag (keep dry) and place on top of ice to keep cool. Do not freeze or submerge in water or freeze amputated part.

**Upper Extremity Destination:**

- Adult patients with isolated, qualifying upper extremity injuries should be referred to Curtis National Hand Center at MedStar Union Memorial Hospital.
- Pediatric patients who have not yet reached their 15th birthday with qualifying upper extremity injuries should be referred to the closest appropriate Pediatric Trauma Center.

**Lower Extremity Destination:**

- Adult patients with qualifying lower extremity injuries should be referred to the closest appropriate Adult Trauma Center.
- Pediatric patients with qualifying lower extremity injuries should be referred to the closest appropriate Pediatric Trauma Center.

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**ALS**

- Establish IV access with Lactated Ringer’s, if appropriate.
- If patient develops hypotension or signs of hemorrhagic shock:
  - Reassess the patient for other injuries. If multiple system trauma or neurotrauma, refer to Multiple/Severe Trauma protocol and transport to closest appropriate trauma center.
  - Treat per Shock: Hypoperfusion protocol.
- Provide pain management per Pain Management protocol.
Clinical Pearls

- Toe injuries from lawn mower are not candidates for reimplantation and patients should be transported to the closest appropriate facility.
- Use time, distance, weather, and proximity to designated trauma center to determine mode of transport. If estimated ground transport time to designated hand center is less than 30 minutes, use ground transport.

Additional fluid beyond 2,000 mL requires medical consultation.
Consultation and acceptance by the Curtis Hand Center (MedStar Union Memorial) is required prior to medevac authorization (before SYSCOM will dispatch the helicopter).
Indications
- Multiple or severe traumatic injuries in patients 15 years of age and older
- Suspected internal bleeding, external bleeding, fractures, or lacerations
- Patients may present with any of the following:
  - Shock or hypotension
  - Hypertension, particularly in head-injured patients
  - Shallow or absent respirations
  - Tachycardia or bradycardia
  - Decreased motor or sensory function in the extremities
- A patient who meets criteria for any category of the Maryland Trauma Decision Tree (Alpha, Bravo, Charlie, Delta)

BLS
- Airway with Cervical Spine Motion Restriction
  - Apply Spinal Motion Restriction protocol for blunt trauma patients. Patients with isolated penetrating trauma should not have spinal immobilization performed.
  - Place an NPA/OPA early, as needed to establish or maintain a patient airway.
- Breathing and Ventilation
  - Provide ventilatory support and oxygen via appropriate method for the patient.
  - Maintain pulse oximetry (SpO₂) greater than or equal to 94%.
  - For a head-injured adult/adolescent (greater than 13 years of age), provide ventilation at a rate of 20 breaths per minute if:
    - Patient has signs of herniation such as unequal pupils, posturing or paralysis, or
    - Patient is manifesting a rapidly decreasing GCS, or
    - With on-line medical consultation
  - Seal open chest wounds with a vented chest seal.
- Circulation with Hemorrhage Control
  - Apply direct pressure to the area of bleeding.
  - If bleeding is life-threatening at any time OR continues despite direct pressure, then attempt wound packing, hemostatic bandages, and/or early tourniquet as appropriate.
  - Apply pelvic stabilization if indicated; use pelvic binder if available.
  - Pregnancy: For pregnant patients greater than 20 weeks gestation with hypotension, provide constant left lateral uterine displacement.

ALS
- Airway with Cervical Spine Motion Restriction as noted in BLS section above
- Breathing and Ventilation
  - Maintain ETCO₂ between 35-40 mmHg for any patient with significant head injury
  - For patients with suspected head injury AND signs of increased intracranial pressure (brainstem herniation), consider adjusting ventilations to achieve an ETCO₂ 30-35 mmHg.
  - If suspected tension pneumothorax, perform needle decompression thoracostomy; once catheters are placed, do not remove.
- Circulation with Hemorrhage Control
  - For patients with a systolic blood pressure greater than or equal to 90 mmHg (greater than or equal to 110 mmHg if injuries include a suspected head injury):
    - Establish IV/IO access.
Clinical Pearls

- While, time, distance, and proximity are all factors to be considered in the triage decision, the trauma decision tree should be used to determine who should be transported to the nearest appropriate trauma center and when the transport should occur.
- For trauma patients who have not reached their 15th birthday, refer to *Trauma: Multiple/Severe (Pediatric)* protocol.

ALS

- For patients with a systolic blood pressure less than 90 mmHg (less than 110 mmHg if injuries include a suspected head injury):
  - Establish IV/IO access.
  - Administer small boluses of *Lactated Ringer’s* (maximum single bolus of 250 mL prior to additional blood pressure check) to achieve and maintain a systolic blood pressure of greater than or equal to 90 mmHg (110 mmHg if injuries include a suspected head injury).
- For patients 15 years of age and older with suspected hemorrhagic shock (SBP less than 90) due to trauma, administer *TXA* 1 gram in 100 mL of approved diluent (normal saline/*Lactated Ringer’s*/D5W) IV/IO over 10 minutes. Injury must have occurred within the past one (1) hour. Do not delay transport to initiate *TXA*.
  - Initiate a second IV for category alpha and bravo patients ONLY if it does NOT delay transport.
  - Treat per *Pain Management* protocol.

MC

- Additional fluid beyond 2,000 mL requires medical consultation
Indications
- Multiple or severe traumatic injuries in patients less than 15 years of age
- Suspected internal bleeding, external bleeding, fractures, or lacerations
- Patients may present with any of the following:
  - Shock or hypotension
  - Hypertension, particularly in head-injured patients
  - Shallow or absent respirations
  - Tachycardia or bradycardia
  - Decreased motor or sensory function in the extremities
- A pediatric patient who meets criteria for any category of the Maryland Trauma Decision Tree (Alpha, Bravo, Charlie, Delta)

### BLS

- **Airway with Cervical Spine Motion Restriction**
  - Apply *Spinal Motion Restriction* protocol for blunt trauma patients. Patients with isolated penetrating trauma should not have spinal immobilization performed.
  - Place an NPA/OPA early, as needed to establish or maintain a patient airway.
- **Breathing and Ventilation**
  - Provide ventilatory support and oxygen via appropriate method for the patient.
  - Maintain pulse oximetry (SpO₂) greater than or equal to 94%.
  - For a head-injured patient who meets any of the following criteria below, provide ventilation as follows:
    - Criteria:
      - Patients who have signs of herniation such as unequal pupils, posturing, or paralysis, or
      - Patient is manifesting a rapidly decreasing GCS, or
      - With on-line medical consultation
    - Ventilatory rate if any criteria above are met:
      - Adult/Adolescent (greater than 13 years of age): 20 breaths per minute
      - Child (1-12 years of age): 30 breaths per minute
      - Infant (less than 1 year of age): 35 breaths per minute
- Seal open chest wounds with a vented chest seal.

- **Circulation with Hemorrhage Control**
  - Apply direct pressure to the area of bleeding.
  - If bleeding is life-threatening at any time OR continues despite direct pressure, then attempt wound packing, hemostatic bandages, and/or early tourniquet as appropriate.
  - Apply pelvic stabilization if indicated

### ALS

- **Airway with Cervical Spine Motion Restriction** as noted in BLS section above
- **Breathing and Ventilation**
  - If suspected *tension* pneumothorax, perform *needle decompression thoracostomy*; once catheters are placed, do not remove.
- **Circulation with Hemorrhage Control**
  - Establish IV/IO access
  - If age-related vital signs and patient’s condition indicate hypoperfusion, administer initial fluid bolus of 20 mL/kg *LR* IV/IO. If patient’s condition does not improve, administer the second bolus of fluid at 20 mL/kg *LR*.
  - Treat per *Pain Management* protocol
Clinical Pearls

- Pelvic fractures in pediatric patients are rare, pelvic binders/splints should be applied with caution.
Indications
- Abrasions, contusions, and/or bleeding
- Signs of forcible restraint
- Petechiae of the face and conjunctiva, secondary to strangulation
- Facial injuries, including eye injuries, broken teeth, swollen jaw, or cheekbone
- Vaginal or rectal bleeding or pain
- Some patients may present without visible signs of trauma

BLS
- If practical, allow patient to speak with a clinician with whom they are most comfortable.
- Maintain a non-judgmental, caring attitude.
- Preserve the crime scene and clothing articles, if practical.
- Do not perform an examination of the genitals or rectum unless necessary to stabilize the patient.
- Dress wounds (do not attempt to clean).
- Discourage any self-treatment (shower, washing, changing clothes, brushing teeth).
- Treat injuries according to appropriate trauma protocol.
- Destination
  - Patients meeting specialty center criteria or in need of time-sensitive emergent care should be transported to the closest appropriate specialty center or emergency department, even if this is not a Maryland Coalition Against Sexual Assault (MCASA) recognized facility.
  - Patients under 13 years of age should be transported to an MCASA-recognized pediatric facility for a Sexual Assault Forensic Exam (SAFE).
  - For patients 13 years of age and older, transport the patient to the appropriate MCASA-recognized facility for a SAFE exam. Use the term “safe patient” when notifying the receiving facility.
- Reporting
  - All EMS clinicians must report cases of suspected child or vulnerable adult abuse or neglect directly to the local police or adult/child protective services. This report is required by law. Do not initiate the report in the presence of the patient, parent, or caregiver.

ALS
- Treat with ketorolac or opioid, as needed, per Pain Management protocol.

MC
- Not applicable.

Clinical Pearls
- EMS clinicians are protected from liability if they make a report of child or vulnerable adult abuse or neglect in good faith.
**Indications**

- Patients who have a blunt trauma with a high-energy mechanism of injury that has potential to cause spinal cord injury or vertebral instability and one or more of the following should receive spinal motion restriction:
  - Midline cervical, thoracic, or lumbar spinal pain, tenderness, or deformity
  - Signs and symptoms of new paraplegia or quadriplegia
  - Focal neurological deficit (sensory or motor)
  - Altered mental status or disorientation
  - Distracting injury: Any injury (e.g., fracture, chest, or abdominal trauma) associated with significant discomfort that could potentially distract from a patient’s ability to accurately discern or define spinal column pain or tenderness.

- Indications for referral to an Adult Neurotrauma Center:
  - 15 years of age or older AND
  - Signs and symptoms of new paraplegia or quadriplegia in the presence of trauma AND
  - Patent airway AND
  - Hemodynamically stable

- If considering referral to Adult Neurotrauma Center, consult with both the nearest Trauma Center and the Adult Neurotrauma Center, when possible.

**BLS**

- Minimize flexion, extension, and rotation of the spinal column.

- **Cervical collar:** The following patients need application of a cervical collar and do **not** need full immobilization with a backboard. These patients should be assisted with minimal movement to the EMS stretcher and allowed to lie supine on their own accord with head elevated at 30 degrees:
  - Patients who are found by EMS clinicians to be standing or ambulatory,
  - Patients who have a GCS of 15 and are able to safely extricate themselves from the environment (e.g., vehicle seat) without gross movement (flexion, extension, rotation) of the spinal column, and
  - Patients who do not have evidence of a neurological deficit.

- **Cervical collar and backboard:** Patients with neurological deficit or a GCS of less than 15 or who are not able to ambulate on their own accord shall be immobilized with cervical collar and a backboard.

- **Extrication:** Backboards may be used for patient extrication and transfer for patients not meeting the *Spinal Motion Restriction* protocol; however, other devices are preferred (e.g., Reeves™, scoop stretcher).
  - If the backboard is used only for extrication from scene to ambulance, remove the backboard as soon as possible and allow the patient to be supported on the EMS stretcher.

- **Interfacility transport:** Patients who have already been removed from the backboard should not be placed back on one for transport.

- **Found on backboard prior to EMS arrival:** If the patient was immobilized on a backboard prior to EMS arrival, EMS should assess continued need for the device using the criteria above.
Helmet Removal
- If patient is wearing a helmet, the goals are assessment and management of the airway, breathing, and circulation followed by protection of the spinal column by maintaining neutral alignment of the spinal column.
- If patient is wearing helmet and no shoulder pads, removal of the helmet is indicated.
- If patient is wearing helmet with shoulder pads, removal of the helmet is acceptable only with concurrent removal of shoulder pads. Under these conditions, removal of the helmet is indicated for management of the airway or other facial trauma.

Helmet Removal
- If the patient presents with hypotension and concern for neurogenic shock, refer to the neurogenic shock section of the Shock/Hypoperfusion protocol.

Not applicable.
Indications
- Patients who have a blunt trauma with a high-energy mechanism of injury that has potential to cause spinal cord injury or vertebral instability and the presence of or inability to assess one or more of the following should receive spinal motion restriction:
  - Midline spinal pain, tenderness, or deformity
  - Signs and symptoms of new paraplegia or quadriplegia
  - Focal neurological deficit
  - Altered mental status or disorientation
  - Distracting injury
  - Neck pain or torticollis
  - High-impact diving incident or high-risk motor vehicle crash (i.e., head-on collision, rollover, ejected from the vehicle, death in the same crash, or speed greater than 55 mph)
  - Substantial torso injury
  - Conditions predisposing to spine injury
- Indications for referral to a Pediatric Trauma Center:
  - Patient is less than 15 years of age AND
  - Signs and symptoms of new paraplegia or quadriplegia in the presence of trauma AND
  - Patent airway AND
  - Hemodynamically stable

BLS
- Minimize flexion, extension, and rotation of the spinal column
- Cervical collar: The following patients need application of a cervical collar and do not need full immobilization with a backboard. These patients should be assisted with minimal movement to the EMS stretcher and allowed to lie supine on their own accord with head elevated at 30 degrees:
  - Patients who are found by EMS clinicians to be standing or ambulatory,
  - Patients who have a GCS of 15 and are able to safely extricate themselves from the environment (e.g., vehicle seat) without gross movement (flexion, extension, rotation) of the spinal column, and
  - Patients who do not have evidence of a neurological deficit.
- Cervical collar and backboard: Patients with neurological deficit or a GCS of less than 15 or who are not able to ambulate on their own accord, shall be immobilized with cervical collar and a backboard.
- Extrication: Backboards may be used for patient extrication and transfer for patients not meeting the Spinal Motion Restriction protocol; however, other devices are preferred (e.g., Reeves™, scoop stretcher).
  - If the backboard is used only for extrication from scene to ambulance, remove the backboard as soon as possible and allow the patient to be supported on the EMS stretcher.
- Interfacility transport: Patients who have already been removed from the backboard should not be placed back on one for transport.
- Found on backboard prior to EMS arrival: if the patient was immobilized on a backboard prior to EMS arrival, EMS should assess continued need for the device using the criteria above.
Helmet Removal

- If the patient is wearing a helmet, the goals are assessment and management of the airway, breathing, and circulation followed by protection of the spinal column by maintaining neutral alignment of the spinal column.
- If the patient is wearing a helmet and no shoulder pads, removal of the helmet is indicated.
- If the patient is wearing a helmet with shoulder pads, removal of the helmet is acceptable only with concurrent removal of shoulder pads. Under these conditions, removal of the helmet is indicated for management of the airway or other facial trauma.

If the patient presents with hypotension and concern for neurogenic shock, refer to the neurogenic shock section of the Shock/Hypoperfusion protocol.

Not applicable.
High-risk mechanism of blunt trauma AND one or more of the following will receive a minimum of a cervical collar
All patients

- Midline cervical, thoracic or lumbar spinal pain, tenderness, or deformity
- New paraplegia or quadriplegia
- Focal neurological deficit (sensory or motor)
- Altered mental status or disorientation or intoxication
- Distracting injury: Any injury (e.g., fracture, chest or abdominal trauma) associated with significant discomfort that could potentially distract from a patient's ability to accurately discern or define spinal pain or tenderness

Additionally, for patients who have not yet reached their 15th birthday
- Neck pain or torticollis
- High-impact diving incident or high-risk MVC
- Substantial torso injury
- Conditions predisposing to spine injury
- Inability to assess any of above

If NO to all

Spinal precautions not indicated.

If yes to any of the above, minimum of cervical collar

Does the patient have one or more of the following?
- Neurological deficit sensory/motor or GCS less than 15
- Inability to ambulate
- Unable to respond during assessment

NO

SPINAL PRECAUTIONS
Apply cervical collar only

YES

SPINAL IMMOBILIZATION
Perform complete spinal immobilization including cervical collar and long backboard
Indications
- Cardiac arrest with suspected traumatic etiology in a patient who is 15 years of age or older

BLS
- Extricate patient if the environment is unsafe, otherwise initiate care on-scene.
- Determine if patient meets the criteria for Pronouncement of Death in the Field protocol. If criteria are not met, then continue resuscitation.
- Perform spinal motion restriction for blunt trauma patients only.
  - Patients with isolated penetrating trauma should not have spinal motion restriction.
  - If the mechanism of injury includes both blunt and penetrating trauma, perform spinal motion restriction.
- Perform CPR with high-quality chest compressions and minimal interruptions and apply an AED.
- Treat reversible causes of traumatic arrest, on-scene, as soon as possible.
  - Open airway and ensure adequate ventilation, insert necessary adjunct; consider the need for advanced airway earlier in the resuscitation of the trauma arrest patient.
  - Seal open chest wounds with vented chest seals.
  - Control life-threatening external hemorrhage.
- Destination
  - For a penetrating trauma patient in a rhythm other than asystole:
    - Within 15 minutes of a trauma center: Treat reversible causes and transport.
    - Greater than 15 minutes from a trauma center: Treat reversible causes and transport to the closest ED or freestanding emergency medical facility.
  - For a blunt trauma patient: All reversible causes of arrest should be performed on-scene before TOR or transport (if ROSC was achieved).
- If ROSC is not achieved, refer to Termination of Resuscitation (Adult Traumatic) protocol.

ALS
- Establish IV/IO access.
- Initiate LR 20 mL/kg rapid bolus IV/IO
- Treat reversible causes of traumatic arrest, on-scene, as soon as possible.
  - BLS interventions as indicated above
  - Bilateral needle decompression thoracostomy: For traumatic arrest due to suspected multi-system blunt trauma, or due to penetrating neck, chest, or abdominal trauma, bilateral needle decompressions should be performed. Catheters should not be removed once placed.
- Identify rhythm and refer to appropriate treatment algorithm.
  - Note: Epinephrine is generally not indicated for adult traumatic arrest.
- If ROSC is not achieved, refer to Termination of Resuscitation (Adult Traumatic) protocol.

MC
- Not applicable.

Clinical Pearls
- Early cardiac arrest secondary to trauma is usually due to severe hypoxia, neurologic injury, or massive hemorrhage. Address the ABCs first.
Indications

- Cardiac arrest with suspected traumatic etiology in a patient who is less than 15 years of age
- Extricate patient if the environment is unsafe, otherwise initiate care on-scene.
- Determine if patient meets the criteria for Pronouncement of Death in the Field protocol. If criteria are not met, then continue resuscitation.
- Perform spinal motion restriction for blunt trauma patients only.
  - Patients with isolated penetrating trauma should not have spinal motion restriction.
  - If the mechanism of injury includes both blunt and penetrating trauma, perform spinal motion restriction.
- Perform CPR with high-quality chest compressions and minimal interruptions and apply an AED.
- Treat reversible causes of traumatic arrest, on-scene, as soon as possible.
  - Open airway and ensure adequate ventilation, insert necessary adjunct; consider the need for advanced airway earlier in the resuscitation of the trauma arrest patient.
  - Seal open chest wounds with vented chest seals.
  - Control life-threatening external hemorrhage.
- Destination
  - For a penetrating pediatric trauma patient in a rhythm other than asystole:
    - Within 15 minutes of a Pediatric Trauma Center: Treat reversible causes and transport.
    - Greater than 15 minutes from a Pediatric Trauma Center: Treat reversible causes and transport to the closest Adult Trauma Center, ED, or freestanding emergency medical facility.
  - For a blunt trauma patient: all reversible causes of arrest should be performed on-scene before TOR or transport (if ROSC was achieved).
- If ROSC is not achieved, refer to Termination of Resuscitation (Pediatric Traumatic) protocol.

Clinical Pearls

- Early cardiac arrest secondary to trauma is usually due to severe hypoxia, neurologic injury, or massive hemorrhage. Address the ABCs first.

{Back to Contents}
Trauma Arrest: Pediatric Termination of Resuscitation
(Patients Under 15 Years of Age)

Cardiac Arrest
(Considering Termination of Resuscitation)

Exclusions:
- Pregnant
- Hypothermia or submersion

NO

Meets Pronouncement of Death criteria?

YES

Should terminate resuscitation

NO

Minimum of 5 two-minute cycles of HPCPR—treat reversible causes of arrest

Rhythm

Asystole AND ETCO$_2$ less than 15 mmHg?

May terminate resuscitation

VF, VT, PEA
OR
Asystole with ETCO$_2$ greater than 15 mmHg?

Less than 15 min. to Pediatric Trauma Center

HPCPR/Treat reversible causes, transport to Pediatric Trauma Center

More than 15 min. to Pediatric Trauma Center

HPCPR/Treat reversible causes, transport to local ED or Adult Trauma Center

Continue resuscitation and safe transport
Trauma Decision Tree

Measure vital signs and level of consciousness and assess for major injury

**Category Alpha**

- GCS less than or equal to 13
- For patients 10 years and older (including adults), systolic blood pressure less than 90 mmHg.
- For patients under 10 years of age, systolic blood pressure less than 70 + 2x age in years mmHg.
- Respiratory rate less than 10 or greater than 29 (less than 20 in infant age less than one year) or need for ventilatory support

**YES**

Transport to trauma center or specialty center per protocol; alert trauma team; consider helicopter transport if quicker and of clinical benefit (refer to GPC Section I).

**NO**

Assess for other injuries.

**Category Bravo**

- 2 or more proximal long-bone fractures
- Crushed, degloved, mangled, or pulseless extremity
- Ampullation proximal to wrist or ankle
- Open or depressed skull fracture
- Paralysis (spine)
- Chest wall instability or deformity (e.g., flail chest)
- Penetrating injuries to head, neck, torso, or extremities proximal to elbow and knee

**YES**

Transport to trauma center or specialty center per protocol; alert trauma team; consider helicopter transport if quicker and of clinical benefit (refer to GPC Section I).

**NO**

Evaluate for evidence of mechanism of injury and high-energy impact.

**Category Charlie**

- High risk Auto Crash
  - Intrusion (including roof) greater than 12 in. occupant site; greater than 18 in. any site
  - Ejection (partial or complete) from vehicle
  - Death in same passenger compartment
  - Vehicle telemetry data consistent with high risk of injury
- Falls
  - Adult: greater than 20 feet (one story is equal to 10 feet)
  - Pediatric: greater than 10 feet or 3 times the child’s height
- Exposure to blast or explosion

**YES**

Transport to Trauma Center; alert trauma team. Patients within a **30-minute drive time** of the closest appropriate trauma/specialty center shall go by ground unless there are extenuating circumstances. Receiving Trauma Center medical consultation required when considering whether helicopter transport is of clinical benefit (refer to GPC Section I).

**NO**

Evaluate for other considerations.

**Category Delta**

- Older adults
  - Risk of injury/death increases after age 55
  - SBP less than 110 may indicate shock after age 65
  - Low-impact mechanisms (e.g., ground-level falls) may result in severe injury
- Children
  - Should be triaged to Pediatric Trauma Center
- Burns
  - Without trauma mechanism, triage to Burn Center
  - With trauma mechanism, triage to Trauma Center
- Pregnancy greater than 20 weeks
- EMS clinician judgment
- Anaphylaxis and bleeding disorders
  - Patients with head injury are at high risk for rapid deterioration

**YES**

Consider medical direction and transport to trauma center. Patients within a **30-minute drive time** of the closest appropriate trauma/specialty center shall go by ground unless there are extenuating circumstances. Receiving Trauma Center medical consultation required when considering whether helicopter transport is of clinical benefit (refer to GPC Section I).

**NO**

Transport according to protocol.
Appendices

Pharmacology
Procedures
Interfacility Protocols
Pilots
Optional Supplemental Protocols
Research Protocols
AED: Automated External Defibrillation or Automated External Defibrillator

**Alternative Airway Device:** An airway adjunct other than an endotracheal tube that may include the laryngeal tube airway device (e.g., King LTS-D™) or laryngeal mask airway with design to facilitate hospital endotracheal intubation

AMI: Acute Myocardial Infarction

**APGAR score:** An acronym and method of scoring to determine the condition of a newly born infant (see *APGAR Chart* in Protocol 5.4)

Apnea: An absence of spontaneous respirations

**Aspiration:** The act of taking fluid (e.g., vomitus, mucus, or blood) from the body via a suction device or the act of taking foreign material or vomit into the lungs

**Asymptomatic:** The lack of any evidence or indication of illness, disease, or physical disturbance of patient’s condition

**AVPU:** A method of determining and recording a patient’s mental status or level of consciousness where “A” stands for Alert, “V” stands for responsive to Verbal stimuli, “P” stands for responsive to Painful stimuli, and “U” stands for Unresponsive

**Barotrauma:** Injury sustained as a result of exposure to excessive environmental pressure changes (e.g., blast injury or underwater pressure injury)

BPM: Breaths per minute

BSI: Body Substance Isolation

BVM: Bag-Valve-Mask

**Carte blanche:** Full discretionary power

**Children with Special Healthcare Needs (CSHN):** Children with chronic illness or conditions requiring specialized assessment, treatment, technology, or transport destination

CISM: Critical Incident Stress Management

**Clinician:** Includes EMR, EMT, CRT-(I), and paramedic

**Commercial ambulance:** Ambulance licensed by the State Office of Commercial Ambulance Licensing and Regulation

**Continuous CPR:** Chest compressions asynchronous with ventilation and infrequent, minimal interruptions (less than 10 seconds each)

**COPD:** Chronic Obstructive Pulmonary Disease (e.g., asthma, emphysema, bronchitis)
Cricothyroidotomy (needle or surgical): A syringe with a needle attached or a scalpel is used to make a puncture hole or surgical incision through the cricothyroid membrane that overlies the trachea. A needle catheter or ET tube is passed into the trachea and then attached to a jet insufflation device or bag-valve device to ventilate the patient.

Critical: Approaching death or having the nature of a crisis (e.g., time-critical, critical injury)

CRT-(I): Cardiac Rescue Technician-Intermediate

CVA: Cerebral Vascular Accident/Stroke

Cyanotic: Bluish color of the skin or mucus membranes caused by lack of oxygen to the tissue

DCAP BTLS: Acronym for signs of injuries to assess during a physical examination of patients: D = Deformity, C = Contusions, A = Abrasions, P = Punctures/penetrations, B = Burns, T = Tenderness, L = Lacerations, S = Swelling

Defibrillation: Administration of electrical current(s) to the heart in an effort to normalize rhythm

Defibrillation set (stacked shocks): Includes a set of three successive shocks either biphasic or monophasic standard 200 J, 300 J, 360 J, or peds 2–4 J/kg

Distracting Injury: Any injury (e.g., fracture, chest, or abdominal trauma) associated with significant discomfort that could potentially distract from a patient’s ability to accurately discern or define spinal column pain or tenderness

DNR: Do Not Resuscitate

Dystonic: Any impairment of muscle tone, which may be manifested by prolonged muscle contractions that may cause twisting and repetitive movements or abnormal posture. These movements may be in the form of rhythmic jerks. Symptoms that “appear” to be of a focal seizure-like nature in an awake and alert person with no history of seizures but who probably has a recent history of anticholinergic medication use (e.g., anti-psychotic, anti-vomiting).

EJ: External Jugular vein of the neck; peripheral IV access site

Emergency Information Form: A two-page form, designed by the American Academy of Pediatrics and American College of Emergency Physicians (AAP and ACEP, respectively), that provides a brief summary of special health care needs including: diagnosis, usual pattern of disease, emergency action plan, primary and specialty doctors and hospitals. Can be downloaded and data entered at http://www.aap.org/advocacy/eif.doc.

Emetic: Referring to a substance that causes vomiting

eMEDS®: electronic Maryland EMS Data System (a patient care reporting system)
EMR: Emergency Medical Responder

EMS: Emergency Medical Services

EMT: Emergency Medical Technician

EOC: Emergency Operations Center

Erythema: Redness or inflammation of the skin or mucous membranes that is the result of dilatation and congestion of superficial capillaries

ETA: Estimated Time of Arrival

ETCO₂: Non-invasive measurement (numeric and/or waveform) of carbon dioxide (CO₂) levels in exhaled breaths (end-tidal CO₂)

Extrapyramidal: Pertaining to tissues and structures outside of the cerebrospinal pyramidal tracts of the brain that are associated with movement of the body, excluding stimulation from the motor neurons, the motor cortex, and the corticospinal and corticobulbar tracts. Symptoms that “appear” to be of a focal seizure-like nature in an awake and alert person with no history of seizures but who probably has a recent history of anticholinergic medication use (e.g., anti-psychotic, anti-vomiting).

Fluid Bolus: The administration of a fluid dose as rapidly as possible, usually over five to twenty minutes, to a patient with clinical signs of shock

GCS: Glasgow Coma Scale (a tool to evaluate injury and illness severity)

Hemodynamically Stable: When a patient’s vital signs (including pulse oximeter or EKG if available) are all within normal for the patient’s age range, the patient does not have active bleeding, and there are no signs of distress (skin conditions or capillary refill are normal) as observed over time

Hemodynamically Unstable: When a patient exhibits any of the following: abnormal vitals signs for age range (including pulse oximeter or EKG if available), active bleeding, or there are signs of distress (skin conditions or capillary refill are abnormal)

Hemostatic Dressing: A bandage or gauze with impregnated hemostatic agent that hastens the hemostasis/clotting process

HTN: Hypertension

Hypoxia: Too little oxygen in the cells

IM: Intramuscular injection

IN: Intranasal administration

IV: Intravenous line or administration of medication through IV
IVP: Intravenous Push

IWMI: Inferior Wall Myocardial Infarction

J: Joules or watts-seconds of electrical energy for defibrillation or cardioversion

JVD: Jugular Vein (external) Distention

kg: Kilogram, metric measure of weight equal to 1,000 grams (1 kg = 2.2 pounds)

KVO: Keep Vein Open. A slow IV flow rate.

Laryngectomy: The removal of the larynx and separation of the airway from the mouth, nose, and esophagus. Patients with a laryngectomy breathe through an opening in the neck called a stoma. Patients with a laryngectomy are not able to breathe or be intubated through the mouth or nose.

Lividity: Venous pooling in dependent body parts

LOC: Level of Consciousness

LR: Lactated Ringer’s (a type of isotonic IV solution)

MCI: Mass Casualty Incident. Occurs when the number of victims exceeds the number of medical personnel or resources immediately available and is declared by the local jurisdiction.

Meconium: The first feces of an infant

Medical Consultation (On-Line Medical Direction): With an atmosphere of courtesy and respect, direct voice/data communication between a clinician and an EMS Base Station physician, or a jurisdictionally affiliated physician, or with an “on-scene physician.” This communication is bi-directional and provides the clinician with medical direction while providing the physician or the receiving hospital with valuable information on the patient. This exchange can take place on-scene, over a telecommunications device, or in the hospital setting.

Medical Protocol: A guideline for the provision of patient care

mL: Milliliter (the symbol for a metric measure of volume)

MOI: Mechanism of Injury

MOLST: Medical Orders for Life-Sustaining Treatment

NDT: Needle Decompression Thoracostomy

Near Drowning: A short duration of submersion under water with possible short-term loss of consciousness

Neonatal (also neonate): A term that describes an infant from 1 hour to 28 days of life
**Newly Born (also called newborn):** A term that describes an infant within the first hour after delivery.

**NOI:** Nature of Illness

**Notification:** An “information only call” directly to the receiving hospital through the jurisdictional EOC or EMS communication system not requiring medical consultation and that may follow local standing operational procedures.

**NRB:** Non-Rebreather Mask

**NTG:** Nitroglycerin

**Nurse Practitioner:** An individual who has been licensed as a Registered Nurse and certified as a Nurse Practitioner by the Maryland Board of Nursing. This does not include individuals who are only Registered Nurses or Licensed Practical Nurses.

**OIC:** Officer in Charge

**On-Scene Physician:** On-Scene Physician may be the patient’s identified private physician or a bystander physician who is physically on location. Care rendered or orders given by the on-scene physician should be documented, including the identification of the physician. All on-scene medical direction shall be consistent with *The Maryland Medical Protocols for Emergency Medical Services*. Any medical procedure that is not consistent with the protocols shall only be rendered by the on-scene physician, who shall also accompany the patient to the hospital. Any extraordinary care by EMS clinicians pursuant to the protocols may be approved only by the EMS Base Station physician or a system medical director (based on COMAR 30.02.03.02A.).

**OPQRST:** Used to recall pertinent questions (Onset, Provocation, Quality, Radiation, Severity, Time) to ask when obtaining a patient history for medical emergencies

**Optional Supplemental Program (OSP):** A voluntary jurisdictional program that requires MIEMSS approval

**Pallor:** An unnatural paleness or absence of color in the skin

**PCM:** Patient Controlled Medications (a medication delivery system under a patient’s control)

**PCR:** Patient Care Report (equivalent to MAIS) document used to record pertinent patient information regarding assessment, treatment, and transport (this is a confidential medical record)

**Pilot Program (PP):** A program designed to test a new project or procedure in order to determine its effect on EMS (requires MIEMSS approval and reporting all uses to MIEMSS)
Plethora: A term applied to the beefy red coloration of a newborn

PMD: Program Medical Director

PO: By mouth (*per os*)

PPE: Personal Protective Equipment

Pulse Oximetry: A non-invasive measurement of arterial oxygen saturation using infrared absorption frequencies

PVC: Premature Ventricular Contraction

Recovery Position: The position (patient flat on left lateral side) or placement of patients to reduce risk of aspiration

RMD: Regional Medical Director

RVMI: Right Ventricular Myocardial Infarction

SAFER: Stabilize, Assess and acknowledge, Facilitate, Encourage, and Recovery OR Referral

SAMPLE: Used to aid in obtaining pertinent patient history (S = Symptoms and signs patient is exhibiting, A = patient Allergies, M = patient Medications (prescription and non-prescription), P = Past medical history, L = what and when was the patient’s Last oral intake, E = Events prior to arrival, or simply, the history of the current emergency)

SC: Subcutaneously

Sign: Any objective evidence or indication of illness, disease, or physical disturbance of patient’s condition

SL: Sublingual (under the tongue)

SMOI: Significant Mechanism Of Injury

SOP: Standard Operational Procedure (defined by local jurisdiction or region)

Spinal Motion Restriction: The act of placing a patient on a backboard with cervical collar for the purpose of trying to prevent excessive movement of the spinal column

Spinal Protection: The act of protecting the spinal cord from further injury

Standing Orders: Orders, rules, regulations, or procedures prepared as guidelines in the preparation and carrying out of medical and surgical procedures
**Sublingually:** Under the tongue

**Symptom:** Any subjective evidence of disease or of a patient’s condition (such as evidence perceived by the patient)

**Symptomatic:** The subjective evidence or indication of illness, disease, or physical disturbance of patient’s condition

**Syncope:** A fainting spell. It usually follows a feeling of lightheadedness and may often be prevented by lying down. Syncope may also result from any number of heart, neurologic, or lung disorders.

**System Medical Director:** Means any of the following: Executive Director of MIEMSS, State EMS Medical Director, Associate State Medical Director for Pediatrics, Regional Medical Directors, Associate Regional Pediatric Medical Directors, EMS Operational Program Medical Directors, and Assistant EMS Operational Program Medical Directors

**TOI:** Type of Incident to which EMS clinicians may be called upon to respond (e.g., ill and/or injured patients, hazardous materials incidents, fires, mass casualty incidents)

**Tracheostomy:** An incision into the trachea (windpipe) that forms a temporary or permanent opening called a stoma. A tube is inserted through the opening to allow passage of air and removal of secretions.

**Vagal:** Pertaining to the vagus nerve (the tenth cranial nerve, which is essential for speech, swallowing, and slowing of the heart rate)

**VF:** Ventricular Fibrillation

**Volume-Sensitive Children:** Children who need smaller fluid bolus volumes due to special needs including: neonates (1 hour to 28 days of age), congenital heart diseases, chronic lung disease, or chronic renal failure

**VT:** Ventricular Tachycardia

**Vulnerable Adult:** An adult who lacks the physical or mental capacity to provide for his or her daily needs (Digest of Criminal Law)
## Appendices – PROCEDURES, MEDICAL DEVICES, AND MEDICATIONS FOR EMS AND COMMERCIAL SERVICES

### 9.2

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**SO** Standing Order

**OSP** Optional Supplemental Program

**MC** Medical Consultation Required

**PP** Pilot Program

**REA** Research
## Appendices – PROCEDURES, MEDICAL DEVICES, AND MEDICATIONS FOR EMS AND COMMERCIAL SERVICES (continued)

<table>
<thead>
<tr>
<th>PROCEDURE</th>
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<td>HALO Cervical Immobilization</td>
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<td>IABP InterAortic Balloon Pump</td>
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<td>Ileostomy Tube (Non-infusing)</td>
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<td>PICC-peripherally inserted central catheter or CVA -</td>
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<td>Peak Expiratory Flow Meter</td>
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<td>Portable Outpatient Fixed Medication Pump/PCA Pump</td>
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<td>Peritoneal Dialysis (Non-active, Capped)</td>
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<td>Surgical Drains</td>
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<td>Transvenous Pacemaker (Temporary Transvenous)</td>
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<td>Ventilators (Acute, Chronic, Scene)</td>
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<td>Ventricul Peritoneal Shunt</td>
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<td>Wound Vacuum Device</td>
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**SO** Standing Order  
**OSP** Optional Supplemental Program  
**MC** Medical Consultation Required  
**PP** Pilot Program  
**REA** Research

*If being discharged home or to rehab, may go by ALS.
### Appendices – PROCEDURES, MEDICAL DEVICES, AND MEDICATIONS FOR EMS AND COMMERCIAL SERVICES (continued)

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<thead>
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<th>Medications</th>
<th>EMR</th>
<th>EMT</th>
<th>CRT</th>
<th>PM</th>
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<td>Acetaminophen</td>
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<td>Activated Charcoal (Without Sorbitol)</td>
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<td>Adenosine</td>
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<td>Droperidol</td>
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<td>Epinephrine (1:1,000) Vial or Syringe</td>
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<td>Etomidate (Amidate)</td>
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**Abbreviations:**
- **SO**: Standing Order
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*Release Date July 1, 2022*
## MEDICATIONS

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<td>Nitroglycerin Paste</td>
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<td>Nitroglycerin (tablet/spray) (patient’s prescribed)</td>
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<td>Oral Glucose</td>
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<td>Oxygen</td>
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<td>Vaccines (Hepatitis B, Influenza, and COVID-19)</td>
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<td>Verapamil</td>
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TRADE NAMES: Tylenol®

a) Indications
Patients 3 months of age and older with:
(1) Mild to moderate discomfort (e.g., 1–5 on FACES scale) or
(2) Fever (EMS-documented temperature greater than or equal to 100.4 F / 38 C)

b) Adverse Effects
Not clinically significant

c) Precautions
(1) Administration of acetaminophen for mild to moderate pain does not eliminate
the need for transport of the patient to an appropriate facility capable of conducting
a comprehensive evaluation of the cause of the pain and appropriate definitive
treatment.
(2) A 3 mL, 5 mL, or 6 mL syringe must be used to measure doses of acetaminophen.

d) Contraindications
(1) Head Injury
(2) Hypotension
(3) Administration of acetaminophen or medications containing acetaminophen within
the previous 4 hours. Many common cold preparations contain acetaminophen.
(4) Inability to swallow or take medications by mouth
(5) Respiratory distress
(6) Persistent vomiting
(7) Known or suspected liver disease (including patients suspected of current alcohol
ingestion)
(8) Allergy to acetaminophen
(9) Patients less than 3 months of age

e) Preparations Use Unit Dose Only
(Do NOT USE MULTIDOSE BOTTLE OF LIQUID)
Unit dose 160 mg/5 mL liquid
Unit dose 325 mg pill or tablet

f) Dosage
(1) Less than 3 months of age: Not indicated
(2) 3 months to 2 years of age:

<table>
<thead>
<tr>
<th>Age</th>
<th>Under 3 months</th>
<th>3 months</th>
<th>4-11 months</th>
<th>12-23 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liquid 160 mg/5 mL</td>
<td>Not indicated</td>
<td>1.25 mL</td>
<td>2.5 mL</td>
<td>3.75 mL</td>
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</table>

(3) 2–4 years: Unit dose 160 mg/5 mL
(4) 5–12 years: TWO unit doses of 160 mg/5 mL each for a total of 320 mg/10 mL
(5) 13 years and above: FOUR unit doses of 160 mg/5 mL each for a total of 640 mg/20 mL OR in a form of 325 mg pill or tablet x2 for a total of 650 mg with sips of
water as tolerated by the patient.
TRADE NAMES: Not Applicable

a) Indications
   Poisoning by mouth

b) Adverse Effects
   May indirectly induce vomiting and cause nausea

c) Precautions
   Does not adsorb all drugs and toxic substances

d) Contraindications
   (1) Altered mental status
   (2) Patients who have received an emetic

e) Preparations
   (1) 25 grams/125 mL bottle
   (2) 50 grams/250 mL bottle

f) Dosage
   (1) Adult: Administer 1 gram/kg PO
   (2) Pediatric: Administer 1 gram/kg PO

POISON INFORMATION CENTER RECOMMENDATIONS SHOULD BE SOLICITED IN CONJUNCTION WITH MEDICAL CONSULTATION, BUT MEDICATION ORDERS CAN ONLY BE ACCEPTED FROM AN APPROVED BASE STATION OR CONSULTATION CENTER.
TRADE NAMES: PROVENTIL®, VENTOLIN®

a) Indications
   (1) Signs and symptoms of respiratory distress
   (2) Bronchospasm/wheezing associated with:
       (a) Asthma
       (b) COPD/emphysema
       (c) Allergic reactions (anaphylaxis)

b) Adverse Effects
   (1) Tachycardia/palpitations
   (2) Hypertension
   (3) Angina
   (4) Nervousness/anxiety
   (5) Tremors
   (6) Dizziness
   (7) Headache
   (8) Sweating
   (9) Nausea/vomiting
   (10) Sore throat

c) Precautions
    May cause severe bronchospasm from repeated excessive use.

d) Contraindications
    Known hypersensitivity

e) Preparations
   (1) Hand-held (unit dose) aerosol inhaler
   (2) Ampule for nebulizer

f) Dosage
   Inhaler
   (1) Adult: Patient may receive a maximum of 2 doses (4 puffs) over a 30-minute period
   (2) Pediatric: Patient may receive a maximum of 2 doses (4 puffs) over a 30-minute period
   Nebulizer
   (1) Adult: 2.5 mg by nebulized aerosol connected to 6–8 lpm of oxygen; may repeat one time
   (2) Pediatric: May repeat one time; connect to 6–8 lpm of oxygen
      (a) Age 2 or older: 2.5 mg by nebulized aerosol
      (b) Age less than 2 years: 1.25 mg by nebulized aerosol
   (3) Additional doses may be administered with medical consultation.
TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Platelet inhibitor
   (2) Anti-inflammatory

b) Pharmacokinetics
   Blocks platelet aggregation

c) Indications
   Suspected Acute Coronary Syndrome and/or ST Elevation MI (STEMI)

d) Contraindications
   (1) Known hypersensitivity.
   (2) Patients who receive a full dose (324 mg) of aspirin prior to EMS arrival.

e) Adverse Effects
   (1) Heartburn
   (2) Nausea and vomiting
   (3) Wheezing

f) Precautions
   GI bleeding and upset

g) Dosage
   (1) Adult: 324 mg or 325 mg chewed
   (2) Pediatric: Not indicated
TRADE NAMES: Not Applicable

a) Indications
   (1) Moderate to severe allergic reaction with respiratory distress or mild allergic reaction with history of life-threatening allergic reaction
   (2) Patients with severe asthma

b) Adverse Effects
   (1) Tachycardia/palpitations
   (2) Angina
   (3) Headache
   (4) Nausea/vomiting
   (5) Dizziness
   (6) Hypertension
   (7) Nervousness/Anxiety
   (8) Tremors

c) Precautions
   Medical consultation must be obtained before administering the EMS service’s manual epinephrine or EMS service’s auto-injector to cardiac (pediatric and adult), pregnant, and adult asthma patients. However, medical consultation is not required for severe allergic reactions with respiratory distress.

d) Contraindications
   None in the presence of anaphylaxis

e) Preparations
   Epinephrine
   (Patient prescribed or EMS supplied)
   (1) Vial: 1 mg in 1 mL
   (2) Preloaded Syringe
      (a) Adult: 0.5 mg in 0.5 mL
      (b) Pediatric: 0.15 mg in 0.15 mL

f) Dosage
   (1) Patients 5 years of age or greater:
      Adult: 0.5 mg in 0.5 mL IM in lateral thigh
   (2) Patients less than 5 years of age:
      Pediatric: 0.15 mg in 0.15 mL IM in lateral thigh
   (3) Additional doses may be administered with medical consultation.
TRADE NAMES: Not Applicable

a) Indications
   (1) Moderate to severe allergic reaction with respiratory distress or mild allergic reaction with history of life-threatening allergic reaction
   (2) Patients with severe asthma

b) Adverse Effects
   (1) Tachycardia/palpitations
   (2) Angina
   (3) Headache
   (4) Nausea/vomiting
   (5) Dizziness
   (6) Hypertension
   (7) Nervousness/anxiety
   (8) Tremors

c) Precautions
   Medical consultation must be obtained before administering the EMS service’s manual epinephrine or EMS service’s auto-injector to asthma patients with pregnancy or cardiac history. However, medical consultation is not required for any patients who have severe allergic reactions with respiratory distress.

d) Contraindications
   None in the presence of anaphylaxis

e) Preparations
   Epinephrine Auto-injector (single or multi-dose) only
   (Patient prescribed or EMS supplied)
   (1) Adult: 0.3 mg
   (2) Pediatric: 0.15 mg

f) Dosage
   (1) Less than 5 years of age: 0.15 mg IM in the lateral thigh via epinephrine auto-injector.
   (2) 5 years and greater: administer 0.3 mg IM in the lateral thigh via epinephrine auto-injector.
   (3) Additional doses may be administered with medical consultation.
TRADE NAMES: Narcan®

a) Pharmacology
   Reverses all effects due to opioid (morphine-like) agents. This drug will reverse the respiratory depression and all central and peripheral nervous system effects.

b) Pharmacokinetics
   (1) Onset of action is within a few minutes with intranasal (IN) administration.
   (2) Patients responding to naloxone may require additional doses and transportation to the hospital since most opioids/narcotics last longer than naloxone.
   (3) Has no effect in the absence of opioid/narcotic.

c) Indications
   To reverse respiratory depression induced by opioid/narcotic agent.

d) Contraindications
   Patients under 28 days of age

e) Adverse Effects
   Opioid withdrawal

f) Precautions
   (1) Naloxone may induce opiate withdrawal in patients who are physically dependent on opioids.
   (2) Certain drugs may require much higher doses of naloxone for reversal than are currently used.
   (3) Should be administered and titrated so respiratory efforts return, but not intended to restore full consciousness.
   (4) Intranasal naloxone must be administered via nasal atomizer.
   (5) Naloxone has a duration of action of 40 minutes; the effect of the opioid/narcotic may last longer than naloxone and patients should be encouraged to be transported.

CLINICIANS MUST CONTACT A BASE STATION PHYSICIAN FOR PATIENTS WISHING TO REFUSE TRANSPORT AFTER BLS ADMINISTRATION OF NALOXONE.

g) Dosage
   (1) Adult: Administer 2 mg IN, dividing administration of the dose equally between the nares to a maximum of 1 mL per nare, OR administer 4 mg/0.1 mL IN in one nare.
   (2) Pediatric (child aged 28 days to adult): Administer 2 mg IN, dividing administration of the dose equally between the nares to a maximum of 1 mL per nare, OR administer 4 mg/0.1 mL IN in one nare.
   (3) Repeat as necessary to maintain respiratory activity.
TRADE NAMES: Not Applicable
(Patient Prescribed, Patient Assisted)

a) Indications
Chest pain

b) Adverse Effects
(1) Hypotension
(2) Headache
(3) Dizziness
(4) Tachycardia

c) Precautions
(1) BLS clinician may only administer patient prescribed sublingual nitroglycerin.
(2) Reassess blood pressure before and after administration.
(3) If systolic blood pressure drops more than 20 mmHg per dose of nitroglycerin given, obtain medical consultation before further administration.

d) Contraindications
(1) Blood pressure below 90 mmHg systolic
(2) Heart rate less than 60 or greater than 150 bpm
(3) Medication not prescribed for the patient
(4) Pediatric patient under age 13
(5) Any patient having taken medication for Pulmonary Artery Hypertension (e.g., Adcirca® or Revatio®) or erectile dysfunction (e.g., Viagra®, Levitra®, or Cialis®) within the past 48 hours. Medical consultation is required to override this contraindication.

e) Preparations
Spray or tablet

f) Dosage
(1) Adult: One tablet or one spray sublingually
   (a) Repeat in 3 to 5 minutes if chest pain persists.
   (b) Maximum of three doses (a combination of patient-administered and EMT-administered) of nitroglycerin

(2) Pediatric: (nitroglycerin contraindicated for children under age 13)

(3) Additional doses may be administered with medical consultation.
TRADE NAMES: Not Applicable

a) Indications
   (1) Altered mental status with known diabetic history
   (2) Unconscious for an unknown reason
   (3) Measured blood glucose less than 70 mg/dL

b) Adverse Effects
   Not clinically significant

c) Precautions
   Patient without gag reflex may aspirate.

d) Contraindications
   Not clinically significant

e) Preparations
   10–15 grams of glucose (contained in 24, 30, or 37.5 gram tube)

f) Dosage
   (1) Adult: Administer 10–15 grams of oral glucose between the gum and cheek. Consider single additional dose of oral glucose if not improved after 10 minutes.
   (2) Pediatric: Administer 10–15 grams of oral glucose between the gum and cheek; this may be accomplished through several small administrations. Consider single additional dose of oral glucose if not improved after 10 minutes.
BLS Pharmacology – OXYGEN

TRADE NAMES: Not Applicable

a) Pharmacology
(1) Increases oxygen content of the blood
(2) Improves tissue oxygenation
(3) Decreases energy expended for respirations

b) Pharmacokinetics
Changing the percentage of inspired oxygen results in an increased blood and tissue level equilibration within 5–20 minutes.

c) Indications
(1) If evidence of hypoxia (Less than 94% SpO₂)
(2) Respiratory distress
(3) Cardiopulmonary arrest
(4) Trauma
(5) Suspected CO exposure
(6) Dyspnea

d) Contraindications
Not clinically significant

e) Adverse Effects
High concentrations of oxygen will reduce the respiratory drive in some COPD patients; these patients should be carefully monitored.

f) Precautions
(1) Never withhold oxygen from those who need it.
(2) Oxygen should be given with caution to patients with COPD.
(3) Simple or partial rebreather face masks must be supplied with a minimum 6 lpm.
(4) Non-breather (NRB) face masks must be supplied with a minimum 12 lpm.

g) Dosage
(1) Adult: Administer 12–15 lpm via NRB mask or 2–6 lpm via nasal cannula, as needed.
   CO exposure: Administer 100% oxygen via NRB mask. Maintain SpO₂ at 100%
(2) Pediatric: Administer 12–15 lpm via NRB mask or 2-6 lpm via nasal cannula, as needed.
   CO exposure: Administer 100% oxygen via NRB mask. Maintain SpO₂ at 100%

<table>
<thead>
<tr>
<th>Percent O₂ Saturation</th>
<th>Ranges</th>
<th>General Patient Care</th>
</tr>
</thead>
<tbody>
<tr>
<td>94–100%</td>
<td>Normal</td>
<td>Give oxygen as necessary</td>
</tr>
<tr>
<td>91–93%</td>
<td>Mild Hypoxia</td>
<td>Give oxygen as necessary</td>
</tr>
<tr>
<td>86–90%</td>
<td>Moderate Hypoxia</td>
<td>Give 100% oxygen</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Assisting Ventilations if necessary</td>
</tr>
<tr>
<td>less than or equal to</td>
<td>Severe Hypoxia</td>
<td>Give 100% oxygen</td>
</tr>
<tr>
<td>85%</td>
<td></td>
<td>Assist Ventilations</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If indicated, Intubate</td>
</tr>
</tbody>
</table>

**INACCURATE OR MISLEADING SpO₂ READINGS MAY OCCUR IN THE FOLLOWING PATIENTS: HYPOTHERMIC, HYPOPERFUSION (SHOCK), CO POISONING, HEMOGLOBIN ABNORMALITY, ANEMIA, AND VASOCONSTRICTION.**
TRADE NAMES: Tylenol®

a) Indications
Patients 3 months of age and older with:
(1) Mild to moderate discomfort (e.g., 1–5 on FACES scale) or
(2) Fever (EMS-documented temperature greater than or equal to 100.4 F / 38 C)

b) Adverse Effects
Not clinically significant

c) Precautions
(1) Administration of acetaminophen for mild to moderate pain does not eliminate
the need for transport of the patient to the hospital to receive a comprehensive
evaluation of the cause of the pain and appropriate definitive treatment.
(2) A 5 cc or 6 cc syringe must be used to measure doses of acetaminophen.

d) Contraindications
(1) Head Injury
(2) Hypotension
(3) Administration of acetaminophen or medications containing acetaminophen within
the previous 4 hours. Many common cold preparations contain acetaminophen.
(4) Inability to swallow or take medications by mouth
(5) Respiratory distress
(6) Persistent vomiting
(7) Known or suspected liver disease (including patients suspected of current alcohol
ingestion)
(8) Allergy to acetaminophen
(9) Patients less than 3 months of age

e) Preparations Use Unit Dose Only
(Do NOT USE MULTIDOSE BOTTLE OF LIQUID)
Unit dose 160 mg/5 mL liquid
Unit dose 325 mg pill or tablet

f) Dosage
(1) Less than 3 months of age: Not indicated
(2) 3 months to 2 years of age:

<table>
<thead>
<tr>
<th>Age</th>
<th>Under 3 months</th>
<th>3 months</th>
<th>4-11 months</th>
<th>12-23 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liquid 160 mg/ 5 mL</td>
<td>Not indicated</td>
<td>1.25 mL</td>
<td>2.5 mL</td>
<td>3.75 mL</td>
</tr>
</tbody>
</table>

(3) 2–4 years: Unit dose 160 mg/5 mL
(4) 5–12 years: TWO unit doses of 160 mg/5 mL each for a total of 320 mg/10 mL
(5) 13 years and above: FOUR unit doses of 160 mg/5 mL each for a total of 640 mg/
20 mL OR in a form of 325 mg pill or tablet x2 for a total of 650 mg with sips of
water as tolerated by the patient.
ALS Pharmacology –
ACTIVATED CHARCOAL (WITHOUT SORBITOL)

TRADE NAMES: Not Applicable

a) Pharmacology
Variable drug or toxin absorption when ingested

b) Pharmacokinetics
Adsorbs poisons and prevents toxins from entering body systems

c) Indications
Poisoning by mouth

d) Contraindications
(1) Altered mental status
(2) Patients who have received an emetic

e) Adverse Effects
Not clinically significant

f) Precautions
Does not adsorb all drugs and/or toxic substances

g) Dose
(1) Adult: Administer 1 gram/kg PO
(2) Pediatric: Administer 1 gram/kg PO

POISON INFORMATION CENTER RECOMMENDATIONS SHOULD BE SOLICITED IN CONJUNCTION WITH MEDICAL CONSULTATION, BUT MEDICATION ORDERS CAN ONLY BE ACCEPTED FROM AN APPROVED BASE STATION OR CONSULTATION CENTER.
TRADE NAMES: ADENOCARD®

a) Pharmacology
   (1) Naturally occurring purine nucleoside
   (2) Used to treat narrow complex tachycardia, PSVT with WPW
   (3) Slows conduction through the AV node
   (4) No effect on ventricular contractility
   (5) Causes peripheral vasodilatation (often dramatic)

b) Pharmacokinetics
   Onset of action within 5–20 seconds following an IV dose; half-life is 10 seconds

c) Indications
   (1) To slow the rate of narrow complex tachycardia
   (2) Is only effective on SVT/PSVT
   (3) No effect on VT, atrial fibrillation, or flutter
   (4) In stable, wide complex tachycardia (possible VT) for pediatric with caution

d) Contraindications
   (1) Known hypersensitivity
   (2) History of moderate to severe asthma or active bronchospasm
   (3) Polymorphic or irregular wide complex tachycardia

e) Adverse Effects
   Flushing, dyspnea, chest pressure, nausea, headache, dizziness, and hypotension

f) Precautions
   (1) Effects antagonized by theophylline
   (2) Effects enhanced by dipyridamole (Persantine®), digitalis, carbamazepine, calcium channel blockers, and benzodiazepines
   (3) Be prepared for up to 40 seconds of asystole.

g) Dosage
   (1) Adult:
      6 mg rapid IVP bolus followed by a rapid flush
      Give 12 mg if no response within 2 minutes.
      Give 12 mg more if no response within another 1–2 minutes.
      REDUCE DOSAGE BY HALF FOR PATIENTS WITH TRANSPLANTED HEARTS AND THOSE TAKING DIPYRIDAMOLE OR CARBAMAZEPINE.
   (2) Pediatric: 0.1 mg/kg rapid IVP/IO; maximum initial dose 6 mg
      Second and third doses: 0.2 mg/kg rapid IVP/IO; maximum single additional dose 12 mg
11.4 ALS Pharmacology – ALBUTEROL

TRADE NAMES: PROVENTIL®, VENTOLIN®

a) Pharmacology
Stimulates beta-2 adrenergic receptors of the bronchioles; bronchodilator

b) Pharmacokinetics
(1) Bronchodilation begins within 5–15 minutes after inhalation.
(2) Peak effect occurs in 30–120 minutes.
(3) Duration of action is usually 3–4 hours.

c) Indications
(1) Signs and symptoms of respiratory distress
(2) Bronchospasm/wheezing associated with:
   (a) Asthma
   (b) COPD/emphysema
   (c) Allergic reaction (anaphylaxis)
(3) Hyperkalemia

d) Contraindications
Known hypersensitivity

e) Adverse Effects
Tachycardia, palpitations, peripheral vasodilation, tremors, nervousness, headache, sore throat, PVCs, nausea, and vomiting

f) Precautions
May cause severe bronchospasm from repeated excessive use

g) Dosage
(1) Bronchospasm
   Inhaler
   (a) Adult: Patient may receive a maximum of 2 doses (4 puffs) over a 30-minute period
   (b) Pediatric: Patient may receive a maximum of 2 doses (4 puffs) over a 30-minute period
   Nebulizer
   (a) Adult: 2.5 mg by nebulized aerosol connected to 6–8 lpm of oxygen; may repeat one time
   (b) Pediatric: May repeat one time; connect to 6–8 lpm of oxygen
      (i) Age 2 or older: 2.5 mg by nebulized aerosol
      (ii) Age less than 2 years: 1.25 mg by nebulized aerosol
(2) Hyperkalemia
   (a) Adult: 20 mg (if available) by nebulized aerosol connected to 6–8 lpm of oxygen
   (b) Pediatric
      (i) **Age 2 or older**: 2.5 mg by nebulized aerosol
      (ii) **Age less than 2 years**: 1.25 mg by nebulized aerosol
TRADE NAMES: Not Applicable

a) Pharmacology
Prolongs duration and refractory period of action potential. Slows electrical conduction, electrical impulse generation from sinoatrial node, and conduction through accessory pathways. Also dilates blood vessels.

b) Pharmacokinetics
Amiodarone primarily alters/blocks the potassium and sodium ion permeability across the myocardial membrane, which in effect, stabilizes the ion channels and changes impulse conduction through the myocardium. Amiodarone also has some effects on beta receptors and calcium channels.

c) Indications
(1) Prevent recurrence of ventricular fibrillation/tachycardia after defibrillation and conversion to supraventricular rhythm
(2) Ventricular tachycardia (VT)
(3) Ventricular fibrillation (VF)

d) Contraindications
(1) Second or third degree AV blocks
(2) Sensitivity to amiodarone
(3) Idioventricular escape rhythms
(4) Accelerated idioventricular rhythm
(5) Sinus bradycardia or arrest or block
(6) Hypotension
(7) Cardiogenic shock
(8) Ventricular conduction defects
(9) Iodine hypersensitivity

e) Adverse Effects
(1) Bradycardia
(2) Hypotension
(3) Prolonged QT interval

f) Precautions
May prolong the QT interval increasing risk of torsades de pointes, and VF. Amiodarone inhibits atrioventricular conduction and decreases myocardial contractility, increasing the risk of AV block or of hypotension with any calcium channel blocker.
g) Dosing

(1) Adult with pulse: 150 mg IV/IO over 10 minutes (mixed in 50 - 100 mL of approved diluent). May repeat once.

(2) Adult without pulse VF/VT/(torsades after magnesium sulfate): 300 mg IV/IO. May repeat one time

(3) Pediatric with pulse: 5 mg/kg IV/IO over 20 minutes (mixed in 50 - 100 mL of approved diluent)

(4) Pediatric without pulse: 5 mg/kg IV/IO; max single dose 300 mg. May repeat twice to a maximum of 15 mg/kg.
TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Platelet inhibitor
   (2) Anti-inflammatory

b) Pharmacokinetics
   Blocks platelet aggregation

c) Indications
   (1) Suspected Acute Coronary Syndrome and/or
   (2) ST Elevation MI (STEMI)

d) Contraindications
   (1) Known hypersensitivity.
   (2) Patients who receive a full dose (324 mg) of aspirin prior to EMS arrival.

e) Adverse Effects
   (1) Heartburn
   (2) Nausea and vomiting
   (3) Wheezing

f) Precautions
   GI bleeding and upset

g) Dosage
   (1) Adult: 324 mg or 325 mg chewed
   (2) Pediatric: Not indicated
TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Parasympatholytic (vagolytic action)
   (2) Anticholinergic (accelerates the heart rate)

b) Pharmacokinetics
   (1) Accelerated heart rate within minutes of IV injection.
   (2) Peak effect is seen within the first 15 minutes.
   (3) Atropine disappears rapidly from the blood.
   (4) Excreted in the urine within the first 12 hours.

c) Indications
   (1) Symptomatic bradycardia
   (2) Organophosphate poisoning
   (3) Nerve agents

d) Contraindications
   (1) Known hypersensitivity
   (2) Dysrhythmias in which enhancement of conduction may accelerate the ventricular rate and cause decreased cardiac output (e.g., atrial fibrillation, atrial flutter, or PAT with block)
   (3) Relative Contraindications (weigh risk/benefits):
      (a) AV block at His-Purkinje level (second-degree Type II AV Block and third-degree AV Block)
      (b) Suspected acute myocardial infarction or ischemia
      (c) Glaucoma

e) Adverse Effects
   (1) Excessive doses of atropine can cause delirium, restlessness, disorientation, tachycardia, coma, flushed and hot skin, ataxia, blurred vision, dry mucous membranes.
   (2) Ventricular fibrillation and tachycardia have occurred following IV administration of atropine.

f) Precautions
   Not clinically significant
g) Dosage
(1) Adult:
  Bradycardia: Administer 0.5–1 mg IVP repeated every 3–5 minutes to a total dose of 0.04 mg/kg
(2) Pediatric:
  Bradycardia: Administer 0.02 mg/kg IV/IO; maximum single dose 0.5 mg; ET 0.04–0.06 mg/kg, dilute 5 mL; repeat once
(3) **Organophosphate poisoning:**
  (a) Adult: Administer 2–4 mg IVP or IM every 5–10 minutes.
  (b) Pediatric: Administer 0.02 mg/kg IVP/IO or IM every 5–10 minutes.
(4) Nerve agent exposure
  See *MARK I / DuoDote* protocol.
ALS Pharmacology –
CALCIUM CHLORIDE (10% SOLUTION)

TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Increase cardiac contractile state and ventricular automaticity
   (2) Is useful in reversing cardiac arrhythmias due to hyperkalemia (often seen in renal dialysis patients)

b) Pharmacokinetics
   Rapid onset of action with IV administration

c) Indications
   (1) Hyperkalemia
   (2) Hypocalcemia
   (3) To treat adverse effects caused by calcium channel blocker overdose
   (4) Hypotension secondary to diltiazem administration
   (5) Respiratory depression, decreased reflexes, flaccid paralysis, and apnea following magnesium sulfate administration

d) Contraindications
   (1) Not indicated in cardiac arrest except when hyperkalemia, hypocalcemia, or calcium channel toxicity is highly suspected
   (2) Patient currently taking digoxin with suspected calcium channel blocker overdose

e) Adverse Effects
   (1) Bradycardia may occur with rapid injection.
   (2) Syncope, cardiac arrest, arrhythmia, bradycardia

f) Precautions
   (1) Use with caution on patients taking digitalis, as calcium may increase ventricular irritability and precipitate digitalis toxicity.
   (2) If given with sodium bicarbonate, calcium will precipitate.
   (3) Calcium salts may produce coronary and cerebral artery spasm.

g) Dosage
   (1) Adult: Administer 0.5–1 gram SLOW IVP over 10 minutes.
       Maximum dose 1 gram
       Administer 500 mg SLOW IVP for: hypotension following diltiazem administration.
       Respiratory depression, decreased reflexes, flaccid paralysis, and apnea following magnesium sulfate administration
   (2) Pediatric: Administer 20 mg/kg (0.2 mL/kg) SLOW IVP/IO (50 mg/min)
       Maximum dose 1 gram
Trade Names: Decadron®

a) Indications
   (1) Moderate to severe asthma/COPD exacerbation
   (2) Croup
   (3) Anaphylaxis

b) Adverse Effects
   (1) Headache
   (2) Edema
   (3) Vertigo
   (4) Fluid retention
   (5) Adrenal insufficiency and immunosuppression with long-term use
   (6) HTN
   (7) CHF
   (8) Nausea and vomiting
   (9) Dyspepsia
   (10) Anaphylaxis

c) Precautions
   (1) Caution with diabetes
   (2) Known TB
   (3) Osteoporosis
   (4) Hepatic impairment
   (5) CHF
   (6) Seizure disorder

d) Contraindications
   (1) Hypersensitivity to drug
   (2) Known systemic fungal infection
   (3) Premature infants

e) Dosage (IV solution used for PO administration)
   (1) Adult: 10 mg IV (preferred, if established) or PO
   (2) Pediatric:
      (a) Asthma: 0.5 mg/kg PO (preferred) or IV to a maximum of 10 mg
      (b) Croup: 0.5 mg/kg PO/IM/IV to a maximum of 10 mg
ALS Pharmacology – DEXTROSE

TRADE NAMES: Not Applicable

a) Pharmacology
   Dextrose is a water-soluble monosaccharide found in corn syrup and honey.

b) Pharmacokinetics
   (1) Dextrose restores circulating blood sugar and is rapidly utilized following IV injection.
   (2) Excess dextrose is rapidly excreted unchanged in the urine.

c) Indications
   Correction of altered mental status due to low blood sugar (hypoglycemia) seizures and cardiac arrest

d) Contraindications
   Known hyperglycemia

e) Adverse Effects
   May worsen hyperglycemia (high blood sugar)

f) Precautions
   (1) May worsen preexisting hyperglycemia
   (2) Tissue necrosis if extravasation occurs

g) Dosage
   (1) Adult:
      (a) If blood glucose is less than 70 mg/dL, administer 10% dextrose in 50 mL (5 grams) boluses, one minute apart, to a maximum of 250 mL OR 25 grams of 50% dextrose IVP, until:
         (i) the patient has a return to normal mental status, and
         (ii) the patient’s blood glucose is at least 90 mg/dL.
         (iii) If, following 250 mL of 10% dextrose or 25 grams of 50% dextrose, patient has persistently altered mental status and blood glucose less than 90 mg/dL, repeat dosing regimen in (a).
   (2) Pediatric:
      (a) Patient less than 28 days - if blood glucose is less than 40 mg/dL administer 2 mL/kg of 10% dextrose IV/IO.
         D10W is prepared by mixing one part of D50W with four parts LR.
         Recheck glucose after first dose.
         If blood glucose is less than 40 mg/dL, obtain medical consultation to administer second dose of D10W.
      (b) Patients 28 days up to 4 years - if blood glucose is less than 70 mg/dL, administer 2–4 mL/kg of 10% dextrose IV/IO to a maximum of 25 grams. Recheck glucose after first dose.
         If blood glucose is less than 70 mg/dL, obtain medical consultation to administer second dose of D10W.
         (i) If unable to start IV and blood glucose is less than 70 mg/dL, administer 0.5 mg glucagon IM/IN.
         (ii) Medical consultation for additional dosing to a maximum of 3 mg IM/IN.
(c) **Patients 5 years up to patient’s 18th birthday** - if blood glucose is less than 70 mg/dL, administer 2–4 mL/kg of 10% dextrose IV/IO to a maximum of 25 grams. 
Recheck glucose after first dose.

(i) If blood glucose is less than 70 mg/dL, obtain medical consultation to administer second dose of D10W.

(ii) If unable to start IV and blood glucose is less than 70 mg/dL, administer 1 mg glucagon IM/IN.

(ii) Medical consultation for additional dosing to a maximum of 3 mg IM/IN
ALS Pharmacology – DIAZEPAM

TRADE NAMES: VALIUM®

a) Pharmacology
   (1) Sedation, hypnosis, alleviation of anxiety, muscle relaxation, anticonvulsant activity
   (2) Little cardiovascular effect

b) Pharmacokinetics
   (1) Onset of action is extremely rapid following IV administration.
   (2) Half-life ranges from 20–90 minutes.

c) Indications
   (1) Sustained and/or recurrent seizures
   (2) Severe nerve agent exposure

d) Contraindications
   (1) Known hypersensitivity, head injury
   (2) Should be used with caution in patients with altered mental status, hypotension, or acute narrow angle glaucoma

e) Adverse Effects
   (1) Lightheadedness, motor impairment, ataxia, impairment of mental and psychomotor function, confusion, slurred speech, amnesia
   (2) Additive effect with ethanol
   (3) Irritability and excitation may be seen paradoxically.

f) Precautions
   (1) Respiratory depression may occur with IV administration, especially if given too rapidly.
   (2) Respiratory support may be required.
   (3) Use with caution in pregnant patients, persons ingesting alcohol, or persons ingesting sedatives.

g) Dosage (paramedic may perform without consultation for patients with active seizures if Midazolam is not available.)
   (1) Adult: Administer 2.5–10 mg in 2.5 mg increments SLOW IVP/IM
      (IM requires all clinicians to obtain medical consultation.)
      Maximum total dose 10 mg
   (2) Pediatric: Administer 0.1 mg/kg in 2.5 mg increments SLOW IVP/IO/IM (IM requires all clinicians to obtain medical consultation.)
      Maximum total dose 5 mg
      Rectal Dose: Administer up to 0.2 mg/kg; maximum total dose 10 mg

Severe nerve agent exposure (clinicians may administer without consultation):
(3) Adult: Administer 10 mg IM.
(4) Pediatric: greater than 30 kg: Administer 10 mg via auto-injector or 0.1 mg/kg IM, maximum of 10 mg.
ALS Pharmacology –  
DILTIAZEM

TRADE NAMES: CARDIZEM®

a) Class
Calcium channel blocker

b) Actions
(1) Inhibits the movement of calcium ions across cardiac muscle cells
(2) Decreases conduction velocity and ventricular rate

c) Indications
Symptomatic atrial fibrillation and atrial flutter

d) Contraindications
(1) Hypotension below 100 mmHg, second or third degree heart block, hypersensitivity to the drug
(2) Patients less than 18 years of age

e) Precautions
Use cautiously in patients with renal failure or congestive heart failure.

f) Side effects
(1) Headache
(2) Nausea
(3) Vomiting
(4) Bradycardia
(5) Hypotension

g) Significant interactions
Congestive heart failure may result if used along with beta blockers.

h) Dosage
(1) Adult
(a) 0.25 mg/kg (maximum dose 20 mg) by IV bolus administered SLOW IV over 2 minutes; if response is not adequate, repeat in 15 minutes with a dosage of 0.35 mg/kg (maximum dose 25 mg) over 2 minutes.
(b) For patients older than 50 years of age, borderline blood pressure, known renal failure, or CHF, consider initial bolus 5–10 mg administered IV over 2 minutes.

(2) Pediatric:
Contraindicated for patients less than 18 years of age. If needed, consult Pediatric Base Station.

i) Overdose or Toxicity Presentation
Generally consists of exaggeration of side effects, including severe hypotension and symptomatic bradycardia
j) **Treatment of Overdose or Other Adverse Reactions**

(1) Give general supportive measures, monitor vitals, administer oxygen.

(2) Hypotension:
   (a) If lungs are clear, administer fluid bolus 20 mL/kg of LR; titrate to a systolic blood pressure of 100 mmHg.
   (b) If rales are present, administer fluid bolus, maximum of 250 mL of LR. Titrate to a systolic of 100 mmHg.
   (c) Administer calcium chloride 500 mg SLOW IVP.

(3) Bradycardia: Consider atropine (0.5 to 1 mg); if necessary, consider pacing.
ALS Pharmacology –
DIPHENHYDRAMINE HYDROCHLORIDE

TRADE NAMES: BENADRYL®

a) Pharmacology
   Antihistamine

b) Pharmacokinetics
   (1) Effect begins within 15 minutes of IV dose.
   (2) Peak effect 1–4 hours
   (3) Metabolized by the liver
   (4) The half-life ranges from 2–10 hours.

c) Indications
   (1) Allergic reaction
   (2) Anaphylaxis
   (3) Dystonic reactions

d) Contraindications
   Known allergy to diphenhydramine

e) Adverse Effects
   Drowsiness, loss of coordination, blurred vision, headache, hypotension, tachycardia, palpitations, thickening of bronchial secretions leading to chest tightness, and wheezing

f) Precautions - Should be used with caution in patients with:
   (1) Severe vomiting
   (2) Alcohol intoxication
   (3) Nursing mothers

g) Dosage
   (1) Adult: Administer 25–50 mg SLOW IVP or IM
   (2) Pediatric: Administer 1 mg/kg SLOW IV or IM
a) Pharmacology
   (1) Alpha and beta adrenergic receptor stimulator
   (2) Dopaminergic receptor stimulator
   (3) Precursor of norepinephrine
   (4) At low doses, less than 2 mcg/kg/min
      (a) Dilates renal and mesenteric blood vessels
      (b) Venoconstricts
      (c) Arterial resistance varies
   (5) At moderate doses, 2–6 mcg/kg/min beta1 stimulating effect on heart
      Results in increased cardiac output
   (6) High dose, 6–10 mcg/kg/min
      Exhibits alpha1 effects; peripheral vasoconstriction including renal and mesenteric vessels, increases left and right ventricular preload
   (7) Doses greater than or equal to 10 mcg/kg/min
      Alpha1 stimulating effects may reverse mesenteric and renal artery dilatation resulting in decreased blood flow, causing increased preload due to effects on venous system

b) Pharmacokinetics
   (1) Extremely rapid onset of action
   (2) Extremely brief duration of action
   (3) The rate of administration may be used to control the effect of dopamine.

c) Indications
   (1) Cardiogenic shock
   (2) Septic shock
   (3) Anaphylactic shock
   (4) Hypovolemic shock (after sufficient volume replacement)

d) Contraindications
   (1) Preexisting tachydysrhythmias
   (2) Uncorrected hypovolemia

e) Adverse Effects
   (1) Anginal pain
   (2) Tachydysrhythmias
   (3) Nausea and vomiting
   (4) Hypertension
   (5) Undesirable degree of vasoconstriction
f) **Precautions**
   
   (1) Extravasation should be reported to the hospital staff on arrival.
   
   (2) Patients receiving monoamine oxidase (MAO) inhibitors are extremely sensitive to the effects of dopamine and should receive a much lower dosage than is usually given.
   
   (3) Patients with pheochromocytoma are extremely sensitive to dopamine and may develop profound hypertension in response to minimal doses.

g) **Dosage**

   (1) For IV/IO infusion only. The preferred route of administration is IV.
   
   (2) In general, the infusion rate is adjusted to blood pressure and clinical response.
   
   (3) Adult: Administer 2–20 mcg/kg/min IV drip titrated to BP of 100 systolic or medical consultation selected BP; initial infusion rate 2–5 mcg/kg/min
   
   (4) Pediatric: Administer 2–20 mcg/kg/min IV drip titrated age specific BP or medical consultation selected BP; initial infusion rate is 2 mcg/kg/min
TRADE NAMES: INAPSINE®

a) Pharmacology
   Antipsychotic

b) Pharmacokinetics
   Onset of action is within 10 minutes of the IM administration.

c) Indications
   Moderate agitation (defined as behavior that puts the patient or clinician at risk of harm) due to suspected psychiatric emergency (e.g., schizophrenia) or medical delirium

d) Contraindications
   (1) Children under 13 years of age
   (2) Pregnancy
   (3) Parkinson’s disease
   (4) CNS depression or acute CNS injury
   (5) Severe agitation (see midazolam and ketamine)

e) Adverse Effects
   (1) Dystonic reaction
      (a) Rarely seen with short-term use
      (b) ADULT: administer a single dose of diphenhydramine 25–50 mg IV/IM
      (c) PEDIATRIC: Administer a single dose of diphenhydramine 1 mg/kg IV/IM (max of 25 mg)
   (2) Hypotension and tachycardia
      (a) Administer Lactated Ringer's fluid bolus if hypotension occurs
      (3) Torsades de pointes (polymorphic ventricular tachycardia)
         (a) Patients receiving droperidol should be monitored for cardiac dysrhythmias.

f) Precautions
   (1) Violent patients may require physical restraint while the medication is administered.
   (2) Patients receiving droperidol should be monitored for cardiac dysrhythmias.

g) Dosage
   (1) ADULT
      (a) Patient 18–68 years of age: 2.5 mg IM
      (b) Patient 69 years and older: 1.25 mg IM
      (c) Medical consultation required for additional dose
   (2) PEDIATRIC
      (a) Patient 13–18 years of age: 2.5 mg IM
      (b) Patient less than 13 years of age: contraindicated
      (c) Medical consultation required for additional dose
ALS Pharmacology –
EPINEPHRINE (0.1 mg/mL and 1 mg/mL)

TRADE NAMES: Not Applicable

a) Pharmacology
(1) The administration of epinephrine causes increases in:
   (a) Systemic vascular resistance
   (b) Systemic arterial pressure
   (c) Heart rate (positive chronotropic effect)
   (d) Contractile state (positive inotropic effect)
   (e) Myocardial oxygen requirement
   (f) Cardiac automaticity
   (g) AV conduction (positive dromotropic effect)
(2) Causes bronchial dilation by smooth muscle relaxation

b) Pharmacokinetics
(1) IV administered epinephrine has an extremely rapid onset of action.
(2) Is rapidly inactivated by the liver
(3) Subcutaneous administration of epinephrine results in slower absorption due to local vasoconstriction.
(4) Local massage will hasten absorption.
(5) Topically applied nebulizer within the respiratory tract, epinephrine has vasoconstrictor properties that result in reduction of mucosal and submucosal edema. It also has bronchodilator properties that reduce airway smooth muscle spasms.

c) Indications
(1) Medical cardiac arrest and pediatric traumatic arrest
(2) Moderate to severe allergic reaction/anaphylaxis
(3) IV push epinephrine should be reserved for cardiac arrest patients
(4) Epinephrine infusion (IV/IO) should be reserved for patients in shock refractory to fluid bolus or for patients in anaphylactic shock
(5) Severe asthma
(6) Respiratory stridor (suspected croup)

d) Contraindications
(1) Hypertension
(2) Preexisting tachydysrhythmias with a pulse (ventricular and supraventricular)
(3) IV push epinephrine should not be administered to any patient with a pulse

e) Adverse Effects
(1) Tachydysrhythmias (supraventricular and ventricular)
(2) Hypertension
(3) May induce early labor in pregnant women
(4) Headache
(5) Nervousness
(6) Decreased level of consciousness
(7) Rebound edema may occur 20–30 minutes after administration to croup patients.
ALS Pharmacology –
EPINEPHRINE (0.1 mg/mL and 1 mg/mL) (continued)

f) Precautions
(1) Do not mix with sodium bicarbonate as this deactivates epinephrine.
(2) Epinephrine causes a dramatic increase in myocardial oxygen consumption.
(3) Its use in the setting of an acute MI should be restricted to cardiac arrest.
(4) Medical consult for IM epinephrine in pregnant and cardiac patients.

g) Dosage
(1) Cardiac Arrest
   (a) Adult:
      (i) Administer epinephrine (0.1 mg/mL) 1 mg IVP/IO every 4 minutes to a maximum of 4 doses for the initial arrest. If arrest recurs following any period of ROSC, administer a maximum of 2 additional doses.
   (b) Pediatric:
      (i) Administer epinephrine (0.1 mg/mL) 0.01 mg/kg IVP/IO every 4 minutes to a maximum of 4 doses for the initial arrest. 0.01 mg/kg is equivalent to 0.1 mL/kg. If arrest recurs following any period of ROSC, administer a maximum of 2 additional doses.
      (ii) ET: Administer epinephrine (1 mg/mL) 0.1 mg/kg, diluted with 5 mL of LR; repeat every 4 minutes to a maximum of 4 doses for the initial arrest. If arrest recurs following any period of ROSC, administer a maximum of 2 additional doses.
   (c) Neonate:
      (i) Administer epinephrine (0.1 mg/mL) 0.01 mg/kg IVP/IO every 4 minutes to a maximum of 4 doses for the initial arrest. 0.01 mg/kg is equivalent to 0.1 mL/kg. If arrest recurs following any period of ROSC, administer a maximum of 2 additional doses.
      (ii) ET: Administer epinephrine (1 mg/mL) 0.1 mg/kg diluted with 5 mL of LR; repeat every 4 minutes to a maximum of 4 doses for the initial arrest. If arrest recurs following any period of ROSC, administer a maximum of 2 additional doses.

(2) Bradyarrhythmias
   (a) Adult: Using epinephrine infusion (1 mg epinephrine in 10 mL LR), administer 1 mL/min (60 drops/min) using a 60 drop-set. If systolic blood pressure remains less than 90 mmHg, obtain medical consultation for further dosing. Infusion pump: 2-10 mcg/min.
   (b) Pediatric:
      (i) Administer epinephrine (0.1 mg/mL) 0.01 mg/kg IVP/IO; repeat every 3–5 minutes. 0.01 mg/kg is equivalent to 0.1 mL/kg.
      (ii) ET: Administer epinephrine (1 mg/mL) 0.1 mg/kg, diluted with 5 mL of LR; repeat every 3–5 minutes.
   (c) Neonate:
      (i) Administer epinephrine (0.1 mg/mL) 0.01 mg/kg IVP/IO; repeat every 3–5 minutes. 0.01 mg/kg is equivalent to 0.1 mL/kg.
      (ii) ET: Administer epinephrine (0.1 mg/mL) 0.03 mg/kg, diluted with 1 mL of LR.
(3) Allergic Reaction/Anaphylaxis/Asthma/COPD
   (a) ADULT
      For patients who are in extremis with severe hypotension or impending respiratory failure:
      (i) Administer epinephrine (1 mg/mL) 0.5 mg IM every 5 minutes up to a total of 3 doses.
      (ii) If patient remains hypotensive or with impending respiratory failure, administer epinephrine infusion per management of shock with epinephrine infusion (below).
   (b) PEDIATRIC
      (i) Administer epinephrine
         Less than 5 years of age: administer epinephrine (1 mg/mL) 0.15 mg IM
         5 years and greater: administer epinephrine (1 mg/mL) 0.5 mg IM
      (ii) May repeat IM dose every 5 minutes for a total of 3 doses for severe reactions
      (iii) Epinephrine infusion (see management of non-traumatic shock with epinephrine infusion below)

(4) Croup
   (a) Adult: not indicated
   (b) Pediatric
      (i) Administer 2.5 mL of epinephrine 1 mg/mL via nebulizer.
         If patient does not improve, administer a second dose of 2.5 mL of epinephrine 1 mg/mL via nebulizer.
      (ii) Severe croup/imminent respiratory arrest: Administer 0.01 mg/kg of epinephrine (1 mg/mL) IM. Max single dose of 0.5 mg IM.
      (iii) Medical consult required for patients less than 1 year of age.

ALL PATIENTS WHO RECEIVE NEBULIZED EPINEPHRINE MUST BE TRANSPORTED BY AN ALS UNIT.

(5) Cardiogenic Shock
   (a) If rales are present, administer fluid bolus, titrate to a systolic blood pressure of 90 mmHg or greater. Maximum single of bolus of 250 mL of LR IV.
   (b) Additional fluid requires medical consultation.
   (c) Initiate epinephrine infusion
      (i) Add 1 mg of epinephrine (either 1 mg/mL or 0.1 mg/mL) in a 100 mL bag of LR or NS.
      (ii) Use a Microdrip set (60 drops/mL) for infusion administration.
      (iii) Adult epinephrine infusion dosage:
         (1) Administer infusion through a free-flowing IV, ideally 20 gauge or larger, or by IO
         (2) Start infusion at 1 mL/min (60 drops/min) IV/IO.
         (3) Check blood pressure every 5 minutes. If MAP is less than 65 mmHg or systolic blood pressure is less than 90 mmHg, increase to a maximum rate of 2 mL/min (120 drops/min).
(4) If above blood pressure goals are not met upon reaching maximum rate, obtain online medical consultation.

(6) Hypovolemic or Septic Shock
(a) If lungs are clear, administer fluid bolus of 20 mL/kg of LR IV. Titrate to a systolic blood pressure of 90 mmHg (or mean arterial pressure of 65 mmHg). Maximum patient dose of 2,000 mL of LR
(b) If hypotension persists after 2 L of LR are provided, consider additional LR up to a maximum of 30 mL/kg total.
(c) Initiate epinephrine infusion if systolic blood pressure remains less than 90 mmHg (or mean arterial pressure less than 65 mmHg) after IV fluid bolus of 30 mL/kg LR.

(7) Anaphylactic shock: Initiate epinephrine infusion for patients who are in extremis with severe hypotension or impending respiratory failure, after having administered 3 doses of IM epinephrine. (Refer to Anaphylaxis protocol.)

(8) Neurogenic shock (suspected spinal cord injury which typically presents with hypotension and bradycardia)
(a) If lungs are clear, administer fluid bolus of 20 mL/kg of LR IV. Titrate to a systolic blood pressure of 110 mmHg (or mean arterial pressure of 85 mmHg). Maximum patient dose of 2,000 mL of LR.
(b) Initiate epinephrine infusion if systolic blood pressure remains less than 110 mmHg (or mean arterial pressure less than 85 mmHg).

h) Pediatric epinephrine infusion dosage
(1) The following dosing chart should be used for pediatric patients less than 50 kg (using approved epinephrine infusion and 60 drop set):

<table>
<thead>
<tr>
<th>Weight range (kg)</th>
<th>Initial epinephrine dose</th>
<th>If goal blood pressure not achieved at 5 min, increase to</th>
</tr>
</thead>
<tbody>
<tr>
<td>LESS than 10 kg</td>
<td>6 drops/min (0.1 mL/min)</td>
<td>12 drops/min (0.2 mL/min)</td>
</tr>
<tr>
<td>10-19 kg</td>
<td>12 drops/min (0.2 mL/min)</td>
<td>24 drops/min (0.4 mL/min)</td>
</tr>
<tr>
<td>20-29 kg</td>
<td>18 drops/min (0.3 mL/min)</td>
<td>36 drops/min (0.6 mL/min)</td>
</tr>
<tr>
<td>30-39 kg</td>
<td>24 drops/min (0.4 mL/min)</td>
<td>48 drops/min (0.8 mL/min)</td>
</tr>
<tr>
<td>40-49 kg</td>
<td>30 drops/min (0.5 mL/min)</td>
<td>60 drops/min (1.0 mL/min)</td>
</tr>
</tbody>
</table>

(2) Blood pressure goal:
(a) For patients 10 years and older (including adults), systolic blood pressure greater than 90 mmHg;
(b) For patients under 10 years of age, systolic blood pressure greater than 70 + 2x age in years mmHg; OR
(c) Systolic blood pressure ordered by the pediatric base station.

(3) If above blood pressure goal not met after 10 minutes, obtain online medical consultation.
**Pharmacology**
(1) Synthetic opioid binds with opiate receptors in the CNS, altering both perception and emotional response to pain.
(2) Fentanyl is significantly more potent than morphine. 100 mcg of fentanyl is equivalent to 10 mg of morphine.

a) **Pharmacokinetics**
Onset of action is 2–3 minutes after IV dose and effects last 30 minutes to 1 hour.

b) **Indications**
(1) The patient reports moderate to severe pain.
(2) In the clinician’s judgment the patient will benefit from treatment with an opioid analgesic, including patients who are MOLST and/or EMS/DNR patients or being pre-medicating for a procedure.

c) **Contraindications**
(1) Hypersensitivity or known allergy to fentanyl
(2) Uncorrected respiratory distress or hypoxemia refractory to supplemental oxygen
(3) Uncorrected hypotension, defined as a persistent systolic pressure less than 90 mmHg

d) **Adverse Effects**
(1) Respiratory depression/arrest
(2) Altered mental status
(3) Increased vagal tone due to suppression of sympathetic pathways (slowed heart rate)
(4) Constricted pupils (pinpoint)
(5) Increased cerebral blood flow

e) **Precautions**
(1) *Naloxone* reverses all effects.
(2) To reduce the risk of chest wall rigidity (especially in children), fentanyl should be administered slowly and titrated to effect.
(3) Vital signs should be monitored frequently.
(4) Hypotension is a greater possibility in volume-depleted patients.
(5) Elderly patients and those with impaired renal function may be more sensitive to the medication’s effects.
f) **Dosage**
   
   (1) **Adult:** Fentanyl IN preferred IV/IO/IM
   
   (a) Administer 1 mcg/kg to a maximum initial dose of 200 mcg (For IN route, dosing may be limited due to volume limitations - administration of max 1mL per nare).
   
   (b) Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of fentanyl 1 mcg/kg to a maximum dose of 200 mcg.
   
   (c) Obtain on-line medical direction for additional doses, if required.

   (2) **Pediatric:** Fentanyl IN. If IN route not accessible, IV/IO/IM
   
   (a) Administer 1 mcg/kg to a maximum initial dose of 200 mcg (For IN route, dosing may be limited due to volume limitations - administration of max 1mL per nare).
   
   (b) Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of fentanyl 1 mcg/kg to a maximum dose of 200 mcg.
   
   (c) Obtain on-line medical direction for additional doses, if required.
ALS Pharmacology –
GLUCAGON

TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Hormone synthesized by the pancreas
   (2) Increases blood glucose concentration
   (3) Inhibits gastric and pancreatic secretions
   (4) May increase heart rate and cardiac output
   (5) May decrease blood pressure
   (6) Increases metabolic rate

b) Pharmacokinetics
   (1) Destroyed by the GI tract and is not effective orally
   (2) Maximum hyperglycemic activity occurs within 30 minutes and disappears after 1–2 hours.
   (3) Relaxation of smooth muscle occurs within 8–10 minutes and persists for 12–27 minutes.
   (4) The half-life is 3–10 minutes.
   (5) Degraded in liver and kidneys

c) Indications
   (1) Patients with altered mental status who are suspected of being hypoglycemic where IV access is not obtainable
   (2) Beta blocker overdose

d) Contraindications
   Known hypersensitivity

e) Adverse Effects
   Nausea and vomiting

f) Precautions
   Glucagon only works if liver has significant glycogen stores.

g) Dosage
   (1) For suspected hypoglycemia without IV access:
      (a) Adult: Administer 1 mg IM/IN (Medical consultation for additional dosing to a maximum of 3 mg IM)
      (b) Pediatric:
         (i) 1 mg IM/IN (5 years of age up to patient’s 18th birthday) (Medical consultation for additional dosing to a maximum of 3 mg IM/IN)
         (ii) 0.5 mg IM/IN (28 days–4 years of age) (Medical consultation for additional dosing to a maximum of 3 mg IM/IN)
   (2) For suspected beta blocker overdose:
      (a) Adult: Administer 1 mg IVP every 5 minutes
      (b) Pediatric: Administer every 5 minutes
         (i) 1 mg IVP (5 years of age up to patient’s 18th birthday) every 5 minutes
         (ii) 0.5 mg IVP (28 days–4 years of age) every 5 minutes
TRADE NAMES: HALDOL®
JURISDICTIONAL OPTION ONLY WHEN APPROVED BY THE STATE EMS MEDICAL DIRECTOR.

a) Pharmacology
   Antipsychotic

b) Pharmacokinetics
   Onset of action is within 10 minutes of the IM administration.

c) Indications
   Moderate agitation (defined as behavior that puts the patient or clinician at risk of harm) due to suspected psychiatric emergency (e.g., schizophrenia) or medical delirium

d) Contraindications
   (1) Children under 5 years of age
   (2) Parkinson’s disease
   (3) CNS depression
   (4) Acute CNS injury
   (5) Severe agitation (see midazolam and ketamine)

e) Adverse Effects
   (1) Dystonic reaction
      (a) Rarely seen with short-term use
      (b) ADULT: administer a single dose of diphenhydramine 25–50 mg IV/IM
      (c) PEDIATRIC: Administer a single dose of diphenhydramine 1 mg/kg IV/IM (max of 25 mg)
   (2) Hypotension and tachycardia
      (a) Administer Lactated Ringer’s fluid bolus if hypotension occurs
      (3) Torsades de pointes (polymorphic ventricular tachycardia)
      (a) Patients receiving haloperidol should be placed on the cardiac monitor to evaluate for dysrhythmias.

f) Precautions
   (1) Violent patients may require physical restraint while the medication is administered.
   (2) Patients receiving haloperidol should be placed on cardiac monitor to evaluate for dysrhythmias.

g) Dosage
   (1) ADULT
      (a) Patient 18–68 years of age: 5 mg IM
      (b) Patient 69 years and older: 2.5 mg IM
   (2) PEDIATRIC
      (a) Child less than 5 years of age: Contraindicated
      (b) Child 5–12 years of age: 0.05 mg/kg IM, max of 2.5 mg IM
      (c) Patient 13 up to 18th birthday: 2.5–5 mg IM
ALS Pharmacology – IPRATROPIUM

TRADE NAMES: ATROVENT®

a) Pharmacology
   (1) Anticholinergic (parasympatholytic) bronchodilator
   (2) Bronchodilator is site-specific, not systemic
   (3) Dries respiratory tract secretions
   (4) Most effective in combination with a beta-adrenergic bronchodilator

b) Pharmacokinetics
   (1) Improved pulmonary function in 15–30 minutes
   (2) Peak effects occur in 1–2 hours.
   (3) Duration of action is usually 4–5 hours.

c) Indications
   (1) Allergic reactions/anaphylaxis
   (2) Bronchial asthma
   (3) Reversible bronchospasms associated with chronic bronchitis and emphysema

d) Contraindications
   (1) Hypersensitivity to the drug
   (2) Hypersensitivity to atropine
   (3) Less than 1 year of age

e) Adverse Effects
   (1) More common: dry mouth, cough, or unpleasant taste
   (2) Less common: vision changes, eye burning or pain, dizziness, headache, nervousness, palpitations, sweating, trembling, chest tightness, rash, hives, or facial sweating

f) Precautions
   (1) Use with caution in patients with congestive heart failure, heart disease, hypertension, glaucoma, and with elderly patients.
   (2) May worsen the condition of glaucoma if it gets into the eyes. Having the patient close their eyes during nebulization may prevent this.
   (3) Not to be used as a single agent—must be used in combination with a beta-agonist.

g) Dosage
   (1) Adult:
      Single administration ONLY, 500 mcg (2.5 mL) by nebulized aerosol connected to 6–8 lpm of oxygen in combination with albuterol 2.5 mg.
(2) Pediatric:
   Single administration ONLY. In combination with *albuterol*,
   nebulized aerosol is connected to 6–8 lpm of *oxygen*.
   (a) **Less than 1 year of age**: contraindicated
   (b) **Age 1 year but less than 2 years**:
       250 mcg (1.25 mL) by nebulized aerosol
   (c) **Age 2 and older**:
       500 mcg (2.5 mL) by nebulized aerosol
TRADE NAMES: KETANEST®, KETASET®, KETALAR®

a) Pharmacology
Sedative-hypnotic; analgesic

b) Pharmacokinetics
A rapid-acting, non-barbiturate, sedative-hypnotic analgesic agent characterized by normal pharyngeal-laryngeal reflexes, normal or enhanced skeletal muscle tone, and possible cardiovascular and respiratory stimulation. It may occasionally produce transient respiratory depression.
ONSET OF ACTION FOR IV/IO KETAMINE MAY BE 5–10 MINUTES.
ONSET OF ACTION FOR IN/IM KETAMINE MAY BE UP TO 15–20 MINUTES.

c) Indications
(1) Moderate to severe pain; musculoskeletal, extremity, and back pain
(2) Severe agitation
(3) Ventilatory difficulty secondary to bucking or combativeness in intubated patients
(4) CPR-induced awareness

d) Contraindications
(1) Known hypersensitivity to ketamine
(2) Penetrating eye injury
(3) Chest pain, abdominal pain, flank pain, or headache
(4) Pregnancy/breastfeeding

e) Adverse Effects
(1) Although respiration is frequently stimulated, respiratory depression may occur with rapid IV administration. Laryngospasm has been known to occur.
(2) Although hypotension may occur, blood pressure and heart rate are frequently stimulated.
(3) Involuntary myoclonus that may mimic seizure activity – these movements should not be confused for seizures of emergence from sedation.
(4) Possible enhanced secretions
(5) Possible unpleasant dreams and delirium upon emergence from sedation

f) Precautions
(1) Cardiac and respiratory function should be continuously monitored when ketamine has been administered.
(2) Some patients who have received ketamine for control of severe agitation may require advanced airway management. ALS clinicians should be prepared to support the patient's airway needs.
(3) Ketamine is supplied in multiple concentrations (10 mg/mL, 50 mg/mL and 100 mg/mL). To avoid dosing errors, ALS clinicians check the concentration carefully and confirm that it is the proper dose for the patient prior to administering ketamine.
(4) Ketamine 100 mg/mL concentration is generally more appropriate for IM or IN administration, as there are maximum volume considerations with these routes. Ketamine 10 mg/mL concentration is preferable for IV administration for analgesia.
ALS Pharmacology – KETAMINE (continued)

11.21

**g) Dosage**

(1) **Pain Management**

(a) Adult: Administer 0.2 mg/kg IV/IO over 1–2 minutes. Maximum single dose 20 mg.
   (i) Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of ketamine 0.2 mg/kg IV/IO over 1–2 minutes. Maximum single dose 20 mg.
   (ii) If IV unavailable, administer 0.5 mg/kg IN/IM (if delivery device is available; divide administration of the dose equally between the nares to a maximum of 1 mL per nare).
   (iii) Reassess in 15 minutes. If pain remains moderate to severe, then administer a second dose of ketamine 0.5 mg/kg IN/IM.

(b) Pediatric: Administer 0.2 mg/kg IV/IO over 1–2 minutes. Maximum single dose 20 mg.
   (i) Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of ketamine 0.2 mg/kg IV/IO over 1–2 minutes. Maximum single dose 20 mg.
   (ii) If IV unavailable, administer 0.5 mg/kg IN/IM (if delivery device is available, divide administration of the dose equally between the nares to a maximum of 1 mL per nare).
   (iii) Reassess in 15 minutes. If pain remains moderate to severe, then administer a second dose of ketamine 0.5 mg/kg IN/IM.

(2) **Severe agitation**

   Medical consultation is required for the first dose of ketamine for severe agitation in all patients, unless there is immediate or imminent danger to the patient or EMS. Additional doses of ketamine always require medical consultation for all patients.

   (a) Adult

   (i) IV dosing: Administer 1 mg/kg IV/IO. Maximum single IV/IO dose 100 mg.
      If severe agitation persists, administer 1 mg/kg IV/IO. Maximum single IV/IO dose 100 mg. Maximum total IV/IO dose 200 mg.
   (ii) IM dosing: 4 mg/kg IM. Maximum total IM dose 400 mg.
      Additional dose of 4 mg/kg IM ketamine for persistent agitation requires medical consultation.

   (b) Pediatric

   (i) IV dosing: For patients who have not yet reached their 18th birthday, administer 1 mg/kg IV/IO. Maximum single IV/IO dose 100 mg. Maximum total IV/IO dose 200 mg.
      If severe agitation persists, administer 1 mg/kg IV/IO. Maximum single IV dose 100 mg.
   (ii) IM dosing: Patients who have not yet reached their 18th birthday, administer 4 mg/kg IM. Maximum IM dose 400 mg.
      Additional dose of 4 mg/kg IM ketamine for persistent agitation requires medical consultation.
(3) Ventilatory difficulty secondary to bucking or combativeness in intubated patients
   (a) Ketamine may be preferred for patients who have hypotension or possible hypovolemia, or if ventilatory difficulty is thought to be the result of pain response.
      (i) Dose: Administer 2 mg/kg IVP/IO over 60 seconds. May repeat 2 additional doses of 1 mg/kg for IVP/IO every 10–15 minutes to a total of 3 doses as needed. Additional doses require medical consultation.

(4) CPR-induced awareness
   (a) Adult
      (i) Administer ketamine 1 mg/kg IV/IO.
      (ii) Repeat doses with medical consultation.
   (b) Pediatric
      (i) Obtain medical consultation from a Pediatric Base Station.
1. Pharmacology
   a) Inhibits synthesis of prostaglandin, which, in turn, reduces pain and inflammation
   b) Antipyretic agent
   c) Does not affect CNS, peripheral acting analgesic, therefore, it does not possess the same sedative properties as a narcotic

2. Pharmacokinetics
   a) Onset: Approximately 30 minutes
   b) Peak effects: 1-2 hours
   c) Half-life: 4-6 hours

3. Indications
   a) Management of moderate to severe acute pain
   b) Consider as a first line medication for renal stones/colic
   c) Burns - mild to moderate
   d) Non-traumatic neuromuscular pain

4. Contraindications
   a) Hypersensitivity to ketorolac, aspirin, and other NSAIDs
   b) Current usage of aspirin or NSAIDs within 6 hours
   c) Severe headache or head injury
   d) Bleeding or clotting disorder
   e) Renal disease or transplant
   f) Active or history of peptic ulcer disease (PUD), active or recent history of GI bleed, and active or history of GI perforation
   g) Pregnancy or breast feeding
   h) Suspected ACS
   i) Trauma with suspected bleeding
   j) Patients who have not yet reached their 2nd birthday

5. Adverse Effects
   a) Burning or pain at the injection site
   b) Rash / itching
   c) GI distress
   d) Nausea / vomiting
6. Dosage

a) Adult: Administer single dose of 15 mg IV only. No repeat doses.
   If IV is unavailable: Administer single dose of 30 mg IM. No repeat doses.

b) Pediatric:
   (1) Newly born to 2 years of age: Contraindicated
   (2) Age 2 to patients who have not reached their 18th birthday: Administer 0.5 mg/kg IV
       only to a maximum total dose of 15 mg. No repeat doses.
       If IV is unavailable: Administer 1 mg/kg IM to a maximum total dose of
       30 mg. No repeat doses.
ALS Pharmacology –
LACTATED RINGER’S

TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Isotonic crystalloid solution
   (2) Lactated Ringer’s (LR) contains:
      (a) Sodium (Na+) ............ 130 mEq/liter
      (b) Potassium (K+) .......... 4 mEq/liter
      (c) Calcium (Ca++) .......... 3 mEq/liter
      (d) Chloride (Cl-) .......... 109 mEq/liter
      (e) Lactate ................. 28 mEq/liter

b) Pharmacokinetics
   Lactated Ringer’s is a water and electrolyte replacement.

c) Indications
   (1) Hypovolemia (limitation in multiple/severe trauma without head injury)
   (2) Keep vein open
   (3) Fluid boluses

d) Contraindications
   Fluid overload states

e) Adverse Effects
   Rare in therapeutic doses

f) Precautions
   (1) Patients receiving Lactated Ringer’s should be monitored to prevent circulatory overload.
   (2) Lactated Ringer’s should be used with caution in patients with congestive heart failure or renal failure.

g) Dosage
   (1) Adult:
      (i) For patients with multiple/severe trauma but without head injury: Administer small boluses of LR (maximum single bolus of 250 mL prior to additional blood pressure check) to achieve and maintain a systolic blood pressure of greater than or equal to 90 mmHg.
      (ii) For multiple/severe trauma with head injury: Administer small boluses of LR (maximum single bolus of 250 mL prior to additional blood pressure check) to achieve and maintain a systolic blood pressure greater than or equal to 110 mmHg.
      (iii) For all other patients: Titrate to a systolic pressure of 90 mmHg.
      (iv) Maximum dose 2,000 mL without medical consultation.
(2) Pediatric:
   (a) KVO
   (b) If age-related vital signs and patient’s condition indicate hypoperfusion, adminis-
       ter initial fluid bolus of 20 mL/kg LR IV/IO. Fluid boluses for neonates and vol-
       ume-sensitive children are 10 mL/kg.
   (c) If patient’s condition does not improve, administer the second fluid bolus of
       20 mL/kg LR IV/IO.
   (d) ☎ Third and subsequent fluid boluses at 20 mL/kg LR IV/IO with
       medical consultation.
TRADE NAMES: XYLOCAINE®

a) Pharmacology
   (1) Anesthesia for IO infusions
   (2) Nasal anesthesia

b) Pharmacokinetics
   (1) Extremely rapid (within minutes) onset following IV administration and lasts approximately 10–20 minutes
   (2) Mucosal anesthesia with onset in 1–5 minutes

c) Indications
   (1) Anesthesia for IO infusions
   (2) Nasal tracheal intubation

d) Contraindications
   (1) AV blocks
   (2) Sensitivity to lidocaine
   (3) Idioventricular escape rhythms
   (4) Accelerated idioventricular rhythm
   (5) Sinus bradycardia or arrest or block
   (6) Hypotension
   (7) Shock
   (8) Ventricular conduction defects

e) Adverse Effects
   (1) Lidocaine may cause clinical evidence of toxicity usually related to the central nervous system.
   (2) Toxicity:
      (a) Early: muscle twitching, slurred speech, altered mental status, decreased hearing, paresthesia (pins and needles), anxiety, apprehension, visual disturbances, nausea, numbness, difficulty breathing or swallowing, decreased heart rate
      (b) Late: convulsions, hypotension, coma, widening of QRS complex, prolongation of the P-R interval, hearing loss, hallucinations

f) Precautions
   (2) Reduce the dosage in patients with decreased cardiac output, liver dysfunction, and the elderly (over age 70).
ALS Pharmacology –
LIDOCAINE (continued)

g) Dosage

(1) Adult/Adolescent with an IO infusion: To prevent or treat pain during an IO infusion in patients greater than or equal to 13 years of age, administer 20–40 mg (1–2 mL) of 2% (preservative-free) lidocaine IO.

<table>
<thead>
<tr>
<th>Pediatric Lidocaine 2%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
</tr>
<tr>
<td>Less than 5 years</td>
</tr>
<tr>
<td>5-12 years</td>
</tr>
<tr>
<td>13-17 years</td>
</tr>
<tr>
<td>Greater than 18 years</td>
</tr>
</tbody>
</table>

(2) IO infusion in patients less than 13 years of age: To prevent or treat pain during an IO infusion for patients under 13 years of age, consult a Pediatric Base Station.

(3) Nasal Pharyngeal Anesthesia (age 13 years and greater)

Draw up 4 mL of lidocaine 4% (40 mg/mL) and using mucosal atomization device, administer 2 mL per nare. The patient IV, gel, and intranasal dosing should not exceed 3 mg/kg.

h) Interfacility Transport Only

(1) IV Infusion

(2) Maintain the IV infusion of lidocaine at the rate established by the sending physician and record vital signs every 15 minutes.

(See Lidocaine Infusion for Interfacility Transport.)
TRADE NAMES: Not Applicable

a) Pharmacology
Physiologic calcium channel blocker and also blocks neuromuscular transmission. Hypomagnesemia can cause cardiac dysrhythmias. It is also a CNS depressant effective in the management of seizures during pregnancy. It does this by decreasing the amount of acetylcholine liberated from motor nerve terminals. Magnesium is necessary for many biochemical processes and plays a role in the transmission of electrical impulses.

b) Pharmacokinetics
With intravenous administration the onset of anticonvulsant action is immediate and lasts about 30 minutes. Magnesium is excreted solely by the kidney at a rate proportional to the plasma concentration and glomerular filtration rate.

c) Indications
(1) Torsades de pointes
(2) Seizures with pregnancy
(3) Refractory VF and VT after amiodarone administration
(4) Moderate to severe asthma/bronchospasm exacerbation

d) Contraindications
(1) Heart blocks
(2) Renal impairment
(3) Hypermagnesemia

e) Adverse Effects
(1) Respiratory depression
(2) Flushing
(3) Sweating
(4) Hypotension
(5) Depressed reflexes

f) Precautions
(1) May exaggerate effects of CNS depressants and neuromuscular blocking agents
(2) Due to concern of hypotension, IV fluid bolus should be initiated if hypovolemia is suspected.
(3) Magnesium toxicity is a concern with higher doses and would present with respiratory depression, decreased reflexes, flaccid paralysis, and apnea. Calcium chloride 500 mg SLOW IVP for above indications of toxicity.
g) Dosage
   (1) Adult:
      (a) Seizure activity associated with pregnancy: 4 grams IV/IO over 10 minutes
          (mixed in 50–100 mL of approved diluent)
      (b) Refractory VT/VF: 1–2 grams IV/IO over 2 minutes
      (c) Moderate to severe asthma/bronchospasm exacerbation: 1–2 grams IV/IO over 10–20 minutes
          (mixed in 50–100 mL of approved diluent)
      (d) Torsades de pointes: 1–2 grams IV/IO over 2 minutes
   (2) Pediatric (under 18-years-old):
      (a) Seizure activity associated with pregnancy: 4 grams IV/IO over 10 minutes (mixed in
          50–100 mL of approved diluent)
      (b) Moderate to severe asthma/bronchospasm exacerbation: consider magnesium sulfate
          50 mg/kg IV/IO (mixed in 50 - 100 mL of approved diluent) to max of 2 grams given over
          10–20 minutes

MAGNESIUM ADMINISTRATION OFTEN CAUSES HYPOTENSION IN CHILDREN. CONSIDER
ADMINISTERING BOLUS 20 ML/KG OF LACTATED RINGER’S WITH THE ADMINISTRATION OF
MAGNESIUM.
   (c) Torsades de pointes: 25 mg/kg to a max of 2 grams IV/IO over 2 minutes

h) Interfacility Transport
   (1) A paramedic may administer continuous infusion established by a sending facility,
       not to exceed the ordered total dose, and monitoring the patient for signs and
       symptoms of magnesium toxicity.
   (2) Magnesium sulfate used for tocolytic control is an RN-level indication.
TRADE NAMES: VERSED®

a) Pharmacology
   (1) Sedative
   (2) Hypnotic
   (3) Anticonvulsant

b) Pharmacokinetics
   (1) A short-acting benzodiazepine with strong hypnotic, anticonvulsant activity, and amnestic properties
   (2) Onset of action is extremely rapid following IV administration; approximately 1.5 minutes, and for IM approximately 15 minutes.
   (3) Duration of effect is 1–4 hours with half-life of 1.5 to 3 hours in healthy adult.

c) Indications
   (1) Sustained and/or recurrent seizures
   (2) Precardioversion to reduce anxiety
   (3) Awake patient requiring transcutaneous pacing (TCP)
   (4) Nasal Tracheal Intubation
   (5) Implanted Cardioverter Defibrillator (ICD) Malfunction
   (6) Nerve/organophosphate exposure
   (7) Bucking Endotracheal Intubated patient
   (8) Moderate to severe stimulant toxicity
   (9) Moderate or severe agitation

d) Contraindications
   (1) Hypotension (See below for ET bucking)
   (2) Known hypersensitivity to midazolam

e) Adverse Effects
   (1) Respiratory depression or apnea
   (2) Hypotension

f) Precautions
   (1) The effects of midazolam can be accentuated and significantly potentiated by CNS depressants, such as opioids or alcohol.
   (2) Midazolam is five times as potent per milligram as diazepam and there is an increased risk of respiratory depression.
g) Dosage

(1) All indications (except for moderate to severe agitation and bucking endotracheal intubated patients)
   (a) Adult:

   REDUCE THE DOSAGE BELOW IV/IO/IN/IM BY 50% FOR PATIENTS 69 YEARS OR OLDER.

   (i) 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes per increment with maximum single dose 5 mg.

   (ii) If IV unavailable, 5 mg IN/IM may be administered. IN administration max 1 mL per nare

   (iii) Additional doses up to a maximum total dose 10 mg require medical consultation for all clinicians.
       For seizures lasting greater than 10 minutes (status), consider IO administration of midazolam.

   (iv) If suspected severe nerve agent exposure, clinicians may administer midazolam 5 mg IM without medical consultation.

(b) Pediatric:
   (i) 0.1 mg/kg in 2 mg increments. SLOW IVP over 1–2 minutes per increment to a maximum single dose of 5 mg.

   (ii) If IV unavailable, 0.2 mg/kg IN/IM
       IN administration max 1 mL per nare
       Maximum total dose 5 mg

   (iii) Additional doses up to a maximum total dose 5 mg require medical consultation for all clinicians.
       For life-threatening conditions, consider IO administration of midazolam.

   (iv) If suspected severe nerve agent exposure, clinicians may administer midazolam as above without medical consultation.

(2) Moderate to Severe Agitation
   (a) Patient 18–69 years: midazolam 5 mg IM/IV
       Patient greater than 69 years: midazolam 2.5 mg IM/IV

   (b) For patients 18 years and older, if severe agitation persists after giving one dose of ketamine, consider midazolam 2.5 mg IV/IO.

   (c) For patients 18 years and older, if IV/IO unavailable:
       If severe agitation persists after IM ketamine dose, administer midazolam 5 mg IM.
(d) For patients aged 5–12 years of age: medical consultation is required for midazolam 0.1 mg/kg IV or 0.2 mg/kg IM/IN (max 5 mg).

   IM route preferred.

(e) For patients age 13 to those who have not yet reached their 18th birthday:

   (i) Medical consultation is required for midazolam 0.1 mg/kg or 0.2 mg/kg IM/IN (max 5 mg). IM route preferred. For severe agitation where there is immediate and imminent risk to the patient or EMS, no consult is required.

   (ii) If severe agitation persists after giving one dose of ketamine IV/IO, consider midazolam 2.5 mg IV/IO.

   (iii) If IV/IO unavailable:

      If severe agitation persists after IM ketamine dose, administer midazolam 5 mg IM.

(3) Bucking Endotracheal Intubated patient

   (a) Adult: Administer 0.1 mg/kg SLOW IVP over 1–2 minutes, while maintaining systolic BP greater than 90 mmHg (110 mmHg if injuries include a suspected head injury). STOP ONCE BUCKING HAS RESOLVED AND VENTILATION IS RELAXED. Maximum single dose is 5 mg.

      Additional doses require medical consultation.

   (b) Pediatric: Administer 0.05 mg/kg SLOW IVP over 1–2 minutes, while maintaining systolic BP greater than 60 in neonates, 70 in infants, [70 + (2 x years) = systolic BP] for patients greater than 1 year of age. Maximum total dose 5 mg.
**TRADE NAMES:** Not Applicable

*(Optional Supplemental Protocol, which allows for jurisdictional selection of both fentanyl and morphine OR replacement of fentanyl by morphine as the opioid of choice)*

**Pharmacology**

1. Decreases pain perception and anxiety
2. Relaxes respiratory effort
3. Causes peripheral dilation, which decreases preload
4. Decreases left ventricular afterload

**a) Pharmacokinetics**

1. Binds with opiate receptors in the CNS, altering both perception and emotional response to pain
2. Onset of action is in less than 5 minutes after IV dose and effects last 4–5 hours.
3. Causes peripheral arterial and venous vasodilation

**b) Indications**

1. The patient reports moderate to severe pain.
2. In the clinician’s judgment the patient will benefit from treatment with an opioid analgesic, including patients who are MOLST and/or EMS/DNR patients or being pre-medicated for a procedure.
3. Pulmonary Edema/Congestive Heart Failure (Pediatric only)

**c) Contraindications**

1. Hypersensitivity or known allergy to morphine
2. Uncorrected respiratory distress or hypoxemia refractory to supplemental oxygen
3. Uncorrected hypotension, defined as a persistent systolic pressure less than 90 mmHg

**d) Adverse Effects**

1. Respiratory depression/arrest
2. Altered mental status (decreased level of consciousness)
3. Increased vagal tone due to suppression of sympathetic pathways (slowed heart rate)
4. Nausea and vomiting
5. Constricted pupils (pinpoint)
6. Increased cerebral blood flow
e) Precautions
   (1) Naloxone reverses all effects.
   (2) Should be administered slowly and titrated to effect.
   (3) Vital signs should be monitored frequently.
   (4) Hypotension is a greater possibility in volume-depleted patients.

f) Dosage
   (1) Adult: IV/IM
      (a) Administer 0.1 mg/kg to a maximum initial dose of 20 mg.
      (b) Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of morphine 0.05 mg/kg to a maximum additional dose of 10 mg.
      (c) Obtain on-line medical direction for additional doses, if required.
   (2) Pediatric: IV/IM
      (a) Administer 0.1 mg/kg to a maximum initial dose of 20 mg.
      (b) Reassess in 5–10 minutes. If pain remains moderate to severe, then administer a second dose of morphine 0.05 mg/kg to a maximum additional dose of 10 mg.
      (c) Obtain on-line medical direction for additional doses, if required.
   (3) Pediatric Pulmonary Edema/CHF
      (a) 0.1 mg/kg SLOW IVP/IO/IM (1–2 mg/min).
      Maximum dose 5 mg.
TRADE NAMES: NARCAN®

a) Pharmacology
Reverses all effects due to opioid (morphine-like) agents. This drug will reverse the respiratory depression and all central and peripheral nervous system effects.

b) Pharmacokinetics
(1) Onset of action is within a few minutes if administered IVP and within 5 minutes if administered IN.
(2) Intramuscular and pediatric/neonatal endotracheal administration results in a slower onset of action.
(3) Patients responding to naloxone may require additional doses and transportation to the hospital since most opioids last longer than naloxone.
(4) Has no effect in the absence of opioids

c) Indications
To reverse respiratory depression induced by opioids

d) Contraindications
Patients under 28 days of age.

e) Adverse Effects
Opioid withdrawal

f) Precautions
(1) Naloxone may induce opioid withdrawal in patients who are physically dependent.
(2) Certain drugs may require much higher doses of naloxone for reversal than are currently used.
(3) Should be administered and titrated so respiratory efforts return, but not intended to restore full consciousness

g) Dosage
(1) Adult: Administer 0.4–2 mg IVP/IO (titrated)/IM/IN (if delivery device is available, divide administration of the dose equally between the nares to a maximum of 1 mL per nare); OR administer 4 mg/0.1 mL IN in one nare. Repeat as necessary to maintain respiratory activity.
(2) Pediatric: Administer 0.1 mg/kg IVP/IO (titrated)IM/IN (if delivery device is available, divide administration of the dose equally between the nares to a maximum of 1 mL per nare); OR administer 4 mg/0.1 mL IN in one nare. May be repeated as necessary to maintain respiratory activity. ET dose: 0.2–0.25 mg/kg
TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Vasodilator—effect on veins more than arteries
   (2) Decreases right heart return (preload) by venous pooling, thereby decreasing myocardial workload and oxygen consumption

b) Pharmacokinetics
   (1) Absorbed through oral mucosa
   (2) Antianginal and vasodilation effects within 1–2 minutes after administration. Half-life is 1–4 minutes.
   (3) Duration of action is less than 5 minutes.

c) Indications
   (1) For treatment of angina
   (2) Congestive heart failure, acute pulmonary edema

d) Contraindications
   (1) Known hypersensitivity
   (2) Pediatric patient under the age of 13
   (3) Any patient having taken medication for Pulmonary Artery Hypertension (e.g., Adcirca® or Revatio®) or erectile dysfunction (e.g., Viagra®, Levitra®, or Cialis®) within the past 48 hours. Medical consultation is required to override this contraindication.
   (4) Asymptomatic hypertension
   (5) Blood pressure below 90 mmHg systolic
   (6) Heart rate less than 60 or greater than 150 bpm

e) Adverse Effects
   Headache, hypotension, nausea, vomiting, dizziness, and decreased level of consciousness

f) Precautions
   (1) May cause hypotension
   (2) If systolic blood pressure drops more than 20 mmHg per dose, obtain a medical consultation for further doses.

g) Dosage
   (1) Adult: Chest pain
      (a) If patient has a prescription or previous history of nitroglycerin use, administer nitroglycerin: 0.4 mg SL (may repeat dose 2 times at 3–5 minute intervals) May be repeated if symptoms persist, BP is greater than 90 mmHg, and pulse is between 60-150 bpm, to a maximum dose of 1.2 mg
      (b) If patient does not have a prescription or previous history of nitroglycerin use, establish IV prior to the administration of nitroglycerin, then administer nitroglycerin as above.
      (c) Additional doses may be administered with medical consultation.
(2) Adult: Pulmonary Edema/Congestive Heart Failure
   (a) Low dose - Administer 0.4 mg SL at 3–5 minute intervals to a maximum dose of 1.2 mg.
   (b) High dose - (until CPAP is applied or if CPAP is not tolerated)
      (i) Administer 1 dose of 0.4 mg SL and apply 1 inch of NTG paste.
      (ii) Administer 1 dose of 0.8 mg SL.
      (iii) Continue 0.8 mg NTG dosing to achieve a 20% reduction in systolic blood pressure.

(3) Pediatric: Requires medical consultation from Pediatric Base Station.
TRADE NAMES: Not Applicable

a) Pharmacology
Nitroglycerin paste contains a 2% solution of nitroglycerin in a special absorbent paste. When placed on the skin, nitroglycerin is absorbed into the systemic circulation. In many cases, it may be preferred over nitroglycerin tablets because of its longer duration of action.

b) Pharmacokinetics
Nitroglycerin is a rapid smooth-muscle relaxant that reduces cardiac work and, to a lesser degree, dilates the coronary arteries. This results in increased coronary blood flow and improved perfusion of the ischemic myocardium. Relief of ischemia causes reduction and alleviation of chest pain. Pain relief following transcutaneous nitroglycerin administration usually occurs within 5 to 10 minutes, and therapeutic effects can be observed up to 30 minutes later. Nitroglycerin also causes vasodilation, which decreases preload. Decreased preload leads to decreased cardiac work. This feature, in conjunction with coronary vasodilation, reverses the effects of angina pectoris.

c) Indications
Patients in respiratory distress with moderate or severe symptoms and elevated systolic blood pressure.

d) Contraindications
(1) Known hypersensitivity
(2) Pediatric patient under the age of 13
(3) Any patient having taken medication for Pulmonary Artery Hypertension (e.g., Adcirca® or Revatio®) or erectile dysfunction (e.g., Viagra®, Levitra®, or Cialis®) within the past 48 hours. Medical consultation is required to override this contraindication.
(4) Asymptomatic hypertension
(5) Blood pressure below 90 mmHg systolic
(6) Heart rate less than 60

e) Adverse Effects
Headache, dizziness, weakness, tachycardia, hypotension, orthostasis, skin rash, dry mouth, nausea, and vomiting.

f) Precautions
Patients taking the drug routinely may develop a tolerance and require an increased dose. Headache is a common side effect of nitroglycerin administration and occurs as a result of vasodilation of the cerebral vessels.

Postural syncope sometimes occurs following the administration of nitroglycerin. This should be anticipated and the patient kept supine when possible. It is important to monitor the blood pressure continuously.

g) Dosage
(1) Adult: 1 inch of the NTG paste is applied. Measuring applicators are supplied.
(2) Pediatric: Requires medical consultation from Pediatric Base Station.
TRADE NAMES: ZOFTRAN®

a) Pharmacology
   A selective blocking agent of the serotonin 5-HT3 receptor type

b) Pharmacokinetics
   Anti-nausea and anti-emetic with onset of action within 5–15 minutes IV and 30 minutes IM

c) Indications
   (1) Prevention and control of nausea and/or vomiting
   (2) Ondansetron can be administered in an effort to reduce the nausea or vomiting complications associated with certain existing injuries, medical illness, or medication side effects (e.g., penetrating eye injury, high risk for aspiration, or following opioid administration).

d) Contraindications
   Known hypersensitivity to ondansetron
   Patients less than 28 days

e) Adverse Effects
   (1) Hypotension
   (2) Tachycardia
   (3) Extrapyramidal reactions
   (4) Seizures
   (5) QT interval prolongation

f) Precautions
   (1) Monitor EKG, pulse oximetry, and blood pressure.
   (2) Have emesis basin and suction ready.

g) Dosage
   (1) Adult: 8 mg SLOW IV over 2–5 minutes OR 4-8 mg IM OR 8 mg orally disintegrating tablet (ODT)
       May repeat once without medical consultation.
       For third repeat dose to a patient with maximum total dose of 24 mg.

   (2) Pediatric:
       Patients 28 days to 12 years old: 0.1 mg/kg SLOW IV over 2–5 minutes
       Patients who are 13 to 18 years old: 8 mg ODT OR 8 mg SLOW IV over 2–5 minutes
       OR
       If no IV: 0.1 mg/kg IM (with max single dose of 8 mg);
       May repeat once without medical consultation.
       For third repeat dose to a patient with maximum total dose of 0.3 mg/kg or 24 mg, whichever is lower.
TRADE NAMES: Not Applicable

a) Pharmacology
   (1) Increases oxygen content of the blood
   (2) Improves tissue oxygenation
   (3) Decreases energy expended for respirations

b) Pharmacokinetics
   Changing the percentage of inspired oxygen results in an increased blood and tissue level equilibration within 5–20 minutes.

c) Indications
   (1) If evidence of hypoxia (Less than 94% SpO₂)
   (2) Respiratory distress
   (3) Cardiopulmonary arrest
   (4) Trauma
   (5) Suspected CO exposure
   (6) Dyspnea

d) Contraindications
   Not clinically significant

e) Adverse Effects
   High concentrations of oxygen will reduce the respiratory drive in some COPD patients; these patients should be carefully monitored.

f) Precautions
   (1) Never withhold oxygen from those who need it.
   (2) Oxygen should be given with caution to patients with COPD.
   (3) Simple or partial rebreather face masks must be supplied with a minimum 6 lpm.
   (4) Non-breather (NRB) face masks must be supplied with a minimum 12 lpm.

g) Dosage
   (1) Adult: Administer 12–15 lpm via NRB mask or 2–6 lpm via nasal cannula, as needed. CO exposure: Administer 100% oxygen via NRB mask. Maintain SpO₂ at 100%
   (2) Pediatric: Administer 12–15 lpm via NRB mask or 2-6 lpm via nasal cannula, as needed. CO exposure: Administer 100% oxygen via NRB mask. Maintain SpO₂ at 100%

<table>
<thead>
<tr>
<th>Percent O₂ Saturation</th>
<th>Ranges</th>
<th>General Patient Care</th>
</tr>
</thead>
<tbody>
<tr>
<td>94–100%</td>
<td>Normal</td>
<td>Give oxygen as necessary</td>
</tr>
<tr>
<td>91–93%</td>
<td>Mild Hypoxia</td>
<td>Give oxygen as necessary</td>
</tr>
<tr>
<td>86–90%</td>
<td>Moderate Hypoxia</td>
<td>Give 100% oxygen</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Assisting Ventilations if necessary</td>
</tr>
<tr>
<td>less than or equal to 85%</td>
<td>Severe Hypoxia</td>
<td>Give 100% oxygen</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Assist Ventilations</td>
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<tr>
<td></td>
<td></td>
<td>If indicated, Intubate</td>
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</tbody>
</table>

INACCURATE OR MISLEADING SpO₂ READINGS MAY OCCUR IN THE FOLLOWING PATIENTS: HYPOTHERMIC, HYPOPERFUSION (SHOCK), CO POISONING, HEMOGLOBIN ABNORMALITY, ANEMIA, AND VASOCONSTRICTION.
ALS Pharmacology –
SODIUM BICARBONATE

TRADE NAMES: Not Applicable

a) Pharmacology
Sodium bicarbonate corrects acidosis; raises blood pH

b) Pharmacokinetics
(1) Rapid onset of action in the blood
(2) Delayed onset of action in the tissues

c) Indications
(1) Acidosis, pre-existing (documented)
(2) Cardiac arrest only with a high clinical suspicion of acidosis, hyperkalemia, or NA channel blocker (including tricyclic antidepressant and phenobarbital) overdose as a cause of the arrest
(3) Hyperkalemia
(4) NA channel blocker (including tricyclic antidepressant and phenobarbital) overdose

d) Contraindications
Preexisting alkalosis

e) Adverse Effects
(1) CHF exacerbation/volume overload
(2) Hypernatremia (high sodium)
(3) Hypokalemia
(4) Intracellular acidosis due to carbon dioxide formation
(5) Metabolic alkalosis
(6) Shifting the oxyhemoglobin dissociation curve, inhibiting the release of oxygen to the tissues

f) Precautions
Inactivates simultaneously-administered catecholamines (epinephrine)

g) Dosage
(1) Cardiac arrest only with a high clinical suspicion of acidosis, hyperkalemia or NA channel blocker (including tricyclic antidepressant and phenobarbital) overdose as the cause:
   (a) Adult: Administer 1 mEq/kg IVP bolus initially; repeat 0.5 mEq/kg every 10 minutes.
   (b) Pediatric: Administer 1 mEq/kg IVP/IO; for patients less than 1 year of age, must be diluted (1:1) with Lactated Ringer’s.
(2) Hyperkalemia:
   Adult: Sodium bicarbonate 50 mEq IV/IO, slow over 5 minutes. Flush the IV with 5 mL of Lactated Ringer’s between calcium and sodium bicarbonate administration.
(3) Crush Syndrome (or patients with functional kidneys by history):
   (a) Adult: Sodium bicarbonate 50 mEq IV/IO, slow over 5 minutes and then initiate drip of sodium bicarbonate 100 mEq in 1,000 mL Lactated Ringer’s to run over 30–60 minutes.
(b) Pediatric: Sodium bicarbonate 1 mEq/kg IV/IO, slow over 5 minutes. For patients less than 1 year of age, must be diluted 1:1 with Lactated Ringer’s.

(4) NA channel blocker (including tricyclic antidepressant and phenobarbital) overdose (sodium channel blockade):
(a) Adult: Administer 1 mEq/kg IVP/IO bolus initially; repeat 0.5 mEq/kg every 10 minutes. Monitor QRS interval for narrowing.
(b) Pediatric: Administer 1 mEq/kg IVP/IO; for patients less than 1 year of age, must be diluted (1:1) with Lactated Ringer’s.
ALS Pharmacology – TERBUTALINE SULFATE

TRADE NAMES: Not Applicable
(JURISDICTIONAL OPTION ONLY WHEN APPROVED BY STATE EMS MEDICAL DIRECTOR)

a) Pharmacology
   (1) Stimulates beta-2 receptors located in the smooth bronchioles
   (2) Causes relaxation of bronchospasm

d) Pharmacokinetics
   Relieves bronchospasm in acute and chronic airway disease with minimal cardiovascular effect

c) Indications
   (1) Asthma
   (2) Reversible airway obstruction associated with bronchitis or emphysema

e) Indications
   (1) Patients under 12 years of age

g) Contraindications
   (1) Patients under 12 years of age

e) Adverse Effects
   (1) Tachycardia
   (2) Palpitations
   (3) Nervousness
   (4) Tremors
   (5) Dizziness
   (6) Nausea
   (7) Vomiting

f) Precautions
   (1) Exercise caution when administering to patients with hypertension or cardiac history
   (2) Monitor EKG

g) Dosage
   (1) Patients 12 years of age and older:
      Administer 0.25 mg IM. May repeat one time after 15 minutes if there is not adequate improvement. Maximum total dose 0.5 mg IM.
   (2) Patients less than 12 years of age: Not indicated
ALS Pharmacology –
TRANEXAMIC ACID (TXA)

a) Pharmacology
   Anti-fibrinolytic medication

b) Pharmacokinetics
   Onset: variable; Peak effect: 2 hours; Duration: 10 hours

c) Indications
   Suspected hemorrhagic shock (SBP less than 90) due to traumatic mechanism;
injury must have occurred within the past one (1) hour

d) Contraindications
   (1) Patients less than 15 years of age
   (2) Hypersensitivity or allergy to TXA
   (3) Known arterial or venous thromboembolism (PE, DVT)
   (4) Patients more than one (1) hour from time of injury

e) Adverse Effects
   (1) Hypotension (if given faster than 100 mg/min)
   (2) Seizures

f) Precautions
   (1) Administer over 10 minutes to reduce the risk of hypotension
   (2) Do not delay transport to administer TXA

g) Dosage
   For patients 15 years of age and older:
   Administer 1 gram in 100 mL approved diluent (normal saline/Lactated Ringer’s/
   D5W) IV/IO over 10 minutes
TRADE NAMES: ISOPTIN®

JURISDICTIONAL OPTION ONLY WHEN APPROVED BY THE STATE EMS MEDICAL DIRECTOR. ADMINISTRATION OF VERAPAMIL REQUIRES MEDICAL CONSULTATION.

a) Pharmacology
   Calcium channel blocker

b) Pharmacokinetics
   (1) Inhibits the movement of calcium ions across cardiac muscle cells
   (2) Decreases conduction velocity and ventricular rate

c) Indications
   (1) Narrow complex symptomatic atrial fibrillation or atrial flutter

d) Contraindications
   (1) Hypotension below 100 mmHg, second or third degree heart block, hypersensitivity to the drug
   (2) Patient with history of Wolff-Parkinson-White syndrome
   (3) Ventricular tachycardia
   (4) Patients less than 18 years of age

e) Precautions
   Use cautiously in patients with renal failure, congestive heart failure, or on beta blockers.

f) Adverse Effects
   (1) Hypotension (see Treatment of Overdose or Other Adverse Reactions)
   (2) Bradycardia
   (3) Vomiting
   (4) Nausea
   (5) Headache

g) Significant Interactions
   Congestive heart failure may result if used along with beta blockers.

h) Dosage
   (1) Adult:
      a) 2.5–10 mg slow IV over 2 minutes; if response is not adequate, repeat in 15 minutes with a dosage of 2.5–10 mg slow IV over 2 minutes with medical consultation.
   (2) Pediatric:
      Contraindicated for patients less than 18 years of age.
ALS Pharmacology – VERAPAMIL (continued)

i) **Overdose or Toxicity Presentation**
   Generally consists of exaggeration of side effects, including severe hypotension and symptomatic bradycardia.

j) **Treatment of Overdose or Other Adverse Reactions**
   (1) Give general supportive measures, monitor vitals, administer oxygen.
   (2) Hypotension:
      (a) If lungs are clear, administer fluid bolus 20 mL/kg of Lactated Ringer's; titrate to a systolic blood pressure of 100 mmHg.
      (b) If rales are present, administer fluid bolus, maximum of 250 mL of Lactated Ring-er's. Titrate to a systolic of 100 mmHg.
      (c) Administer calcium chloride 500 mg SLOW IVP.
   (3) Bradycardia: Consider atropine (0.5 to 1 mg); if necessary, consider pacing.
a) PURPOSE
Accessing a preexisting central venous catheter or device may be required for fluid volume resuscitation and/or medication administration for critically ill/injured patients when peripheral IV access cannot be established.

b) INDICATIONS
Life-Threatening Emergency
A preexisting central venous access catheter or device may be accessed by a paramedic for resuscitation medication administration or fluid volume administration.

A CRT-I may access these devices WITH MEDICAL CONSULTATION.

Non–Life-Threatening Emergency
Medical consultation is required for all ALS (CRT-I and paramedic) clinicians.

c) CONTRAINDICATIONS
None

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
(1) Infection (local site and in the central bloodstream)
(2) Air in the catheter line (air embolism)
(3) Damage to catheter line
(4) Obstruction in the line
(5) Dislodge the catheter

e) PROCEDURE: PORTS (e.g., Port-a-Cath®, Mediport®, Bard®, Infuse-a-Port®)
A port (reservoir) is a disc about an inch in diameter that is just under the skin, usually on the upper chest. Under the skin, it is connected to a catheter line that lies in a large vein just above the heart.

(1) Explain the procedure to the patient whenever possible.
(2) Obtain assistance as needed.
(3) Position the patient supine.
(4) Using a 10 mL syringe or larger, draw up TWO 5 mL flushes with NS/RL.
   NOTE: 10 mL syringes are used because they have lower pressure when flushing fluids than smaller volume syringes (1 mL, 3 mL, or 5 mL). The smaller volume syringes may deliver enough pressure to break the catheter.
(5) Open the right-angle, non-coring (Huber® or Gripper®) needle package and flush with NS/RL. Be sure there are no air bubbles in the tubing.
(6) Clean the skin site at the port with cleaning material from patient/family, or use alcohol or other approved antibacterial agent (e.g., ChloraPrep®), using a circular motion.
(7) Use sterile latex-safe gloves. Using the non-dominant hand, palpate the area over the port to stabilize the port and locate the center.
(8) With other hand, insert the non-coring needle into the center of the port with firm, steady pressure until you feel the needle reach the back of the port. Do not rock the non-coring needle back and forth in the port.
Procedures – ACCESSING CENTRAL VENOUS CATHETERS AND DEVICES (continued)

(9) Aspirate 5 mL of blood and/or heparinized solution and discard. If unable to aspirate blood, verify needle position by gently pushing the needle farther against the backstop of the port.
   If you are still unable to aspirate blood or fluid, contact MEDICAL CONSULTATION prior to use.

(10) Flush with 5 mL NS/RL while assessing for swelling at the site. **Be sure there are no air bubbles in the syringe or tubing.** Do not force flush if resistance is met. Verify the non-coring needle position by gently pushing the needle further against the backstop of the port, and attempt to flush again.

(11) After assessing patency, clamp the tubing, and remove the syringe.

(12) Apply needleless injection cap, if available, and cleanse with alcohol.

(13) IV fluids, tubing, and connectors must be assembled and primed in the cleanest area possible with **all air eliminated** prior to connecting to the patient.

(14) Attach the completely flushed IV line, unclamp the needle tubing, and begin infusion of fluid/medication. **NOTE:** IV fluids may not infuse by gravity.

(15) Secure the non-coring needle with sterile 2x2 or 4x4 and tape or occlusive dressing, being careful not to tape over the insertion site.

(16) Tape or loop extension tubing to outside of dressing.

f) PROCEDURE: TUNNELED AND NON-TUNNELED LINES

TUNNELED LINES (e.g., Hickman®, Groshong®, Broviac®, Cook®)
A tunneled central line is a catheter that is inserted under the skin of the chest, and the tip of the catheter is in a large vein just above the heart. A tunneled catheter has a cuff below the skin that the soft tissue grows into, reducing the risk of dislodgement and infection. These can be single or multiple-lumen catheters.

NON-TUNNELED LINES: PICC and MLC (e.g., Cook®, Neo-PICC®)
A PICC (Peripherally Inserted Central Catheter) line is a thin catheter that is inserted into one of the large veins, usually in the arm near the bend of the elbow, but may be in the neck or a lower extremity, and is threaded in a large vein just above the heart. A MLC (Mid-Line Catheter) is a thin peripheral catheter that is inserted into a large vein in the elbow and ends in the vein before the shoulder. Both of these catheters have a very small lumen and are considered “low volume lines” and not appropriate for volume resuscitation.

(1) Explain the procedure to the patient whenever possible.

(2) Obtain assistance as needed.

(3) Position the patient supine.

(4) Using a 10 mL syringe or larger, draw up 5 mL flushes with NS/RL. **Be sure there are no air bubbles in the syringe.** Attach a stopcock if available. **NOTE:** 10 mL syringes are used because they have lower pressure when flushing fluids than smaller volume syringes (1 mL, 3 mL, or 5 mL). The smaller volume syringes may deliver enough pressure to break the catheter.

(5) Use sterile latex-safe gloves.

(6) If multiple lumens or ports, determine from patient/family which catheter is most appropriate for use, if possible, or refer to the EIF Form. This is usually the **white** port.
Procedures – ACCESSING CENTRAL VENOUS CATHETERS AND DEVICES (continued)

12.1 Procedures: Accessing Central Venous Catheters and Devices

(7) Clean the existing cap on catheter with alcohol for 30 seconds.
(8) Clamp all lines with special clamps that do not have teeth, which might damage the catheter.
(9) Access the appropriate catheter port with a 10 mL syringe.
(10) Unclamp the catheter line to be accessed and aspirate 5 mL of blood/heparinized solution and discard to confirm placement and access patency. Delete this step if less than 2 Fr PICC catheter, as this may damage the catheter (the lumen is very small and the catheter wall may collapse and any blood in the catheter will form a clot).

**NOTE:** Contact MEDICAL CONSULTATION if unable to aspirate blood/fluid, or less than 2 Fr catheter.

(11) Reclamp the catheter any time you are changing lines or syringes. Remember that regular clamps may damage the central line tubing.
(12) Attach the flush syringe and unclamp.
(13) Flush with 5 mL NS/RL. **Be sure there are no air bubbles in the syringe or tubing.**
(14) Clamp this line again with the special clamp.
Procedures – AIRWAY MANAGEMENT:
BAG-VALVE-MASK VENTILATION

a) PURPOSE

(1) Bag-valve-mask (BVM) ventilation is the technique of providing rescue breathing for patients with inadequate respiratory effort or cardiac arrest. Patients in respiratory failure may respond to BVM ventilation and not require endotracheal intubation.

(2) A BVM may also be used to administer inhaled medications for patients with severe respiratory failure.

b) INDICATIONS

(1) Inadequate respiratory rate
   (a) Adult less than 8
   (b) Adolescent (13–18 years of age) less than 12
   (c) Child (1–12 years of age) less than 16
   (d) Infant/Toddler (less than 1 year of age) less than 20

(2) Inadequate respiratory effort
   (a) Absent or diminished breath sounds
   (b) Paradoxical breathing (chest and abdomen moving in opposite directions)
   (c) Cyanosis or oxygen saturation less than 90% on 100% oxygen by nonrebreather with the exception of patients with chronic hypoxemia

(3) Symptomatic Bradycardia
   (a) Adult/Adolescent Heart rate less than 60
      (greater than 13 years of age)
   (b) Child (1–12 years of age) Heart rate less than 80
   (c) Infant (less than 1 year of age) Heart rate less than 100

(4) Cardiac arrest

(5) Altered mental status
   Glasgow Coma Scale of 8 or less

c) CONTRAINDICATIONS

None

d) POTENTIAL ADVERSE EFFECTS / COMPLICATIONS

(1) Gastric distension
(2) Vomiting
(3) Increased intracranial pressure as a result of increased vagal stimulation if mask applied over the patient’s eyes

e) PRECAUTIONS

(1) Have suction available since vomiting may occur.
(2) Use an appropriate size airway adjunct with BVM.
(3) Use an appropriate size mask to avoid pressure over the eyes (pediatric patient), which may cause vagal stimulation.

(4) For single clinician BVM use the “E-C clamp” technique to achieve an adequate seal and avoid pressure on the soft tissues of the face or neck: Place the third, fourth, and fifth fingers along the jaw to provide a chin lift (forming an E); use the thumb and index finger to hold the mask on the child’s face (forming a C).

(5) If the patient does not have adequate chest rise and breath sounds with BVM, consider the following interventions:
(a) Use 2-hand jaw lift and oral airway to relieve tongue obstruction.
(b) Use a larger bag to increase the volume of air delivered into the patient.
(c) Evaluate and treat the patient for gastric distension. Clinicians may manually decompress the stomach and/or open an existing gastric tube or button and/or place NG or OG tube.

f) SUGGESTED SIZES FOR RESUSCITATION MASKS

<table>
<thead>
<tr>
<th>Age</th>
<th>Mask Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature infants</td>
<td>Neonatal</td>
</tr>
<tr>
<td>Newborn to 1 year</td>
<td>Infant</td>
</tr>
<tr>
<td>1–4 years</td>
<td>Toddler</td>
</tr>
<tr>
<td>5–12 years</td>
<td>Pediatric</td>
</tr>
<tr>
<td>Greater than 13 years</td>
<td>Small adult</td>
</tr>
<tr>
<td>Adult</td>
<td>Adult</td>
</tr>
</tbody>
</table>

g) SUGGESTED SIZES FOR RESUSCITATION BAGS

<table>
<thead>
<tr>
<th>Age</th>
<th>Bag Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infant to less than 1 year of age</td>
<td>Infant (450–500 mL)</td>
</tr>
<tr>
<td>Child 1-12 years</td>
<td>Pediatric (750 mL)</td>
</tr>
<tr>
<td>Adolescent/Adult</td>
<td>Adult (1,000–1,200 mL)</td>
</tr>
</tbody>
</table>
Procedures – AIRWAY MANAGEMENT:
CONTINUOUS POSITIVE AIRWAY PRESSURE (CPAP)

12.3

a) INDICATIONS

(1) Respiratory distress or failure, due to cardiogenic pulmonary edema or COPD/asthma, in which the patient demonstrates spontaneous respirations and a patent, self-maintained airway
(2) Patients who are 13 years of age or older
(3) Exception: EMT may transport a patient who is chronically on CPAP who is going for routine medical care and has in attendance a patient provided attendant who can manage the patient’s own CPAP.

CLINICIAN MUST ASSURE THAT THE CPAP MASK FITS THE PATIENT APPROPRIATELY.

b) CONTRAINDICATIONS

(1) Circumstances in which endotracheal intubation or a surgical airway is indicated to secure a patient airway
(2) Facial deformity or trauma which prevents the use of the device
(3) Patient has significantly decreased level of consciousness or an inability to protect their airway
(4) Patient has a tracheostomy
(5) Patient is vomiting
(6) Circumstances in which the patient does not improve or continues to deteriorate despite CPAP administration
(7) Known intolerance for noninvasive airway procedures (CPAP)

c) PROCEDURE

(1) Assure patent airway.
(2) Administer 100% O₂ via appropriate delivery system.
(3) Perform appropriate patient assessment, including obtaining vital signs, pulse oximeter (SpO₂) reading, and cardiac rhythm.
(4) Apply CPAP device per manufacturer’s instructions.
(5) Continuously reassess the patient.
(6) Monitor continuous pulse oximetry.
(7) Monitor continuous ETCO₂ with nasal prongs (if available).
(8) Follow the appropriate set of standing orders for continued treatment.
(9) Contact the medical control as soon as possible to allow for prompt availability of hospital CPAP equipment and respiratory personnel.

FOR CIRCUMSTANCES IN WHICH THE PATIENT DOES NOT IMPROVE OR CONTINUES TO DETERIORATE DESPITE CPAP AND/OR MEDICATIVE THERAPY, TERMINATE CPAP ADMINISTRATION AND PERFORM BVM VENTILATION AND ENDOTRACHEAL INTUBATION IF NECESSARY.

CPAP MAY BE CONSIDERED FOR NON-CARDIOGENIC PULMONARY EDEMA.
Procedures – AIRWAY MANAGEMENT: Extraglottic Airway (EGA) Devices

a) INDICATIONS

To provide ventilation and oxygenation to a patient with a compromised airway, including:

(1) Patient without a gag reflex and unable to maintain own airway
(2) Patients who cannot be intubated following the administration of paralytic medication
(3) Pediatric patients; EGA is preferred over endotracheal tube placement for pediatric patients

b) CONTRAINDICATIONS

(1) Responsive patients with an intact gag reflex
(2) Known esophageal disease or ingestion of caustic substances

c) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) The EGA does not protect against the effects of regurgitation and aspiration.
(2) High airway pressures may divert gas either to the stomach or to the atmosphere.
(3) A potential complication of the King LTS-D airway is inadvertent tracheal intubation. After placement, perform standard checks for breath sounds and utilize an appropriate carbon dioxide (ETCO₂) monitor.

d) PROCEDURE

King LTS-D

(1) Inspect all components of the LTS-D for visible damage.
(2) Select appropriately sized LTS-D airway as specified by manufacturer.
(3) Test cuffs by injecting the maximum volume of air (by size) as specified by manufacturer and lubricate with water-soluble jelly.
(4) Maintain cervical immobilization (if indicated) and lift tongue and jaw upward with one hand. Ideal position of the head is in the “sniffing position”; however, the LTS-D airway can be inserted with the head in neutral position.
(5) Insert LTS-D airway using a lateral approach and advance the tip behind the base of the tongue while rotating the tube back to midline so the blue line faces the patient’s chin.
(6) Without exerting excessive force, advance tube until base of connector is aligned with teeth and gums.
(7) Inflate cuff and ventilate patient. Gently withdraw the tube until ventilation becomes easy and free-flowing.
(8) Adjust cuff inflation to obtain a seal of the airway.
(9) Ventilate and evaluate for appropriate placement (bilateral lungs sounds, absence of gastric sounds, chest rise, ETCO₂ waveform/value, oxygen saturation).
(10) Once effective ventilation has been confirmed, secure the LTS-D and continue to monitor oxygen saturation and ventilate to desired ETCO₂ level.
**Procedures – AIRWAY MANAGEMENT: Extraglottic Airway (EGA) Devices (continued)**

**Supraglottic Airway (LMA®, Air-Q®, iGel®)**

1. Inspect all components of the supraglottic airway for damage.
2. Select appropriately sized supraglottic airway as per manufacturer’s specifications.
3. Lubricate with water-soluble jelly per manufacturer’s specifications.
4. Maintain cervical immobilization, if indicated, and lift tongue.
5. Inflate cuff, if applicable, per manufacturer specifications.
6. Ventilate and evaluate for appropriate placement (bilateral lungs sounds, absence of gastric sounds, chest rise, ETCO₂ waveform/value, oxygen saturation).
7. Adjust cuff inflation (if applicable) and position as needed to obtain a seal of the airway.
8. Once effective ventilation is confirmed, secure the supraglottic airway and continue to monitor oxygen saturation and ventilate to desired ETCO₂ level.

**Inability to Ventilate with EGA Devices**

If unable to achieve adequate ventilation using an EGA, remove the device and resume BVM ventilation with an NPA/OPA, then attempt placement again. If unable to place the EGA a second time, repeat the steps above and consider obstructed airway maneuvers, if not already performed, and refer to *Cricothyroidotomy* protocol.

**EMSOP Requirements**

An EMSOP must carry at least one type of extraglottic airway referenced in this protocol in sizes appropriate for all patients from newborn through adult.
a) PURPOSE

A naso/orogastric tube is passed to relieve the gastric distension or pressure in an effort to reduce the risk of aspiration and increase the intrathoracic volume.

b) INDICATIONS

(1) All pediatric intubated patients
(2) Pediatric patients with gastric distension complicating Bag Valve Mask Ventilation
(3) Intubated adult patients exhibiting signs and symptoms of gastric distension that compromise ventilation or circulation

c) CONTRAINDICATIONS

(1) History of esophageal varices
(2) Esophageal or gastric surgery within the past 6 weeks
(3) Anatomical deformity complicating nasal passage of the tube (nasogastric)
(4) Suspected basilar skull fracture

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) Tracheal intubation with gastric tube
(2) Epistaxis
(3) Coiling or knotting of tube in the stomach or esophagus
(4) Trauma to the nose, esophagus, or stomach
(5) Triggering vomiting
(6) Intracranial placement of gastric tube in patients with unidentified skull fractures

e) PRECAUTIONS

Have suction available since vomiting may be induced.
PARAMEDIC ONLY

a) PURPOSE

Nasal intubation is the technique of passing an endotracheal tube through the nose and pharynx into the trachea. This is done without using a laryngoscope to visualize the vocal cords (blind technique). The procedure is limited to breathing patients in whom oral intubation is difficult.

b) INDICATIONS

(1) Use is primarily for hypoxemic CHF and COPD patients and is allowed for closed head injury patients with clenched teeth
(2) An oxygen saturation of less than or equal to 90% in a patient on 100% oxygen by face mask and respiratory distress
(3) A respiratory rate of 8 or less per minute or 35 or greater per minute
(4) A Glasgow Coma Score of 8 or less, or
(5) Loss of gag reflex

c) CONTRAINDICATIONS

(1) Patient receiving anticoagulants, such as Coumadin (warfarin)
(2) Patient with upper airway hemorrhage, significant mid-facial trauma, or laryngeal trauma
(3) Patient with cerebral spinal fluid leakage or evidence of basilar skull fracture

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) Epistaxis
(2) Intubation of the esophagus
(3) Trauma to the oral pharynx, vocal cords, esophagus, or trachea
(4) Right mainstem bronchus intubation
(5) Vomiting
(6) Increased intracranial pressure, as result of increased vagal stimulation
(7) Pneumothorax/tension pneumothorax from high pressure ventilation or underlying preexisting trauma
(8) Intracranial tube placement through basal skull fracture

e) PRECAUTIONS

(1) Topical anesthesia (lidocaine 4% spray or gel) should be applied to both nares to minimize discomfort.
(2) Confirmation of ET placement
   (a) Utilization of the Beck Airway Airflow Monitor (BAAM) device when available
   (b) Auscultation of all lung fields to confirm air exchange
Procedures – AIRWAY MANAGEMENT:
NASOTRACHEAL INTUBATION (continued)

(c) Auscultation of the epigastrium to deny disturbance of gastric fluids upon ventilation
(d) Observation of bilateral expansion of the thorax
(e) ETCO₂ detection device required. At a minimum, use colorimetric devices.
(f) The esophageal detection device
(g) Documentation of tube depth at the nares
(h) Other clinical signs of improved perfusion and ventilation
   (e.g., pupillary response, skin color, etc.)

(3) Nasal intubation may require facilitation with sedation. When hypovolemia is unlikely and hypotension is not present, morphine/fentanyl or midazolam, or a combination of both, may be given by direct medical consultation to achieve mild sedation.
a) PURPOSE

Needle Decompression Thoracostomy is the procedure of introducing a needle/catheter with a minimum length of 3.25 inches and a minimum diameter of 14 gauge (with optional add-on flutter valve attached) into the pleural space of the chest to provide temporary relief for the patient suffering from a tension pneumothorax.

b) INDICATIONS

1) Patients who are assessed to have a life-threatening tension pneumothorax in extremis with absent lung sounds AND clear evidence of hemodynamic compromise to include hypotension (systolic blood pressure less than 100 mmHg), and/or arrest

2) If traumatic arrest is suspected due to multi-system blunt trauma, or due to penetrating neck, chest, or abdominal trauma, bilateral needle decompression should be performed. Once catheters are placed, do not remove.

3) Allowable site:
   (a) Adults and children 15 years of age and older:
      (i) Fifth (5th) intercostal space, anterior axillary line
      (ii) If fifth (5th) intercostal space site is not available, use second (2nd) intercostal space at the mid-clavicular line on the anterior chest wall
   (b) Children under 15 years of age:
      (i) Use the second (2nd) intercostal space, at the mid-clavicular line on anterior chest wall

c) CONTRAINDICATIONS

   Patients whose tension pneumothorax can be relieved by the removal of an occlusive dressing from an open chest wound

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

   1) Intercostal vascular or nerve injury
   2) Pneumo/hemothorax
   3) Direct damage to the lung
   4) Pericardial/cardiac injury
   5) Infection

e) PRECAUTIONS

   1) Reassessment of catheter patency
   2) Second decompression may need to be performed if reaccumulation, catheter occlusion, or dislocation is evident.
Procedures – AIRWAY MANAGEMENT: OBSTRUCTED AIRWAY
FOREIGN BODY REMOVAL: DIRECT LARYNGOSCOPY

a) PURPOSE

The attempted correction of a foreign-body airway obstruction through direct laryngoscopy should be accomplished only by a Maryland licensed CRT-(I) or paramedic. This is accomplished after the ALS clinician has determined (by noting repeated unsuccessful attempts at dislodging the object by applying the standard basic method of foreign body removal by BLS clinicians or the ALS clinician) that the object cannot be dislodged by these means. The patient must be unconscious and supine before this method is attempted.

b) INDICATIONS

Patient must be unconscious due to foreign body upper airway obstruction that has not resolved with standard basic methods for foreign body removal.

c) CONTRAINDICATIONS

None

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

Trauma to the oral pharynx, vocal cords, esophagus, or trachea

e) PRECAUTIONS

It is important to distinguish the foreign body from portions of the patient’s anatomy.
Procedures – AIRWAY MANAGEMENT:
OROTRACHEAL INTUBATION

a) PURPOSE

(1) Endotracheal intubation involves the passage of an endotracheal tube with direct visualization or digital manipulation through the larynx and into the trachea to provide direct maximum ventilatory support for a patient.
(2) Blind digital intubation is accomplished without the laryngoscope.

b) INDICATIONS

(1) Cardiac arrest
(2) Respiratory arrest, patient without gag reflex
(3) Deep coma, patient without gag reflex
(4) Patient in extremis, in severe respiratory distress with extremely poor air exchange, or agonal respirations (gag reflex may be present)

c) CONTRAINDICATIONS

Upper airway obstruction due to foreign objects

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) Intubation of the esophagus
(2) Trauma to the oral pharynx, vocal cords, esophagus, or trachea
(3) Right mainstem bronchus intubation
(4) Vomiting
(5) Increased intracranial pressure as a result of increased vagal stimulation
(6) Pneumothorax/ tension pneumothorax from high pressure ventilation or underlying preexisting trauma

e) PRECAUTIONS

(1) When the patient cannot be intubated (following no more than two tracheal intubation attempts), avoid future intubation attempts until the patient reaches the hospital, unless otherwise directed by the physician.
(2) Confirmation of ET placement
As it has been determined that no single method of assessment is 100% reliable, the position of the endotracheal tube must be assessed to be properly in the trachea by all means available to the EMS clinician. The following methods may be used to confirm proper placement of the endotracheal tube:
(a) Visualization of the ET tube protruding adequately past the vocal cords and into the trachea
(b) Auscultation of all lung fields to confirm adequate air exchange
(c) Auscultation of the epigastrium to deny disturbance of the gastric fluids upon ventilation
(d) Observation of the bilateral expansion of the thorax
Procedures – AIRWAY MANAGEMENT: OROTRACHEAL INTUBATION (continued)

(e) ETCO$_2$ detection device. At a minimum, utilize colorimetric devices (required for all intubated patients).
(f) The esophageal detection device
(g) Documentation of tube depth at the lip
(h) Other clinical signs of improved perfusion and ventilation (e.g., pupillary response, skin color, etc.)

3. Once initial placement is confirmed:
(a) The tube must be adequately secured
(b) The patient must be prepared for transport in such a fashion as to minimize movement of the head and neck.

4. Placement of the tube should be verified by all means possible (as in (2) above) and as often as possible as part of the clinicians’ ongoing assessments. It has been further noted that flexion of the neck can cause 3–5 cm displacement of the ET tube dislodging the tube from the trachea. At a minimum this reconfirmation should occur:
(a) Once the patient is prepared for transport,
(b) Anytime the patient is moved,
(c) Anytime dislodgment of the tube is suspected, and
(d) When responsibility for care is transferred to any other clinician.

5. During routine reporting procedures, documentation of proper placement should include which methods were utilized and at which points, in the care of the patient, verification was accomplished.

6. Maintain neutral alignment of head and neck with cervical stabilization when intubating trauma patients.

7. The Blind Digital method may be utilized for intubation of a patient in whom hyper-extension of the cervical spine may be contraindicated. It may also benefit patients with severe facial trauma. However, it must be emphasized that this can be a difficult procedure, and the clinician must be certain that the patient cannot bite.
### Equipment Sizes

<table>
<thead>
<tr>
<th>Age</th>
<th>Oral Airway</th>
<th>Bag-Valve-Mask</th>
<th>ETT Size</th>
<th>ETT Blade</th>
<th>Suction Catheter</th>
<th>Gastric Tube</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature</td>
<td>0</td>
<td>Neonatal</td>
<td>2.5–3.0</td>
<td>0</td>
<td>6F</td>
<td>5F</td>
</tr>
<tr>
<td>Newborn</td>
<td>0</td>
<td>Neonatal</td>
<td>3.0–3.5</td>
<td>0–1</td>
<td>6F</td>
<td>5–8F</td>
</tr>
<tr>
<td>3 mo.</td>
<td>1</td>
<td>Infant</td>
<td>3.5</td>
<td>1</td>
<td>6–8F</td>
<td>5–8F</td>
</tr>
<tr>
<td>6 mo.</td>
<td>1</td>
<td>Infant</td>
<td>3.5–4.0</td>
<td>1</td>
<td>8F</td>
<td>8F</td>
</tr>
<tr>
<td>1 yr.</td>
<td>1</td>
<td>Infant</td>
<td>4.0</td>
<td>1</td>
<td>8F</td>
<td>8F</td>
</tr>
<tr>
<td>2 yrs.</td>
<td>2</td>
<td>Child</td>
<td>4.0–4.5</td>
<td>1–2</td>
<td>8–10F</td>
<td>8–10F</td>
</tr>
<tr>
<td>3 yrs.</td>
<td>2</td>
<td>Child</td>
<td>4.5</td>
<td>2</td>
<td>10F</td>
<td>10F</td>
</tr>
<tr>
<td>4 yrs.</td>
<td>3</td>
<td>Child</td>
<td>4.5–5.0</td>
<td>2</td>
<td>10F</td>
<td>10–12F</td>
</tr>
<tr>
<td>6 yrs.</td>
<td>4</td>
<td>Child</td>
<td>5.0–5.5</td>
<td>2</td>
<td>10F</td>
<td>12–14F</td>
</tr>
<tr>
<td>8 yrs.</td>
<td>4</td>
<td>Child</td>
<td>5.5–6.0</td>
<td>2</td>
<td>10–12F</td>
<td>14F</td>
</tr>
<tr>
<td>10 yrs.</td>
<td>5</td>
<td>Child</td>
<td>5.5–6.5</td>
<td>3</td>
<td>12F</td>
<td>14F</td>
</tr>
<tr>
<td>12 yrs.</td>
<td>5</td>
<td>Adult Small</td>
<td>6.5–7.0</td>
<td>3</td>
<td>12F</td>
<td>14–18F</td>
</tr>
<tr>
<td>14 yrs.</td>
<td>5</td>
<td>Adult</td>
<td>6.5–7.5</td>
<td>3</td>
<td>12–14F</td>
<td>16–18F</td>
</tr>
<tr>
<td>Adult</td>
<td>5</td>
<td>Adult</td>
<td>7.0–10.0</td>
<td>4</td>
<td>12–14F</td>
<td>16–18F</td>
</tr>
</tbody>
</table>

**Alert:**

Endotracheal tube selection for a child should be based on 16 plus child’s age divided by four \((16 + \text{ Year}) \div 4 = \text{Tube Size}\) or size recommended by length-based resuscitation tape (e.g., Broslow tape).

Age in the chart is a quick reference. Given individual variations in airway size, may consider one tube size larger and one tube size smaller. Use a length-based tape if available.
a) PURPOSE
Changing a tracheostomy tube may be required to reestablish a patent airway in patients who present with respiratory distress secondary to tracheostomy tube occlusion or obstruction that has not been relieved through suctioning.

b) INDICATIONS
(1) Inability to ventilate with BVM
(2) Ineffective spontaneous ventilations (poor chest rise, decreased breath sounds bilaterally)
(3) Hypoxia, cyanosis, or decreased O₂ saturation levels, not relieved by suctioning
(4) Increased work of breathing
(5) Altered mental status secondary to hypoxia

c) CONTRAINDICATIONS
None

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
(1) Inability to reinsert a tracheostomy tube
(2) Edema at stoma site
(3) Inability to maintain adequate chest rise and fall with assisted ventilations due to air leak around uncuffed tracheostomy tube

Patients greater than eight years of age who require assisted ventilations will need to have a cuffed tube inserted to prevent air leak around the tube and ensure adequate chest rise. If an appropriate sized cuffed tracheostomy tube is not available, then ALS clinicians may use an ET tube.

e) PROCEDURE
(1) Two clinicians or clinician and trained family member
(2) Use latex-safe sterile gloves and equipment.
(3) Position patient with the head and neck hyperextended to expose the tracheostomy site.
(4) Explain procedure to patient/family.
(5) Have new tracheostomy tube nearby.
(6) To remove the tracheostomy tube:
   (a) If a double cannula tracheostomy tube is in place, attempt to change inner cannula first and reassess the patient to see if the obstruction is relieved. If the patient continues to have respiratory distress, change the entire tracheostomy tube. If cuffed, deflate using a 10 mL syringe.
   (b) Carefully cut the tracheostomy ties.
   (c) Remove the tracheostomy tube, outward and backward towards the chest.
   (d) Lubricate the new tracheostomy tube with lubricating jelly or saline/water.
   (e) Insert new tracheostomy tube into stoma, inward and downward towards the lungs.
   **NOTE: STOP IF YOU MEET RESISTANCE (see (7) below).**
(f) If cuffed tracheostomy tube is used, once the tube has been inserted, inflate the cuff with an appropriate amount of air to avoid air leak around the tube (1–3 mL for pediatric tubes and 5–10 mL for adult tubes).

(g) Reassess the patient.

(h) With good chest rise and fall and improved skin color, secure the tracheostomy tube with ties or Velcro® at the back of the neck, so only one fingertip fits between the neck and the ties.

(7) If you meet resistance inserting the tracheostomy tube, do NOT force the tube into the stoma. Request ALS rendezvous, if appropriate. Assess the patient:

(a) Reposition the patient, hyperextend the neck area.

(b) Reoxygenate using BVM to stoma site, with infant mask and appropriate size reservoir bag for the patient’s size. Assess for chest rise and fall.

(c) If inadequate rise and fall of the chest, AND the patient has not had a laryngectomy, attempt BVM orally while placing an occlusive dressing over the stoma site. If a laryngectomy patient, you will only be able to ventilate with BVM at the stoma site.

(d) Attempt to insert a half-size smaller tracheostomy tube after lubricating with lubricating jelly or saline/water.

(e) Proceed with (6) f-g-h above.

(f) If you meet resistance, reassess the patient. Reoxygenate as needed.

(g) Insert a suction catheter through the tracheostomy tube, and use the suction catheter as a guide to insert the tracheostomy tube.

(h) Proceed with (6) f-g-h above.

(i) If ALS, attempt to insert a similar sized endotracheal tube into the stoma. If cuffed endotracheal tube is used, inflate the cuff with an appropriate amount of air to avoid air leak around the tube (1–3 mL for pediatric tubes and 5–10 mL for adult tubes).

(j) If ALS and unable to insert the ET tube into the stoma, AND the patient has not had a laryngectomy, attempt to intubate orally and apply an occlusive dressing over the stoma site.

(k) If you continue to have problems, STOP, consult the Base Station and continue BVM ventilations orally, or BVM to tracheostomy site ventilations if a laryngectomy patient, while en route to the closest appropriate hospital.
Procedures – AIRWAY MANAGEMENT: TRACHEOSTOMY SUCTIONING

a) PURPOSE
Tracheostomy suctioning may be required to maintain a patent airway in patients who present with respiratory distress secondary to tracheostomy tube occlusion or obstruction.

b) INDICATIONS
(1) Increased secretions from tracheostomy site or a mucous plug
(2) Hypoxia, cyanosis, or decreased oxygen saturation levels
(3) Increased work of breathing
(4) Altered mental status secondary to hypoxia

c) CONTRAINDICATIONS
None

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
(1) Bleeding at tracheal stoma site
(2) Dislodgment of tracheostomy tube
(3) Exaggerated cough reflex with introduction of saline
(4) Increased hypoxia/respiratory distress
(5) Infection

e) PROCEDURE
(1) Two clinicians or clinician and trained family member
(2) Use latex-safe sterile gloves and equipment.
(3) Position patient with the head and neck hyperextended to expose the tracheostomy site.
(4) Pre-oxygenate patient at the tracheostomy site:
   (a) NRB mask if patient has adequate effective spontaneous respirations
   (b) BVM if ventilator-dependent or there are ineffective spontaneous respirations
(5) Select appropriately sized suction catheter (2 x internal diameter of tracheostomy tube).
(6) Insert suction catheter:
   (a) Measure from the tracheostomy site to the sternal notch. OR
   (b) Insert until there is a cough reflex.
(7) Apply suction ONLY as the catheter is withdrawn, rotating the catheter in a twisting motion between thumb and finger.
(8) Suction for maximum of 10 seconds.
(9) Reoxygenate and reevaluate patient.
(10) Repeat suction procedure as needed (for thick secretions instill 3–5 cc sterile saline/water prior to repeat suctioning).
Procedures – AIRWAY MANAGEMENT: VENTILATORY DIFFICULTY SECONDARY TO BUCKING OR COMBATIVENESS

12.12

a) **INDICATIONS**

Patients successfully intubated with an endotracheal tube, an approved alternative airway device, or cricothyroidotomy, for whom the ability to provide manual or mechanical ventilation is impaired secondary to bucking or combativeness.

b) **CONTRAINDICATIONS**

Unsecured airway.

c) **PROCEDURE**

1. Ventilatory difficulty secondary to bucking or combativeness in intubated patients. Ketamine may be preferred for patients who have hypotension or possible hypovolemia, or if ventilatory difficulty is thought to be the result of pain response.
   
   (i) Dose: Administer 2 mg/kg IVP/IO over 60 seconds. May repeat 2 additional doses of 1 mg/kg for IVP/IO every 10-15 minutes to a total of 3 doses as needed. Additional doses require medical consultation.

2. Midazolam up to 0.1 mg/kg IVP/IO/(over 1-2 minutes)/IM, titrated to abate bucking and relax ventilation while maintaining systolic blood pressure greater than 90 mmHg (110 mmHg if injuries include a suspected head injury). Maximum single dose is 5mg.

3. If ventilatory difficulty is thought to be the result of pain response, opioid may be used per Pain Management protocol in addition to or instead of midazolam: Titrate to abate bucking and relax ventilation while maintaining systolic BP greater than 90 mmHg.

4. Continue to monitor oxygen saturation and ventilate to desired ETCO₂ level.

5. Obtain on-line medical direction if further problems present.

6. Midazolam up to 0.05 mg/kg IVP over 1–2 minutes, titrated to abate bucking and relax ventilation while maintaining systolic BP: greater than 60 in neonates, 70 in infants, and [70 + (2 x years) = systolic BP] for patients greater than 1 year of age. Maximum single dose is 5 mg.

7. If ventilatory difficulty is thought to be the result of pain response, opioid may be used per Pain Management protocol in addition to or instead of midazolam: Titrate to abate bucking and relax ventilation while maintaining systolic BP: greater than 60 in neonates, 70 in infants, and [70 + (2 x years) = systolic BP] for patients greater than 1 year of age.

8. Continue to monitor oxygen saturation and ventilate to desired ETCO₂ level.

9. Obtain on-line medical direction if further problems present.


**Procedures – AIRWAY MANAGEMENT:**

**VENTILATORY MANAGEMENT**

**a) PURPOSE**

(1) Manual ventilation using a bag-valve-mask (BVM) or mechanical (machine) ventilation can be an effective method for managing a patient in the prehospital environment when performed correctly. Ventilatory management is important at both the BLS and ALS levels.

(2) Special considerations such as etiology of respiratory failure and method of achieved airway management, including intubation (e.g., rapid sequence intubation), may require the advanced life support clinician to provide additional care.

**b) INDICATIONS**

(1) Any condition requiring assisted or artificial ventilation with a bag-valve-mask or mechanical (machine) ventilation

(2) All patients will require manual ventilation after the placement of an advanced airway. Inadequate respiratory rate may be secondary to underlying respiratory pathology or the result of pharmacologic intervention secondary to medications used in rapid sequence intubation.

**c) CONTRAINDICATIONS**

None

**d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS**

(1) Gastric distension, vomiting, and/or aspiration

(2) Hypoxemia

(3) Secretions and tube/bag obstruction

(4) Barotrauma

(5) Patient agitation

(6) Equipment failure

**e) PROCEDURE/PRECAUTIONS:**

(1) Have suction available and ensure a patent airway using a BLS airway adjunct (OPA or NPA).

(2) Rate of *initial* ventilation by single hand bag-valve technique should generally be the following:

   (a) For all ages except neonates, 1 breath every 5 seconds
       (8–12 breaths/min)

   (b) For a neonate, 1 breath every 3 seconds (higher rates may be required)

(3) AVOID hyperventilating unless patient exhibits signs of brainstem herniation (e.g., unequal pupils, posturing). Hyperventilation is associated with increased mortality.

(4) In the absence of contraindications (e.g., CPR or spinal trauma), consider elevating the head of the bed to 30 degrees.

(5) Continuous pulse oximetry shall be used. If a sudden drop in SpO₂ is observed, assess airway patency and consider obstruction (e.g., tongue, vomitus, blood), poor seal around BVM, and flow of oxygen being administered (LPM).
(6) A gastric tube should be considered for gastric decompression whenever distention is caused by BVM ventilation. Gastric distention can reduce effectiveness of ventilations.

(7) Waveform capnography and patient-specific considerations:
   (a) Continuous ETCO₂ shall be used whenever an advanced airway has been placed.
   (b) Continuous ETCO₂ monitoring is encouraged for all other manually-ventilated patients.
   (c) The waveform shape and reading can contribute to an understanding of the underlying pathology.
   (d) Waveform capnography is utilized to optimize manual ventilation. Deliver ventilations to achieve a target ETCO₂ level of 35–40 mmHg if patient has a pulse.
   (e) ETCO₂ can be used to assess trends during a cardiac arrest and may contribute to understanding the pathology. A sudden substantial increase in ETCO₂ may indicate ROSC.
   (f) Hypercapnia is seen in patients experiencing respiratory failure as a result of obstructive disease, such as asthma and COPD. Chronic baseline hypercapnia should be considered when ventilating to a target ETCO₂.
   (g) A target ETCO₂ of 30–35 mmHg should be used for the rare patient who exhibits signs of brainstem herniation. Lower ETCO₂ has been associated with increased mortality.

(8) If advanced airway is placed and patient does not have adequate chest rise, absent or significantly diminished breath sounds, or decreased SpO₂ or abnormal ETCO₂ levels, consider the DOPES mnemonic:
   “D”: Is the tube displaced? Assess for bilateral breath sounds and reassess tube depth and compare to initial depth noted after insertion.
   “O”: Is an obstruction present? Suction the tube with a flexible suction catheter.
   “P”: Are there signs of a tension pneumothorax? If present, perform needle decompression thoracostomy.
   “E”: Is there an equipment malfunction? Check oxygen flow in tubing and level in portable cylinder, determine whether SpO₂ and ETCO₂ devices are working correctly, and ensure the cuff is adequately inflated.
   “S”: If history of asthma or COPD is known, consider extending the interval between ventilations to avoid stacked ventilations.

(9) Consider using a positive end expiratory pressure (PEEP) valve on the BVM, especially if the patient is hypoxemic (start at 5 cm H₂O).

(10) If combativeness or bucking prevents the delivery of adequate ventilations, management shall be guided by the Ventilatory Difficulty Secondary to Bucking protocol.
Procedures – ELECTRICAL THERAPY: AUTOMATED EXTERNAL DEFIBRILLATION (AED)

a) INDICATIONS

Sudden cardiac arrest (patients with no pulse and not breathing).

| Neonate (1 hour to 28 days of life) to less than 1 year of age | Manual defibrillator preferred. (If unavailable, an AED with pediatric capability is preferred over an adult AED.) |
| 1 year of age to 8 years of age | AED with pediatric capability, using the pediatric capability, is preferred over an adult AED. |
| Child 8 years of age or greater | Adult AED |

b) CONTRAINDICATIONS

Patient exhibiting signs of life
Newly born patients (up to one hour after birth)

USE OF THE AED IN THE MANUAL MODE IS RESERVED FOR ALS.

c) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) Burns to skin
(2) Deactivation of patient’s implanted pacemaker
(3) Injury to patient, self, and/or bystanders

d) PRECAUTIONS

(1) Make sure the patient and the environment are dry.
(2) Avoid placing pads over cardiac pacemakers/defibrillators or nitroglycerin patches.
(3) DO NOT touch the patient while the AED is analyzing the patient or discharging energy.
(4) ENSURE that no one is touching the patient when the shock button is pushed.
(5) Never defibrillate while moving the patient or when in a moving ambulance.

e) PROCEDURE

(1) Initiate analysis of rhythm.
(2) If shock is indicated:
   (a) Ensure all individuals are clear of the patient.
   (b) Initiate shock to the patient.
   (c) Immediately perform 5 cycles of CPR between shocks, then initiate analysis of rhythm.
   (d) If patient remains pulseless, continue this cycle of CPR and shocks until the AED prompt states “no shock advised,” or ROSC is achieved or ALS arrives or the patient is transported or the Termination of Resuscitation protocol is initiated.
(3) If shock is not indicated and the patient remains in cardiac arrest:
   (a) Perform 5 cycles of CPR.
   (b) Initiate analysis of rhythm.
   (c) If shock is indicated, see “If shock is indicated” section above.
(d) If shock is not indicated, continue CPR until ALS arrives or ROSC is achieved or the patient is transported or the *Termination of Resuscitation* protocol is initiated.

(4) If shock is not indicated and patient regains pulse, treat per *Return of Spontaneous Circulation* (ROSC) protocol.

f) **SPECIFIC DOCUMENTATION**

(1) Document the number of analyses and shocks delivered, times of assessments and treatments, and the patient’s response to shocks/CPR. Specify the type of AED, location of AED, bystander and clinician contact, and the triggering event.

(2) If using an AED with EKG strip recorder, generate 2 recordings.

(3) Give one to the ALS clinician or hospital and attach the other to your patient care report.

(4) Record the name of the contact for accessing AED data download summary.

(5) Consider bringing the AED to the hospital for downloading.
Procedures – ELECTRICAL THERAPY: 
CARDIOVERSION

a) INDICATIONS

Hemodynamically unstable with life-threatening, rate-related signs and symptoms including hypotension, acutely altered mental status, signs of shock, ischemic chest discomfort/AMI, or acute heart failure.

b) DOSAGE

(1) Adult
   (a) For SVT or VT or atrial flutter:
      (i) Initial 100 J, or follow manufacturer's recommendations
      (ii) Subsequent 200 J, 300 J, 360 J, or follow manufacturer's recommendations
   (b) For atrial fibrillation:
      (i) Initial 200 J, or follow manufacturer’s recommendations
      (ii) Subsequent 300 J, 360 J, or follow manufacturer's recommendations

(2) Pediatric

Symptomatic tachydysrhythmias
   (a) Initial 0.5 J/kg; if the calculated joules setting is lower than the defibrillation device is able to deliver, use the lowest joules setting possible or obtain medical consultation.
   (b) Subsequent 1 J/kg; repeat at 2 J/kg

(3) If the patient exhibits ventricular fibrillation following emergency cardioversion, immediately turn off the synchronizer and defibrillate with appropriate delivered energy (200 to 360 J for adults and 2 to 4 J/kg for pediatric patients) and refer to defibrillation and/or other appropriate protocol.

c) CONTRAINDICATIONS

Tachydysrhythmias due to digitalis toxicity

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

An unsynchronized shock can result in ventricular fibrillation.
e) PRECAUTIONS

(1) If the calculated joules setting is lower than the cardioversion device is able to deliver, use the lowest joules setting possible or obtain medical consultation.

(2) Pre-procedural sedation or analgesia
   (a) Patient may experience moderate to severe discomfort during cardioversion. Consider pre-medication by administering opioid per Pain Management protocol.
      OR
   (b) Administer midazolam 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes per increment, with maximum single dose 5 mg. (Reduce by 50% for patients 69 years or older.)

(3) Pre-procedural sedation or analgesia
   (a) Patient may experience moderate to severe discomfort during cardioversion. Consider pre-medication by administering opioid per Pain Management protocol.
      OR
   (b) Administer midazolam 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes per increment, with maximum single dose 5 mg.
a) PURPOSE

Defibrillation involves the delivery of non-synchronized direct electric current (monophasic or biphasic) to the myocardium of a patient exhibiting ventricular fibrillation or ventricular tachycardia without palpable pulses/blood pressure. The objective of defibrillation is to depolarize the entire myocardium, which, it is hoped, will result in allowing a single reliable pacemaker site to assume pacemaker control at a rate capable of producing an adequate cardiac output.

b) INDICATIONS

(1) Ventricular fibrillation
(2) Ventricular tachycardia without palpable pulse or BP

c) DOSAGE

(1) Adult
   (a) Initial delivered energy monophasic 360 J or biphasic 120–200 J
   (b) Subsequent delivered energy monophasic 360 J or biphasic increasing joules setting, if device allows

(2) Pediatric
   (a) Initial delivered energy 2 J/kg (monophasic or biphasic)
   (b) Subsequent delivered energy at increasing dosage: 4 J/kg, 6 J/kg, 8 J/kg, to a maximum of 10 J/kg.

d) CONTRAINDICATIONS

None

e) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) Burns to the skin
(2) Deactivation of patient’s implanted pacemaker

f) PRECAUTIONS

(1) Patients who are fully digitalized may require less than the normal recommended delivered energy.
(2) If the calculated joules setting is lower than the defibrillation device is able to deliver, use the lowest joules setting possible or obtain medical consultation.
a) PURPOSE

Non-invasive cardiac pacing, also referred to as external or transcutaneous pacing, involves the temporary application of externally applied electrodes to deliver an adjustable electrical impulse directly across an intact chest wall for the purpose of rhythmically stimulating the myocardium to increase the mechanical heart rate.

b) INDICATIONS

(1) It is indicated for the treatment of hemodynamically compromised patients in settings where cardiac output is compromised due either to the complete failure of cardiac rhythm or to an insufficient rate of the patient's intrinsic pacemaker.

(2) Bradycardia (EKG other than second-degree Mobitz Type II or third-degree AV Block)

(3) Second-degree Mobitz Type II and third-degree AV block with a systolic BP of less than 80 mmHg, or 80–100 mmHg with shock-like signs or symptoms. In the presence of Mobitz II and third-degree AV block, medical consultation is required for atropine administration.

(4) Pacing may be indicated in certain instances in which the heart rate is 60–75 BPM and shock-like symptoms persist. Pacing in these instances requires medical consultation from a physician.

(5) Pediatric patients with profound symptomatic bradycardia unresponsive to optimal airway management, oxygenation, epinephrine, and atropine.

c) DOSAGE

Start pacemaker at age appropriate heart rate:
Infant (less than 1 year): 120 beats per minute
Child (1 through 12 years): 100 beats per minute
Adult/Adolescent (13 years and greater): 80 beats per minute

Start milliamperes (m.a.) as low as possible and gradually increase m.a. until palpable pulse to confirm capture or 200 m.a.

CONTINUE CHEST COMPRESSIONS FOR PEDIATRIC PATIENTS WHO REMAIN POORLY PERFUSED DESPITE PACEMAKER CAPTURE.
d) CONTRAINDICATIONS

(1) Non-witnessed cardiopulmonary arrest with asystole
(2) Patient not meeting blood pressure criteria

e) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

(1) Patient may experience moderate to severe discomfort during pacing. Consider pre-medication by administering opioid per Pain Management protocol.
   (a) Administer opioid per Pain Management protocol.
   OR
   (b) Administer midazolam 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes per increment, with maximum single dose 5 mg. (Reduce by 50% for patients 69 years or older.)

(2) Patient may experience moderate to severe discomfort during pacing. Consider pre-medication by administering opioid per Pain Management protocol.
   (a) Administer opioid per Pain Management protocol.
   OR
   (b) Administer midazolam 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes per increment, with maximum single dose 5 mg.

f) PRECAUTIONS

When properly applied, chest compressions can be performed directly over the insulated electrodes while the pacer is operating.
a) PURPOSE
The University of Maryland Medical System, R Adams Cowley Shock Trauma Center (STC) maintains a deployable advanced surgical team (Go-Team) that includes an attending physician with surgical skills and an anesthetist capable of assisting EMS clinicians with the care of seriously injured patients when extrication times are anticipated to be more than 1 hour. On-scene incident commanders may request the Go-Team by contacting SYSCOM.

b) INDICATIONS
The on-scene incident commander may contact SYSCOM and request the Go-Team for seriously injured patients with potentially life or limb threatening injuries when extrication times are anticipated to be more than 1 hour and who may require advanced resuscitative or surgical services that are beyond the scope of prehospital emergency services.
Examples include:
(1) During a prolonged extrication, assist rescue personnel with planning the type and pace of the rescue by assessing the extent of injury and determine potential consequences that delays in time to definitive care might have on patient outcome.
(2) A patient trapped in heavy machinery requiring anesthesia/pain management to perform extrication
(3) A patient surviving a building collapse requiring an amputation to enable extrication
(4) A patient with a prolonged extrication requiring advanced fluid resuscitation including the administration of blood products
(5) Insertion of chest tubes or gastric and urinary catheters during the course of prolonged extrication

c) PROCEDURE
(1) On-scene incident commander will request the Go-Team by contacting SYSCOM. SYSCOM will coordinate the Go-Team’s transport to and from the scene with Maryland Express Care.
(2) If the Go-Team is dispatched by air, then SYSCOM will notify the Go-Team when the aircraft is landing on the STC helipad. If the Go-Team is dispatched by land, then Maryland Express Care will coordinate the Team’s response.
(3) Prior to the Go-Team’s departure to the scene, SYSCOM will notify the on-scene incident commander for the Go-Team’s ETA and reconfirm the need for the Go-Team.
(4) If the Go-Team is dispatched, the EMS medical commander will contact them using the “Trauma Line” (or other radio) to update them about the circumstances of the entrapment and the patient’s condition.
(5) When the Go-Team arrives on the scene, they are to report to the on-scene incident commander and operate within the Incident Command System.
(6) Once the patient is extricated, the EMS system will transport the patient to the appropriate facility under established EMS guidelines with consultation by the Go-Team physician.
(7) The Go-Team will document the care they provide and file a patient care report with the State EMS Medical Director at MIEMSS.
a) PURPOSE

The external jugular vein is a large vessel in the neck that may be used by a CRT-(I) or paramedic for intravenous cannulation.

b) INDICATIONS

EJs are appropriate when IV access is emergently indicated, but an extremity vein cannot be catheterized.

c) CONTRAINDICATIONS

(1) Inability to visualize the vein
(2) Suspected spinal trauma

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS

Hematoma, pain, infiltration, infection, dislodged catheter, nerve injury, thrombosis, air embolism, airway occlusion, and pneumothorax

e) PRECAUTIONS

Carefully secure EJ catheter and tubing.
a) PURPOSE
To improve survival of sudden out-of-hospital cardiac arrest patients in Maryland. High Performance Cardio-Pulmonary Resuscitation (HPCPR) employed with Code Resource Management (CRM) is a proven concept based on a team approach that ensures effective and efficient use of EMS resources. This systematic change in treatment and management of cardiac arrest patients has demonstrated effectiveness in Maryland, and provides an example for systems embarking on measuring and improving care that is based upon proven research and practices.

b) INDICATIONS
Patients in cardiac arrest who are greater than 24 hours old.

c) CONTRAINDICATIONS
(1) Patients meeting the criteria for Pronouncement of Death in the Field protocol
(2) Patients who are less than 24 hours old

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
None

e) PRECAUTIONS
None

f) IMPORTANT ROLE OF DISPATCHER TELEPHONE CPR (T-CPR)
(1) Immediate recognition of unresponsiveness, activation of EMS system response via 9-1-1, and initiation of CPR by the lay rescuer is essential to maximize survival.
(2) In an unresponsive patient, rapid recognition of agonal (gasing) respirations, or no respirations should prompt dispatcher-directed compressions to the caller (Dispatch-directed T-CPR).
(3) Dispatch-directed T-CPR delivers CPR prior to EMS system arrival and presents a patient more responsive to EMS interventions, thus providing the ability to improve survival.

g) PROCEDURE FOR HIGH PERFORMANCE CPR
(1) The first clinician at the patient’s side will assess and initiate compressions.
(2) Effective Compressions - Manual chest compressions should be initiated immediately upon identification of cardiac arrest, as long as the scene is safe. When compressions are done manually, compressors should be rotated every 2 minutes in order to maintain high-quality compressions. Ideally, one compressor is on each side of the patient’s chest: one person compressing the chest and the other person ready to start. Chest compressions will be performed at a depth of at least 2 inches allowing for complete recoil of the chest after each compression.
   For patients less than one year of age, compressions will be performed at a depth of 1 ½ inches. For patients greater than one year old up to age 13, compressions will be at a depth of 2 inches.
(3) Compressions should be accomplished with equal time given for the down and up motion and achieve a rate of 100–120 per minute.
Procedures – HIGH PERFORMANCE CPR (continued)

(4) **Continuous Compressions** - Chest compressions will be performed at a rate of 100–120 per minute and will NOT be interrupted during the two-minute cycle for any reason. Other treatments such as ventilations, IV access, or intubation attempts will be done while compressions are ongoing. After completion of a two-minute cycle, a brief pause to assess pulses and/or defibrillate will be limited to less than 10 seconds.

(5) **Defibrillation** – placement of the defibrillator pads will not interrupt chest compressions

(a) **Automatic External Defibrillation**
   
   The AED will be powered on as soon as the cardiac arrest is confirmed. Do not interrupt chest compressions to remove clothing or place defibrillation pads. If the AED charges after analyzing, chest compressions will be performed while the device charges, then the patient will be “cleared” and defibrillated. Compressors will hover over the patient with hands ready during defibrillation so compressions can start immediately after a shock. Another two-minute cycle of compressions will be immediately performed. Pulse checks will not occur after a shock, but only after the AED prompts “no shock advised.” If no pulse is palpated, or if unsure, immediately perform another two minutes of CPR.

(b) **Cardiac Monitor/Defibrillator**
   
   When a manual defibrillator is in use, it will be charged to the appropriate energy level as the end of the compression cycle nears (approximately 1 minute and 45 seconds into a two-minute cycle). At the end of the two-minute cycle, the patient will be cleared, the rhythm will then be interpreted rapidly, and the patient will either be defibrillated or the defibrillator energy charge will be cancelled. This sequence must be performed within 10 seconds. During this sequence, the compressors will hover over the patient with hands ready. If a shock is delivered, the compressor will immediately resume CPR. Rhythm interpretation will not occur after a shock, but only occur after the two-minute cycle of CPR is performed. If a shock is not indicated, check for a pulse. If patient remains pulseless, immediately resume HPCPR.

(6) **Ventilations** - Ventilations will be performed without stopping chest compressions. Ventilations are important but can impede the cardiac output from compressions. Thus, rescuers should not provide too many breaths or use excessive force. One ventilation will be given every 10th compression during recoil (upstroke). Once an advanced airway is in place, ventilations will be interposed asynchronously with uninterrupted compressions (1 ventilation every 6 seconds, for all ages). Ventilation volume should be low volume (approximately 500 cc), best approximated by a three finger or end of bag squeeze. High performance, continuous compressions remain the priority. Ensure ventilations are adequate with bag-valve-mask attached to 100% oxygen. Clinicians will not interrupt compressions to obtain an advanced airway.

For children **up to age 13**, maintain a ratio of 2 ventilations every 30th compression for single rescuer CPR or 2 ventilations every 15th compression for two or more rescuer CPR (one ventilation on the recoil of the 14th compression and one ventilation on the recoil of the 15th compression).
Procedures –
HIGH PERFORMANCE CPR (continued)

<table>
<thead>
<tr>
<th>Rescuers Should</th>
<th>Rescuers Should Not</th>
</tr>
</thead>
<tbody>
<tr>
<td>Perform chest compressions at a rate of 100-120/min</td>
<td>Compress at a rate slower than 100/min or faster than 120/min</td>
</tr>
<tr>
<td>Compress to a depth of at least 2 inches (5 cm)</td>
<td>Compress to a depth of less than 2 inches (5 cm) or greater than 2.4 inches (6 cm)</td>
</tr>
<tr>
<td>Allow full recoil after each compression</td>
<td>Lean on the chest between compressions</td>
</tr>
<tr>
<td>Minimize pauses in compressions</td>
<td>Interrupt compressions for greater than 10 seconds</td>
</tr>
<tr>
<td>Ventilate adequately (2 breaths after 30 compressions, each breath delivered over 1 second, each causing chest rise)</td>
<td>Provide excessive ventilation (ie, too many breaths or breaths with excessive force)</td>
</tr>
</tbody>
</table>

(7) **Advanced Life Support** - ALS clinicians will address defibrillation, IV/IO access, medication administration, and advanced airway placement, as indicated within these protocols; however, the placement of an advanced airway is no longer an early focus of cardiac arrest management and will not interrupt chest compressions. Nasal capnography may be utilized to optimize CPR performance and evaluation of ROSC, with use of bag-valve-mask ventilation.

(8) **Return of Spontaneous Circulation (ROSC)** – Refer to ROSC protocol.

(9) **Quality Improvement/Performance Metrics** – Time to CPR, time to defibrillation, and quality of CPR are all factors that have been shown to have a positive impact on survival. One metric that field crews can use to evaluate performance is CPR Fraction.

(a) CPR Fraction – The time CPR is being performed divided by the total time of the cardiac arrest. This fraction is typically reported as a percentage.

(i) A target goal for crews, that has been associated with improvements in survival, is a CPR fraction of equal to or greater than 80%.

(ii) Minimizing pre-shock pauses (e.g., charging defibrillator while clinicians performing chest compressions)

(iii) Feedback is best provided in real time or as close to the provision of care as possible.

(b) CPR compression rates should be between 100 and 120 per minute.

(c) Compression pauses should always be less than 10 seconds.

**h) PROCEDURE: CODE RESOURCE MANAGEMENT (CRM)**
Crews should coordinate their duties keeping the call priorities in mind. Intervention priorities are (in order of highest to lowest):

- Compressions
- Defibrillation
- BLS Airway Adjuncts/Ventilations
- IV/IO Access
- Medications
- ALS Airway
The number of personnel on a given incident and the qualifications of those personnel can vary; however, the priorities remain the same. Appropriate crew roles are outlined below:

2 clinician crew:
Clinician 1 – Chest compressions
Clinician 2 – Ventilate, attach/operate AED/defibrillator, assume crew leader responsibilities (clinicians rotate positions every two minutes)

Roles remain the same even if clinicians are ALS equipped

3 clinician crew:
Clinician 1 – Chest compressions
Clinician 2 – Ventilate
Clinician 3 – Crew Leader, attach/operate AED/defibrillator
(Clinicians 1 and 2 rotate every two minutes)

Roles remain the same even if clinicians are ALS equipped

4 clinician crew:
Clinician 1 – Chest compressions
Clinician 2 – Ventilate
Clinician 3 – Attach/operate AED/defibrillator
Clinician 4 – Crew leader
(Clinicians 1, 2, and 3 rotate every two minutes)

** Once first two roles have begun treatment, ALS clinicians will establish IV/IO and administer medications.

Greater than 4 clinicians - Utilize the same initial assignments as the four clinician crew. The crew leader will assign additional roles such as informing the family of patient status, gathering patient information, and documenting the medical interventions performed on the call. If resources allow, rotate additional clinicians to do chest compressions to achieve optimal performance.

Crew leader - The crew leader will keep time, record interventions performed during the arrest, give compression feedback and ensure rotation of personnel doing compressions every two minutes. Verbal announcements of time should occur at one minute, 30 seconds before reassessment, 15 seconds left, and countdown to reassessment at 10 seconds.
PEDIATRIC HIGH PERFORMANCE CPR (HPCPR)

Assess Patient (less than 10 seconds)

Unresponsive
Not Breathing
No pulse

Clinician #1
Start Chest Compressions (100-120/min)
Ventilations 2 Breaths: 30 Compressions
Call for AED/Defibrillator

Clinician #2
Ventilations 2 Breaths: 15 compressions
Place Airway Adjunct
Suction
Attach AED/Defibrillator

Clinician #3 or More
• Obtain IO Access
• Administer Medication
• Establish ALS Airway
• Family Support

Coordinated pause Activities (complete in less than 10 seconds)
• Check pulse
• Check rhythm
• Shock if indicated
• Rotate compressors
• Resume CPR

2 minute cycles

Pediatric HPCPR Team Member Initial Roles

**Clinician #1:**
• Chest compressions at 100-120 per minute
• Call for AED

**Clinician #2:**
• Ventilate at 2 breaths: 15 compressions
• Attach AED

**Clinician #3 or MORE:**
• Assume timekeeper role
• Assume AED role
• IO Access
• Medications
• Establish ALS Airway
• Family Support

Essentials of High Performance CPR for Pediatrics

1. Ensure proper chest compression rate
   • 100-120/min
2. Ensure proper compression depth
   • Less than 1 year – 1 ½ inches (4 cm)
   • Greater than or equal to 1 year – 2 inches (5 cm)
3. Minimize interruptions (less than 10 second pause)
4. Ensure full chest recoil
5. Coordinate 2 minute cycles
6. Rotate Compressor

* Once an advanced airway is in place, one ventilation every 6 seconds interposed asynchronously
Procedures –
INTRAOSSEOUS INFUSION

a) PURPOSE
The administration of fluids and medications via intraosseous (IO) infusion has long been known to be a relatively safe and effective procedure in the treatment of critically ill patients.

b) INDICATIONS
Patients in which the following conditions are present:
(1) Cardiac arrest, OR
(2) Profound hypovolemia, OR
(3) No available vascular access, or following two unsuccessful peripheral IV attempts for patients with any other life-threatening illness or injury requiring immediate pharmacological or volume intervention OR
(4) In pediatric patients in cardiac arrest, go directly to IO if no peripheral sites are obvious and without having to attempt peripheral access.

c) PROCEDURES
Allowable sites for IO:
(1) Sites for manual placement of IO needle
   (a) IO needle with 18 gauge should be used in patients less than 3 kg.
   (b) Patients 6 years of age or less, use the proximal tibial site: locate the preferred site of 1–3 cm distal to the tibial tuberosity on the anteromedial surface of the tibia.
   (c) Patients greater than 6 years of age, use the distal tibial site: locate the medial surface of the distal tibia just proximal to the medial malleolus.
(2) Sites for mechanical placement of IO needle
   (a) Select appropriate site:
      (i) Patients 3–39 kg or who have not yet reached their 13th birthday: use the proximal tibial site. Extend the leg. Insertion site is approximately 1 cm medial to the tibial tuberosity, or just below the patella (approximately 1 cm or one finger width) and slightly medial (approximately 1 cm or one finger width), along the flat aspect of the tibia. Pinch the tibia between your fingers to identify the center of the medial and lateral borders. Aim the needle set at a 90-degree angle to center of the bone.
      (ii) Patients 40 kg and greater and who have reached their 13th birthday:
         a. Preferred site: use the proximal humerus site: Place the patient’s hand over the abdomen (elbow adducted and humerus internally rotated). Secure the arm in place across the abdomen.
            i. Place your palm on the patient’s shoulder anteriorly. The area that feels like a “ball” under your palm is the general target area. You should be able to feel this ball, even on obese patients, by pushing deeply.
            ii. Place the ulnar aspect of your hand vertically over the axilla.
Procedures – INTRAOSSEOUS INFUSION (continued)

iii. Place the ulnar aspect of your other hand along the midline of the upper arm laterally.

iv. Place your thumbs together over the arm. This identifies the vertical line of insertion on the proximal humerus.

v. Palpate deeply up the humerus to the surgical neck. This may feel like a golf ball on a tee. The spot where the “ball” meets the “tee” is the surgical neck.

vi. The insertion site is 1 to 2 cm above the surgical neck, on the most prominent aspect of the greater tubercle. Point the needle set tip at a 45-degree angle to the anterior plane and posteromedial.

b. If proximal humerus site is not available, use the proximal tibial site. Extend the leg. Insertion site is approximately 2 cm medial to the tibial tuberosity, or approximately 3 cm (two finger widths) below the patella, and approximately 2 cm medial, along the flat aspect of the tibia. Aim the needle set at a 90-degree angle to the center of the bone.

c. If proximal site is not available, use the lower extremity distal tibia site. Insertion site is located approximately 3 cm (2 finger widths) proximal to the most prominent aspect of the medial malleolus. Palpate the anterior and posterior borders of the tibia to assure that your insertion site is on the flat center aspect of the bone. Aim the needle set at a 90-degree angle to the center of the bone.

(b) Select the appropriate needle:

(i) There are three lengths of 15 gauge mechanical IO needles.

(ii) Estimate tissue depth at selected site and select appropriate needle (15 mm, 25 mm, or 45 mm). Always use the 45 mm needle for the proximal humerus site. Point the needle set tip at a 45-degree angle to the anterior plane and posteromedial.

(iii) Insert so needle is touching bone.

(iv) Check the IO needle hub to assure that the 5 mm mark on the needle is visible when the tip of the needle touches the bone. The black line closest to the hub should be visible.

(v) Gently drill into the humerus 2 cm or until the hub is close to the skin. Gently drill, into the tibia approximately 1-2 cm after entry into the medullary space or until the needle set hub is close to the skin. Hold the hub in place and pull the driver straight off. Continue to hold the hub while twisting the stylet off the hub with counter-clockwise rotations. The catheter should feel firmly seated in the bone (1st confirmation of placement).

a. Place the stylet in a sharps container.

b. Place the dressing over the hub.

c. Attach an extension set to the hub if available; firmly secure by twisting clockwise.

d. Aspirate for blood/bone marrow (2nd confirmation of placement).

For patients unresponsive to pain:

e. Flush the IO catheter with 5-10 mL IV fluid.

TWO ATTEMPTS WITHIN FIVE MINUTES ARE PERMITTED. MEDICAL CONSULTATION SHOULD BE OBTAINED FOR FURTHER ATTEMPTS.
(3) Pain due to infusion via IO
   (a) To prevent or treat pain during an IO infusion for adults, administer 20–40 mg of 2\% (only 1–2 mL preservative free/cardiac) lidocaine IO.

   (b) To prevent or treat pain during an IO infusion for an adolescent patient (13–18 years of age), administer 20–40 mg of 2\% (only 1–2 mL preservative free/cardiac) lidocaine IO.

<table>
<thead>
<tr>
<th>Pediatric Lidocaine 2%</th>
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<tbody>
<tr>
<td><strong>Age</strong></td>
</tr>
<tr>
<td>Less than 5 years</td>
</tr>
<tr>
<td>5-12 years</td>
</tr>
<tr>
<td>13-17 years</td>
</tr>
<tr>
<td>Greater than 18 years</td>
</tr>
</tbody>
</table>

(c) Medical consultation is required for patients under 13 years of age.
(d) Slowly infuse lidocaine IO. Allow lidocaine to dwell in IO space 60 seconds. Flush with IV fluid.

d) CONTRAINDICATIONS
   (1) Conscious patient with stable vital signs
   (2) Peripheral vascular access readily available
   (3) Suspected or known fractures in the extremity targeted for IO infusion
   (4) Previous attempt in the same bone within 48 hours
   (5) Cellulitis at the intended site of the procedure
   (6) Patient with known bone disorder
   (7) Prior knee or shoulder joint replacement
   (8) Inability to identify landmarks

e) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
   (1) Extravasation of fluid
   (2) Infection
   (3) Compartment syndrome

f) PRECAUTIONS
   Humeral site: Stabilize the needle prior to any attempt at removing the driver. The humeral cortex can be considerably less dense, and failure to stabilize the needle may cause inadvertent dislodgement. Also, as patients advance in age, bone density continues to decrease and the proximal humeral needle’s stability must be routinely assessed.
Procedures – INTRAVENOUS MAINTENANCE
THERAPY FOR EMT

12.22

Procedures: Intravenous Maintenance Therapy for EMT 12.22

a) CLINICIAN-CONTROLLED IV SOLUTIONS
(1) The EMT is authorized to be the primary caregiver for patients with established intravenous (IV) therapy ONLY when the reason for transport is not related to complications associated with the IV line, and:
(a) The IV Solution DOES NOT contain:
   (i) MEDICATIONS,
   (ii) WHOLE BLOOD, or
   (iii) BLOOD PRODUCTS (such as plasma, platelets, or packed red blood cells)
(b) The IV catheter is placed in a PERIPHERAL LIMB VEIN, or
(c) The IV catheter is a capped (e.g., heparin-locked) peripheral or central line, and
(d) No other ALS interventions are required.

(2) IV fluids
   The EMT is authorized to perform IV maintenance of NON-MEDICATED IV solutions that contain only:
   (a) LR solution
   (b) 2.5%–10.0% dextrose in water
   (c) 0.25%–0.9% saline solution
   (d) Potassium chloride (KCL) added to the solution. The amount of KCL in solution shall not exceed 20 milli-equivalents (mEq)/liter OR
   (e) Peripheral Parenteral Nutrition (PPN) or Total Parenteral Nutrition (TPN)

   IF IV FLUIDS OR PPN ARE BEING ADMINISTERED VIA INFUSION PUMP AND NOT PATIENT-CONTROLLED, THE PATIENT MUST BE ACCOMPANIED BY A NURSE OR APPROPRIATELY TRAINED ALS CLINICIAN.

b) PATIENT-CONTROLLED MEDICATIONS OR IV SOLUTIONS
The EMT is authorized to be the primary caregiver for patients with established intravenous (IV) therapy ONLY when the reason for transport is not related to complications associated with the IV line or the medications being infused and the patient has been caring for the line, IV fluids, and/or IV medications at home without the assistance of a health care clinician.

UNDER NO CIRCUMSTANCES SHALL THE EMT CLINICIAN ATTEMPT TO MAKE ANY ADJUSTMENTS TO IV INFUSION PUMPS, NOR SHOULD THE EMT CLINICIAN ADMINISTER ANY ADDITIONAL MEDICATIONS OR IV FLUIDS.

c) Provide patient care according to appropriate protocol.
d) Routine IV maintenance procedures
(1) Ensure IV solution and catheter placement meets criteria above.
   (a) Request assistance of appropriate level health care clinician if IV solution and/or IV catheter placement do not meet criteria above, or
   (b) Request authorized personnel at health care facility to:
      (i) Replace IV solution with an appropriate IV solution, or
      (ii) Discontinue the IV prior to departing the scene.
Procedures – INTRAVENOUS MAINTENANCE THERAPY FOR EMT (continued)

(2) Confirm appropriate IV solution drip rate prior to transport.
(3) Ensure IV bag contains adequate volume of solution for duration of patient transport.
   If IV solution is not adequate, request authorized personnel at health care facility to:
   (a) Replace IV solution with an adequate volume, or
   (b) Discontinue the IV prior to departing the scene.
(4) Ensure IV solution is flowing at appropriate rate.
(5) Ensure patient has no signs or symptoms specifically related to complications of IV therapy prior to transport.
   If patient has signs or symptoms related to complications of IV therapy: Request authorized personnel at health care facility to correct the complication.

e) COMPLICATIONS OF IV THERAPY
(1) During patient transport, many possible complications of IV therapy may occur that the EMT must be prepared to manage.
   (a) Local complications may include: pain, hematoma, infiltration, infection, dislodged catheter, and tissue sloughing.

**DO NOT ATTEMPT TO REINSERT DISLODGED IV CATHETER.**

(b) Central complications may include: syncope, sepsis (infection), air embolism, pulmonary edema, pulmonary thromboembolism, congestive heart failure, overhydration, and catheter embolism.
(c) General complications may include: restricted flow (e.g., bent tubing, fluid-filled air chamber, inappropriate bag placement), and empty IV solution bag.
(2) Obtain medical direction and prepare to discontinue the IV if any of the complications described above are assessed and/or observed.
(3) If medical direction is genuinely not obtainable, the EMT shall discontinue the IV as soon as possible.

**THE EMT IS AUTHORIZED TO DISCONTINUE PERIPHERAL LIMB VEIN IVs ONLY.**

(4) Specific documentation includes:
   (a) Type of clinician-controlled IV solution
   (b) Type of patient-controlled IV solution
   (c) Type of patient-controlled IV medication
   (d) Volume administered
   (e) Complications encountered
Procedures – MEDEVAC UTILIZATION

12.23

a) PURPOSE
Summarize Medevac Utilization Protocol indications, contraindications, principles for consideration of medevac request, medevac request process, standardized medevac request dataset, optimal landing zone setup, and safety recommendations when interacting with helicopters

b) INDICATIONS FOR “MEDEVAC REQUEST”
The following indications must meet the specific criteria of the indicated protocol(s)
(1) Trauma Category Alpha, Bravo, Charlie*, Delta*
(2) Specialty Category
   (a) Burn
   (b) Hand*
   (c) Eye
   (d) Head
   (e) Spinal
(3) Medical Category
   (a) Stroke
   (b) STEMI
   (c) Hyperbaric (CO, Toxic Inhalation, or SCUBA)
(4) Consult-Approved Critical/Unstable (Time-critical illness or disease requiring specialized care)*

All of the above requests containing an asterisk (*) (adult or pediatric) require acceptance at the Trauma/Medical/Specialty Center for medevac authorization before SYSCOM can dispatch the helicopter.

c) PRINCIPLES FOR CONSIDERATION OF MEDEVAC TRANSPORT MEETING ABOVE INDICATIONS:
(1) Priority 1 Patients (critically ill or injured person requiring immediate attention: unstable patients with life-threatening injury or illness)
   (a) Consider air transportation if the patient will ARRIVE at the appropriate receiving facility more quickly than could be accomplished by ground transportation.
   (b) The clinician should consider all of the following:
      (i) Time for helicopter response
      (ii) Patient turnover (loading time)
      (iii) Flight time to appropriate facility
      (iv) Weather conditions
(2) Priority 2 Patients (less serious condition yet potentially life-threatening injury or illness, requiring emergency medical attention but not immediately endangering the patient’s life)

Consider medevac transport if drive time is greater than 30 minutes.

Special Consideration:
Consider medevac transport if ground transport greater than 60 minutes to a trauma or specialty center would deplete limited EMS resources in the community.
d) CONTRAINDICATION FOR MEDEVAC REQUEST
EMS/DNR-B or MOLST B patients are not candidates for field medevac transport.

ALL REQUESTS FOR SCENE HELICOPTER TRANSPORTS SHALL BE MADE THROUGH SYSCOM.

e) FORMAL REQUEST PROCESS
The Systems Communications Center (SYSCOM) at MIEMSS serves as the communications center for the dispatching and management of Maryland’s public safety helicopter resources. This mission is accomplished through the partnership between jurisdictional 9-1-1 call-centers and SYSCOM operations at MIEMSS. All helicopter requests must be routed through SYSCOM. The Medevac Request Data form is designed to provide a consistent standard by which SYSCOM receives “request” information. Considering the variety in the types of requests received by SYSCOM (e.g., medevac, search-and-rescue, law enforcement tracking) the information requested will vary, depending on the nature of the request. The county communications centers and the EMS clinicians that make the request should be familiar with the Medevac Data Request form to provide essential data to SYSCOM for prompt dispatch of the requested helicopter support.

EMS clinician and 9-1-1 center medevac request process:
(1) Decision made to request medevac based on indication and principles above (if 9-1-1 center has enough information from phone interrogation of call, and trauma indications meet Trauma Decision Tree Category Alpha or Bravo, the 9-1-1 center operator does not have to wait for EMS clinician to arrive on scene to make medevac request).
(2) If indicated, consult with trauma/specialty center for physician authorization to use medevac for transport and acceptance of the patient.
(3) Essential information gathered to complete the Medevac Data Request form (most of this is handled by 9-1-1 center).
(4) Contact SYSCOM for formal medevac request.
(5) Select and secure landing zone following optimal landing zone setup and safety tips.
## Procedures – MEDEVAC UTILIZATION (continued)

### Medevac Data Request Form

**Maryland Helicopter Dispatch Request**

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<table>
<thead>
<tr>
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<tbody>
<tr>
<td>1</td>
<td>Identify Call Origin &amp; Operator ID</td>
</tr>
<tr>
<td>2</td>
<td>Identify Request Type: <strong>Medevac, Search &amp; Rescue, Airborne Law Enforcement</strong></td>
</tr>
<tr>
<td>3</td>
<td>Jurisdictional Incident Number &amp; 9-1-1 Dispatch Time</td>
</tr>
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**Medevac Dispatch**

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<tbody>
<tr>
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<td>Incident Type</td>
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<td>ALS Unit &amp; LZ Contact Info</td>
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<tr>
<td>10</td>
<td>Additional Relevant Information</td>
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**Search & Rescue Dispatch**

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<td>ADC Map Grid OR Lat/Lon Info for LZ</td>
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<td>Ground Contact Unit</td>
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<td>Additional Relevant Information</td>
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**Airborne Law Enforcement Dispatch**

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<td>1</td>
<td>Incident Type</td>
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<td>2</td>
<td>Incident Location: Community &amp; Site</td>
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<td>3</td>
<td>ADC Map Grid OR Lat/Lon Info for LZ</td>
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<td>4</td>
<td>Primary Target</td>
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<td>Time Last Observed</td>
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Procedures – MEDEVAC UTILIZATION (continued)

12.23

f) HELICOPTER SAFETY

(1) OPTIMAL LANDING ZONE (LZ) SETUP

(a) 150 x 150-foot area close to the incident scene and free from obstructions is the minimum required, with a 175 x 175-foot area preferred.

(In mass casualty incident, identify a large enough area to land multiple large helicopters.)

(b) The landing zone should be a flat surface that is firm, free of overhead obstructions, and free of any debris that can blow up into the rotor system. The maximum allowable slope is 10 degrees.

(c) Obstacles such as wires, poles, signs, etc. can be difficult to see from the aircraft. If wires are present at or near the scene, this information must be relayed to the flight crew prior to landing.

(d) Advise the flight crew on overhead radio contact if there are any obstructions in the area, obstructions at the edge of the LZ, or any obstructions in-line with the departure or approach path.

(e) The landing zone will not be located near fixed objects that may be susceptible to wind damage or unsecured objects (e.g., patio furniture, small boats) that may become airborne as the AW-139 aircraft produces a significant amount of main and tail rotor wash.

(f) If the roadway is too narrow, or numerous trees or other obstacles are present, another area must be selected as an alternate LZ and checked for obstacles and other unsafe conditions. After the LZ Officer has evaluated all areas, the best unobstructed landing site must be secured and the flight crew advised of any unsafe conditions they may encounter during the landing.

NOTE: In determining landing zones, be aware that helicopter take-offs and landings can be done in a vertical manner; however, these landings limit the pilot’s visibility of the LZ. Increased power requirements on the helicopter may eliminate land-back areas should an engine malfunction occur, making the approach slower and causing extended periods of rotor wash.

(2) ADDITIONAL LANDING ZONE TIPS

(a) The LZ Officer should walk the area on both sides of the LZ and check for hazards. During night operations, walk the LZ with a flashlight that is directed up and down to detect wires in and around the LZ.

(b) 45-Degree Test—The LZ Officer should stand in the middle of the LZ with one arm extended at a 45-degree angle in front of him/her. Any objects at or above this line are obstacles and need to be reported to the incoming aircraft. This test is done for the full 360 degrees.

(c) Do not recommend landing zones that contain loose material such as gravel. The rotor wash will cause stones or gravel to become airborne, striking personnel and/or damaging vehicles.

(d) When a roadway is to be used as an LZ, all traffic must be stopped in both directions of the roadway, even on multi-lane highways or interstates.
Procedures – MEDEVAC UTILIZATION (continued)

(e) The LZ Officer will ensure that enough personnel is available to prevent any breach of LZ security by pedestrians while the helicopter is approaching, on the ground, or while departing. Failure to do so may cause injuries and/or delay patient transport.

(f) Do not allow traffic to use the roadway until after the aircraft has departed. Traffic will be stopped at least 200 feet in both directions from the landing zone.

(g) Do not use flares or cones to mark the landing zone: they will become airborne during the landing. (Weighted cones/lights that are designed for aircraft operations are generally acceptable.)

(h) The flightcrew is the final authority when selecting an LZ. On some occasions, the flightcrew may not choose to utilize the ground personnel’s suggested LZ and choose an alternate LZ. This decision is usually based on information that is unknown to the ground personnel (e.g., wind, aircraft performance limitations).

(3) APPROACHING THE AIRCRAFT

Personnel should only approach MSP aircraft under the following conditions:

(a) Hearing and eye protection shall be utilized at all times when approaching the aircraft.

(b) Only when accompanied by an MSP flight crew member to the aircraft Response personnel are usually limited to four when loading patients. The crew will provide additional guidance prior to these personnel approaching the aircraft.

(c) In an emergency situation when it becomes necessary to render assistance or rescue occupants of the helicopter. In such cases: **DO NOT APPROACH THE AIRCRAFT UNLESS THE MAIN ROTOR HAS STOPPED!**

(d) Only approach the aircraft from the Safe Zone (see diagram).

(i) Never approach the aircraft from the rear areas due to the hazards existing from the tail rotor.

**REMAIN CLEAR OF THE REAR AND TAIL ROTOR AT ALL TIMES!**
(ii) If it becomes necessary to go from one side of the aircraft to the other, this will be done by walking around the front of the aircraft; however, do not walk under the rotor blades.

(iii) Personnel shall not wear hats and loose clothing when approaching the aircraft. Do not lift anything above shoulder height (e.g., IV bags).

(e) If the aircraft has landed on a slope or hill, care must be taken when approaching the aircraft from the downhill side. Uphill side approaches should be avoided, as the main rotor blade is spinning and is lower to the ground on one side of the aircraft. The Trooper/Flight Paramedic will provide additional guidance in this situation.

(f) Never bring the patient to the aircraft prior to advising the Trooper/Flight Paramedic of the patient’s information. Very high noise levels found in the general proximity of the aircraft make communication and patient turnover impossible.

(g) If debris gets in the eyes and it impairs the vision, do not continue to approach or egress from the aircraft. Personnel will immediately “take a knee,” and the Trooper/Flight Paramedic will provide assistance.

(4) MISCELLANEOUS SAFETY TIPS

(a) Aircraft Doors
   Personnel should not attempt to open or close any aircraft doors. If a person is in the aircraft, they should remain inside until the flight crew member opens the door, thus preventing damage to the door and greatly reducing the risk of an aircraft door opening inadvertently in flight.

(b) Vehicles
   (i) No vehicles or personnel shall be permitted within 200 feet of the aircraft.
   (ii) Do not direct spotlights onto the landing area or at the aircraft, but keep vehicle’s emergency lights displayed until the aircraft is overhead. Once the LZ has been confirmed and verified by the flight crew, vehicle lighting can be reduced to running lights or parking lights for night vision purposes.
Procedures –
PATIENT-INITIATED REFUSAL OF EMS

12.24

a) Initiate General Patient Care.
For the purposes of this protocol, a patient is defined as any person encountered by
in-service rescue or emergency medical personnel with an actual or potential injury or
medical problem. (The term “patient,” in this protocol only, refers both to patients and to
persons who are potential patients. This protocol is not intended to determine the legal
status of any person, the establishment of a clinician-patient relationship, or a legal
standard of care.)

A minor patient is defined as a patient who has not reached their 18th birthday and is
not
(1) Married, OR
(2) Parent of a child, OR
(3) Requesting:
   (a) Treatment for drug abuse or for alcoholism,
   (b) Treatment for Sexual Transmitted Infection (STI) or for contraception,
   (c) Treatment of injuries from alleged rape or sexual offense, OR
(4) Living separate and apart from the minor’s parent, parents, or guardian, whether
with or without consent of the minor’s parent, parents, or guardian, and is not
self-supporting, regardless of the source of the minor’s income.
An authorized decision maker for minor patients is defined as an adult who identifies
themselves as the parent or guardian, or has written authorization for medical decision
making or states that they have written authorization for medical decision making. Clinici-
ans may request the parent or guardian to present identification and will document the
name of the individual who identifies themselves as the decision maker.

IN CASES OF ALLEGED RAPE OR SEXUAL OFFENSE, LAW ENFORCEMENT OR SOCIAL
SERVICES SHALL BE NOTIFIED.

b) These persons may have requested an EMS response or may have had an EMS re-
response requested for them. Because of the hidden nature of some illnesses or injuries,
an assessment must be offered and performed, to the extent permitted, on all patients.
For patients initially refusing care, attempt to ask them, “Would you allow us to check
you out and evaluate whether you are OK?”

IF THE AUTHORIZED DECISION MAKER REFUSES TO PERMIT THE EMS CLINICIAN TO
EXAMINE A MINOR PATIENT TO DETERMINE THE SEVERITY OF THE ILLNESS OR INJU-
RY, THEN CONSIDER CONTACTING LAW ENFORCEMENT FOR ASSISTANCE. CONSID-
ER CONSULTATION WITH PEDIATRIC BASE STATION.

c) Each patient’s assessment shall include:
   (1) Visual assessment - injuries, responsiveness, level of consciousness, orientation,
       respiratory distress, gait, skin color, diaphoresis
   (2) Primary survey - airway, breathing, circulation, and disability
   (3) Vital signs - pulse, blood pressure, respiratory rate and effort, pulse oximeter when
       available
Procedures –
PATIENT-INITIATED REFUSAL OF EMS (continued)

(4) Secondary survey - directed by the chief complaint
   (a) Medical calls - exam of lungs, heart, abdomen, and extremities. Blood glucose
testing for patients with Diabetes Mellitus. Neurological exam for altered con-
sciousness, syncope, or possible stroke.
   (b) Trauma calls - for patients meeting criteria in the Maryland Medical Protocols
      Trauma Decision Tree recommending transport to a Trauma Center: exam of
      neck and spine, neurological exam, palpation and auscultation of affected body
      regions (chest, abdomen, pelvis, extremities).

(5) Capability to make medical decisions (complete questions 1 through 4 on the Pa-
tient-Initiated Refusal of EMS form):
   (a) Disorientation to person, place, time, situation
   (b) Evidence of altered level of consciousness resulting from head trauma, medical
      illness, intoxication, or other cause
   (c) Evidence of impaired judgment from alcohol or drug ingestion
   (d) Language communication barriers were removed by assuring “language line”
      translation when indicated
   (e) The patient understands the nature of the illness

d) Following the assessment, complete items 5 through 9 on the Patient-Initiated Refusal
   of EMS Form, noting the presence of conditions that may place the patient at higher
   risk of hidden illness/injury or of worse potential outcome.

Management

(1) Patients at the scene of an emergency who meet criteria to allow self-determination
    shall be allowed to make decisions regarding their medical care, including refusal of
    evaluation, treatment, or transport. These criteria include:
    (a) Medical capacity to make decisions - the ability to understand and discuss and
        understanding of the nature and consequences of the medical care decision
    (b) Adult (18 years of age or greater)
    (c) Those patients who have not reached their 18th birthday and are:
       (i) Married, OR
       (ii) Parent of a child, OR
       (iii) Requesting:
          a. Treatment for drug abuse or for alcoholism,
          b. Treatment for STI or for contraception,
          c. Treatment of injuries from alleged rape or sexual offense, OR
       (iv) Living separate and apart from the minor’s parent, parents, or guardian,
          whether with or without consent of the minor’s parent, parents, or guardian,
          and is self-supporting, regardless of the source of the minor’s income.
    (d) A patient who has been evaluated by EMS clinicians as having ‘no’
        answers to questions 1, 2, 3a, 3b, and 4 on the Patient-Initiated Refusal of EMS
        form shall be considered to be medically capable to make decisions regarding
        their own care.
Procedures –
PATIENT-INITIATED REFUSAL OF EMS (continued)

(e) Patients with ‘no’ answers to questions 1, 2, 3a, 3b, and 4 on the Patient-Initiated Refusal of EMS form but one or more ‘yes’ answers to questions 5 through 8 (medical conditions) have a higher risk of medical illness. The EMS clinician should consider consulting medical direction if the patient does not wish transport. The purpose of the consultation is to obtain a “second opinion” with the goal of helping the patient realize the seriousness of their condition and accept transportation.

(f) If the EMS clinician is unsure whether the patient has adequate ability to make medical decisions, they should seek medical consultation.

(g) At any time the EMS clinician identifies patient conditions that indicate that the patient should be transported to a hospital, and the patient is refusing transport, then the clinician should seek medical consultation.

(2) Any person at the scene of an emergency requesting an EMS response, or for whom an EMS response was requested, and who is evaluated to have any one of the following conditions, shall be considered incapable of making medical decisions regarding care and shall be transported, with law enforcement involvement, to the closest appropriate medical facility for further evaluation:

(a) Continued altered mental status from any cause including altered vital signs, influence of drugs and/or alcohol, metabolic causes (CNS or hypoglycemia), head trauma, or dementia

(b) Attempted suicide, danger to self or others, or verbalizing suicidal intent

(c) Acting in an irrational manner, to the extent that a reasonable person would believe that the medical capacity to make decisions is impaired

(d) Judgment impaired by severe illness or injury to the extent that a reasonable and medically capable person would seek further medical care

(e) On an Emergency Petition

(3) Further care should be provided according to Maryland Medical Protocols, Agitation protocol or other protocol sections as appropriate, based on patient’s condition.

e) Base Station Hospital Physician Consultation

Patient refusals are one of the highest risk encounters in clinical EMS. Careful assessment, patient counseling, and appropriate base hospital physician consultation can decrease non-transport of high-risk refusals. Patients who meet any of the following criteria require Base Station hospital physician consultation:

(1) The clinician is unsure if the patient is medically capable of refusing transport.

(2) The clinician disagrees with the patient’s decision to refuse transport due to unstable vital signs, clinical factors uncovered by the assessment, or the clinician’s judgment that the patient may have a poor outcome if not transported.

(3) The patient was involved in any mechanism included in the Trauma Decision Tree of the Maryland Medical Protocols that would recommend transportation to a Trauma Center.
Procedures –
PATIENT-INITIATED REFUSAL OF EMS (continued)

(4) Minor patients: No parent, guardian, or authorized decision maker is available or the clinician disagrees with decision made by the parent, guardian, or authorized decision maker.

For patients with significant past medical history, consider consultation with the specialty center that follows the patient if possible.

Patients who do not meet the criteria above but have one or more positive answers to questions 6 through 10 on the Patient-Initiated Refusal of EMS form may have a higher risk of illness. In these situations, clinicians shall consult with the Base Station hospital physician.

f) Documentation

(1) Complete Section One of the Patient-Initiated Refusal of EMS form, documenting the patient’s medical decision-making capability and any “At-Risk” criteria.

(2) Complete Section Two, which documents clinician assessment and actions.

(3) Following patient counseling and Base Station hospital consultation, when indicated, complete Section Three: Initial Disposition, Interventions, and Final Disposition.

(4) Have the patient and witness sign the refusal statement as determined by your jurisdiction.

(5) Document your assessment, the care provided, elements of the refusal, medical decision-making capability, and “At-Risk” criteria on the jurisdictional form.

(6) Submit copies of the Patient-Initiated Refusal of EMS form and the documentation form to the EMS Supervisor.

(7) If the patient/authorized decision maker refuses to sign the refusal statement:

   (a) Contact a supervisor.

   (b) Explain the need for a signature and again attempt to have the patient sign the refusal statement.

   (c) If not already done, have a witness sign the refusal statement.

   (d) Transmit the patient’s unwillingness to sign the refusal statement on a recorded channel and document all steps taken to convince patient to sign.
Section One:
When encountering a patient who is attempting to refuse EMS treatment or transport, assess their condition and record whether the patient screening reveals any lack of medical decision-making capability (1, 2, 3a, 3b, and 4) or high risk criteria (5–8):

1. Disoriented to:  
   Person? □ yes □ no  
   Place? □ yes □ no  
   Time? □ yes □ no  
   Situation? □ yes □ no

2. Altered level of consciousness? □ yes □ no

3. Alcohol or drug ingestion by history or exam with:  
   a. Slurred speech? □ yes □ no  
   b. Unsteady gait? □ yes □ no

4. Patient does not understand the nature of illness and potential for bad outcome? □ yes □ no

4A. Judgment impaired by severe illness or injury? □ yes □ no

5. Abnormal vital signs  
   For Adults  
   Pulse greater than 120 or less than 60? □ yes □ no  
   Systolic BP less than 90? □ yes □ no  
   Respirations greater than 30 or less than 10? □ yes □ no  
   For minor/pediatric patients  
   Age inappropriate HR or □ yes □ no  
   Age inappropriate RR or □ yes □ no  
   Age inappropriate BP □ yes □ no

6. Serious chief complaint (chest pain, SOB, syncope) □ yes □ no

7. Head Injury with history of loss of consciousness? □ yes □ no

8. Significant MOI or high suspicion of injury □ yes □ no

9. For minor/pediatric patients: ALTE, significant past medical history, or suspected intentional injury  
   If yes, consult

10. Clinician impression is that the patient requires hospital evaluation □ yes □ no

Section Two:  
For clinicians: Following your evaluation, document information and care below:

1. Did you perform an assessment (including exam) on this patient? □ yes □ no  
   If yes to #1, skip to #3

2. If unable to examine, did you attempt vital signs? □ yes □ no

3. Did you attempt to convince the patient or guardian to accept transport? □ yes □ no

4. Did you contact medical direction for patient still refusing service? □ yes □ no
Section Three: (CHECK ALL THAT APPLY)

Initial Disposition:

- Patient refused exam
- Patient refused treatment
- Patient refused transport
- Patient accepted exam
- Patient accepted treatment
- Patient accepted transport
- ADM refused exam
- ADM refused treatment
- ADM refused transport

Interventions:

- Attempt to convince patient
- Attempt to convince family member/ADM
- Contact Medical Direction (Facility: ________________________)
- Contact Law Enforcement
- None of the above available

Final Disposition:

- Patient refused exam
- Patient refused treatment
- Patient refused transport
- Patient accepted exam
- Patient accepted treatment
- Patient accepted transport
- ADM refused exam
- ADM refused treatment
- ADM refused transport

Section Four: (MUST COMPLETE)

Provide in the patient’s own words why they refused the above care/service:

_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________

Jurisdiction ______________________ Incident: ______________________ Date: __________
Unit #: _________________________ Clinician Name/EID: _______________ Time: __________
a) PURPOSE
IV access is an invasive skill reserved for ALS clinicians and “Program Approved Option” EMTs with IV Technician training. The purpose of establishing an IV line, or a saline-lock, is to provide direct venous access for the possible administration of fluids and ALS medications (ALS only), if necessary and appropriate.

b) INDICATIONS
1. See treatment protocols for initiation of IV.
2. If the protocol indicates to start an IV, the “Program Approved Option” EMT may initiate an IV or saline-lock, if appropriate.
3. Saline locks may be substituted for IV KVO anywhere in the protocol with the understanding that if the patient needs a fluid bolus or medication, the saline lock is converted to an IV of LR.
4. ALS clinicians, in the event of a life-threatening emergency (CRTs with medical consult) or cardiac arrest, may access indwelling or implanted, central or peripheral venous catheters for medication administration.
5. When a patient is a Hemophiliac A or B (Factor VIII or IX) and the family or patient states that the patient must have factor concentrate administered, the ALS clinician may assist the patient in the IV administration of the patient’s own factor concentrate (VIII or IX). Notify the receiving hospital of the administration of blood factor concentrate.
6. All ALS clinicians may access lower extremity IV sites. The CRT-(I) and paramedic should consider lower extremity IV sites prior to IO attempts (EMT-IV technicians may not access lower extremity IV sites).
7. The ALS clinician may establish a peripheral IV in a patient whose vasoactive medication has been interrupted due to a malfunctioning long-term access device that cannot be repaired by the home health caregiver. The ALS clinician can assist in re-establishment of an existing vasoactive infusion at the same dose or setting. Patient shall be transported to the nearest appropriate facility to access patient’s long-term device. When in doubt, obtain medical consultation.
8. Maximum 2,000 mL LR without medical consultation.
9. Second IV requires medical consultation except when initiating the Sepsis protocol and for ALS clinicians who have Priority 1 patient. Initiation of the second IV shall not delay transport.

c) CONTRAINDICATIONS
See treatment protocols.

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
See IV Maintenance Therapy for EMT.

e) PRECAUTIONS
All sharps must be properly disposed of in an appropriate container.
a) PURPOSE
To prevent harm to patient or others

b) INDICATIONS
(1) Patient physical restraints should be utilized only when necessary and only in situations where the patient is exhibiting behavior that the EMS clinician believes will present a danger to the patient or others.
(2) The procedure does apply to patients treated under implied consent.

c) PROCEDURE
(1) The physical restraint procedure applies to patients greater than 1 year of age.
   (a) Ensure that the scene is safe.
   (b) Ensure sufficient personnel are present to control the patient while restraining. Use police assistance when available.
   (c) Position the patient for safe transport:
      (i) Place patient face up or on their side, if at all possible.
      (ii) Secure extremities:
          ADULT: For adults, use 4-point restraints (ideally with one arm up and the opposite arm down) or use a sheet to carefully wrap the patient before applying a Reeves™-type stretcher.
          PEDIATRIC: For patients less than 13 years of age, use 3-point restraints (two arms, one leg) or use a sheet to carefully wrap the patient before applying a Reeves™-type stretcher.
      (iii) If police have handcuffed the patient, reposition the patient in face-up position with hands anterior and secured to the stretcher (jointly with police).
      (iv) If necessary, utilize cervical-spine precautions to control violent head or body movements.
      (v) Place padding under patient’s head. Pad any other area needed to prevent the patient from further harming himself or herself or restricting circulation.
      (vi) Secure the patient onto the stretcher for transport, using additional straps if necessary. Be prepared at all times to logroll, suction, and maintain airway, especially in the event of vomiting.
   (d) Monitor airway status continuously, utilize pulse oximetry, vital signs, and reassess pulse/capillary refill, motor, and sensory status distal to the restraints. Document findings every 15 minutes, along with reason for restraint.
   (e) Assess for traumatic or medical causes for the patient’s agitation. Refer to Agitation protocol.
   (f) For interfacility transfers, obtain a written physician’s order for use of restraints.

d) PHYSICAL RERAINT GUIDELINES:
(1) Use the minimum restraint necessary to accomplish necessary patient care and ensure safe transportation; soft restraints may be sufficient in some cases. If law enforcement or additional personnel are needed, call for assistance prior to attempting restraint procedures. Do not endanger yourself or your crew.
(2) Avoid placing restraints in such a way as to preclude evaluation of the patient’s medical status (airway, breathing, and circulation). Consider whether placement of restraints will interfere with necessary patient-care activities or will cause further harm.

(i) Patient positioning should be modified when restraining patients with limited mobility, previous injury, or preexisting conditions (e.g., osteoporosis or contracture) to maintain extremities in a neutral position.

(3) Patients shall not be restrained in a prone, hobbled, or “hog-tied” position.

(4) Once restraints have been placed, do not remove them until you arrive at the hospital unless there is a complication from their use. If possible, take extra personnel during transport to hospital to deal with potential complications.
a) PURPOSE

Coronary heart disease is the single largest cause of death in US men and women. Early identification and treatment of patients with acute myocardial infarction (AMI) has proven to reduce myocardial damage and decrease morbidity and mortality. Clinicians should be aware of both typical and atypical presentations.

b) INDICATIONS

(1) Chest pain that may radiate to the arm, shoulders, jaw, or back. Generally described as a crushing pain or toothache. May be accompanied by shortness of breath, sweating, nausea, or vomiting.
(2) Chest discomfort. Some heart attacks involve discomfort in the center of the chest that lasts for more than a few minutes or that goes away and comes back. This discomfort can feel like uncomfortable pressure, squeezing, or fullness.
(3) Discomfort in other areas of the upper body. Symptoms can include discomfort in one or both arms or in the back, neck, jaw, or stomach.
(4) Shortness of breath. This symptom often accompanies chest discomfort. However, it can also occur prior to the chest discomfort.
(5) Other anginal equivalents, such as: unexplained diaphoresis, nausea, lightheadedness, syncope, or a sense of impending doom.
(6) Post cardiac arrest with ROSC.

c) PROCEDURE

(1) Position patient.
(2) Place chest and limb leads.
(3) Acquire 12-lead (15-lead, if trained) and document the patient's last name, first initial, age, and gender. These identifiers should be on the transmission copy (if able to transmit) and shall be on the delivered printed copy.
(4) Continue patient care.

EARLY ASSESSMENT OF PATIENTS WITH ACS LEADS TO EARLY IDENTIFICATION OF A STEMl AND INFLUENCES DOWNSTREAM CARE AND PATIENT DESTINATION. 12-LEAD EKG SHOULD BE ACQUIRED WITHIN 10 MINUTES OF CONTACT WITH AN EKG-CAPABLE CLINICIAN.

SHOULD AN ALS CLINICIAN INITIATE TRANSFER OF CARE/RENDEZVOUS AND TRANSITION OF PATIENT CARE ALS TO BLS, IT SHALL BE IMPORTED INTO THE PATIENT CARE REPORT AND A COPY SHALL BE SENT WITH THE BLS UNIT TO THE RECEIVING FACILITY.
A Multi-Casualty Incident (MCI) or Unusual Event is any event where the number of injured persons exceeds the normal capabilities of the EMS Operational Program in whose jurisdiction the event takes place. Due to the size of the incident, the responding EMS Operational Program may require additional resources and/or must distribute patients to multiple hospitals.

Local EMS Operational programs should have a plan or operational procedures that address response to multiple patient incidents or unusual events. This protocol does not supersede those plans. There are some general practices and procedures that must be followed to ensure the EMS system can be prepared to respond appropriately to support a local response.

**Procedure**

a) Assess scene and recognize that the incident is an MCI or Unusual Event. The definition of MCI or Unusual Event for the purposes of this protocol is an incident that causes more than 5 patient encounters or that involves unusual circumstances that suggest it could place an extraordinary strain on EMS or health care resources. The following events are examples of an MCI or Unusual Event.

1. More than five patients from one or related incidents
2. Multi-patient events that require specialized rescue
3. Three or more immediate (Priority 1) patients
4. Multiple pediatric patients requiring specialty resources
5. More than one burn patient meeting burn center referral criteria
6. Use of more than two medevac helicopters
7. Use of Medical Ambulance Bus (MAB)
8. Multiple patients with unusual signs and symptoms
9. Unresolved WMD related activity that could result in multiple patients (active shooter, bomb threat, intentional WMD agent release, etc.)
10. Decontamination of more than 5 patients resulting in at least one transport
11. Unresolved hazardous material incident that has the potential to affect multiple patients
12. Evacuation of a licensed health care facility or housing complex for individuals requiring special assistance

b) Notify EMRC or the Regional EMRC as soon as the incident is recognized to be an MCI or Unusual Event. Use the specific terms “MCI” or “Unusual Event” when communicating with EMRC to be clear this protocol is being enacted. This should be done as early in the incident as possible when there is a strong suspicion that such an event has occurred so that EMRC may begin to notify hospitals and response partners of the
incident. Responding units can request their dispatchers notify EMRC before the scene is fully assessed if there is reasonable information to suggest that the incident meets the criteria above. As soon as available, the following information should be relayed to EMRC.

1. Type and general description of the incident
2. General location or address of the incident
3. Age range of patients
4. Estimated number of patients by priority
5. Approximate number of patients involved
6. Any hazardous agents involved

c) Initiate the incident command structure according to local SOPs and/or the National Incident Management System. Update EMRC with more details about the incident as they become available.

d) Consider utilization of the MCI Communications protocol.

e) Triage patients using the START / JumpSTART methods.
   1. Identify the patient’s triage category by utilizing the appropriately colored triage ribbon and securely attach a MIEMSS-approved Triage Tag.

f) Do not delay transport of patients for extensive patient care procedures. Provide only the care required to sustain life and limb during transport to the hospital.

g) Track the care, movement, and disposition of EVERY patient utilizing the locally approved triage/treatment/transport logs. Patient information should be written on the triage tag.

h) Consider the need for and request specialty resources through the local dispatch center and/or emergency management as per local procedures. These may include,
   1. Mass Casualty Support Units (MCSU) – (Medical Supply Caches)
   2. Medical Ambulance Buses
   3. CHEMPACK (Organophosphate antidotes - contact EMRC)
   4. Ambulance Strike Teams or EMS Taskforces
   5. Shock Trauma Go-Team

i) The Transportation Group Supervisor and Medical Communications Coordinator responsibilities should be assigned as early as possible. They are the critical link to EMRC, hospitals, and the health care system. Their duties include:
   1. Establish a final checkpoint through which all transport units MUST pass to ensure accountability of all patients.
   2. EMRC will have notified hospitals and acquired their bed availability based on the information originally received and will transmit that information to the scene when requested.
(3) Coordinate through EMRC the patient destination, and communicate the number of patients, general illnesses, ages, and triage category on each transport unit as they leave the scene to the receiving facilities.

(4) If a central point of contact cannot be established, individual transport units MUST communicate the above information individually through EMRC to the receiving hospitals during transport. Those units must announce that they are associated with the MCI or Unusual Event.

j) Coordinate with law enforcement and, if requested, assist the Coroner or Medical Examiner with identification and disposition of deceased casualties.

k) After the last patient has been transported, notify 9-1-1 dispatch center and EMRC that last patient has been transported. Demobilize scene, stand down or release resources dedicated to incident, and complete appropriate documentation. Cooperate with local officials, EMRC, hospitals, and emergency management to complete a final accounting of the disposition of all the patients.
Procedures – POTENTIALLY VOLATILE ENVIRONMENT WITH LIFE-SUSTAINING INTERVENTIONS

a) BACKGROUND
(1) A review of past active assailant incidents has shown that the conventional prehospital practice of not entering the scene until it is deemed safe by law enforcement (LE) has been associated with additional loss of life.
(2) This protocol is designed to be all-hazards in nature. It is meant to provide a clinical concept of operations that empowers trained and equipped, but not necessarily tactical, EMS prehospital clinicians, to access casualties and expedite life-sustaining interventions closer to the point and time of injury. For active assailant and other LE-related incidents, EMS clinicians shall be under LE escort. EMS clinicians shall use appropriate personal protective equipment as defined by local jurisdiction.
(a) Examples of such potentially volatile environments include, but are not limited to:
   (i) Active assailant (active shooter/IED) situations
   (ii) Post-blast detonations
   (iii) Intentional release of a chemical agent
   (iv) Industrial accident/explosion
   (v) Hazardous materials incident
   (vi) Structural collapse/urban search and rescue situations
   (vii) Transportation mishaps with limited scene access
   (viii) In the immediate aftermath of a natural disaster such as a tornado

b) INTRODUCTION
(1) This protocol provides guidelines for the type of intervention and care that should be rendered at various proximities to a threat in a potentially volatile environment.
(2) By definition, potentially volatile environments are dynamic in nature. Scene conditions may change and emergent evacuation of responders and patients may interfere with the delivery of interventions described in this protocol.

c) INDICATIONS
(1) This protocol does not replace or supersede the general patient care practices in The Maryland Medical Protocols for Emergency Medical Services, which are still to be followed once the concern of active threat has been mitigated.
(2) Use of this protocol is an acknowledgement by the EMS clinician that the situation is:
   (a) Unique, austere, and different than the conventional environment of care in which EMS medicine is usually rendered AND
   (b) The application of standard prehospital emergency practices could unnecessarily jeopardize the safety of the patient and/or medical clinician.
(3) An active assailant incident or Potentially Volatile Environments with Life-Sustaining Interventions (PVE/LSI) protocol is declared.

d) CONTRAINDICATIONS
(1) Absent the presence of perceived or actual threat, standard general patient care practices should be followed.
e) ZONES OF CARE/OPERATIONS

(1) The zones described below are intended to standardize the terminology used by responding emergency medical clinicians in Maryland and to establish a common understanding of the interventions to be performed within each zone.

(2) Hot Zone (Direct Threat): (Integrated Tactical EMS) Operational area with a direct and immediate threat to personal safety or health

(a) The overarching priority in the Hot Zone is mitigation of active threat. Medical care is a secondary function to threat mitigation.

(b) Medical clinicians must be an integrated tactical medic (i.e., TEMS) to operate in this environment. Medical priorities are to prevent casualties and responders from sustaining additional injuries and include prompt evacuation to a more secure zone.

(i) If at all possible, casualties should self-evacuate.

(ii) Goals of care include keeping the response team engaged in neutralizing the threat, minimizing public harm, and controlling life-threatening extremity hemorrhage.

a. Control of severe hemorrhage in the direct threat environment is best accomplished with commercially available tourniquets.

b. Tourniquet should be placed as high up on the limb as possible without taking the time to expose the area.

c. For full or partial amputation, immediately place a tourniquet if possible.

d. Cardiopulmonary resuscitation (CPR) is not indicated in this environment.

(iii) In circumstances of chemical agent exposure, administration of Nerve Agent Antidote Kits (NAAK/MARK-1) might be warranted if available.

(3) Warm Zone (Indirect Threat): (Limited LSI) Area with a potential threat to personal safety or health

(a) Evacuation of patients to a completely safe area is the primary objective of care in this area. The following care guidance is dependent on the availability of equipment, supplies, and the appropriate level clinicians. Extrication should NOT be delayed to provide advanced or involved treatment measures.

(i) The Warm Zone typically exists between the Hot Zone and Cold Zone, but is not geographic and depends on the evolving situation.

(ii) Responders must remain cognizant that scene security can change instantly.

(iii) A focused and deliberate approach to providing patient care should occur.

(iv) The potential benefits of providing medical care in these zones must outweigh the risks of the ongoing tactical operation and/or delaying opportunity to evacuate the patient.

(v) Care in the Warm Zone typically occurs at or near the point of injury once scene stabilizing measures have occurred. Care may also take place at a casualty collection point (CCP).

(vi) A CCP is a location concealed and covered from immediate threat where victims can be assembled for movement from areas of risk to the triage/
treatment area. Multiple CCPs may be required, which may be located in the Warm or Cold Zone. CCPs should be established and locations communicated as early as possible through operations to ALL responders.

(vii) If possible, an abbreviated triage system should be set up to identify the priority for the extrication of patients. The use of ribbons or markers to clearly identify immediate and delayed (red and yellow, respectively) patients is highly recommended. Deceased individuals should also be labeled/tagged appropriately to prevent repeat assessments by multiple clinicians.

(viii) Medical care in the Warm Zone should be limited to essential interventions only and is guided by the mnemonic “MARCHED”
  a. M – Massive Hemorrhage Control
     i. Massive hemorrhage remains the greatest threat to life in most trauma patients. Attaining hemorrhage control is the top priority.
     ii. Tourniquets remain the preferred means of hemorrhage control for life-threatening bleeding in this environment.
        1. If a tourniquet was applied in the Hot Zone, it should be reassessed.
        2. Tourniquets applied over clothing are not as effective and may need to be adjusted.
        3. Tourniquets should only be discontinued by an appropriately trained ALS clinician in consultation with medical control.
        4. Other methods of hemorrhage control include deep wound packing with either sterile gauze or hemostatic impregnated gauze.
        5. Vascular injuries in the neck, groin, and axilla (i.e., junctional zones) are not amenable to traditional extremity tourniquets. In addition, effective pressure dressings are often extremely difficult to apply. Hemostatic impregnated dressings with direct pressure (minimum 5 minutes with continuous pressure is preferred) have shown useful in such situations.

(b) A – Airway management
  (i) Patients in the Warm Zone with airway issues are high priority for evacuation due to their often intense resource requirements.
  (ii) Consider applying oxygen if available and indicated.
  (iii) Unconscious casualty without airway obstruction:
     a. Chin lift or jaw thrust maneuver
     b. Nasopharyngeal airway
     c. Place casualty in the recovery position
  (iv) Casualty with airway obstruction or impending airway obstruction:
     a. Chin lift or jaw thrust maneuver
     b. Nasopharyngeal airway
     c. Allow casualty to assume position that best protects the airway, including sitting up or leaning forward
     d. Place unconscious casualty in the recovery position
(v) If previous measures unsuccessful, if time and resources permit, consider per protocol:
   a. Supraglottic Devices (e.g., King LT™, EASYTube®, or Combitube™).
   b. Oro/nasotracheal intubation
   c. Surgical cricothyroidotomy

(c) R – Respirations
   (i) The chest/upper abdomen should be assessed for any evidence of an open chest wound and an occlusive dressing should be applied accordingly.
   (ii) Tension pneumothorax remains a significant cause of preventable death in trauma patients.
      a. In suboptimal environments that interfere with complete physical assessment, any patient with significant blunt or penetrating chest trauma who displays dyspnea should be treated as a developing tension pneumothorax and receive needle decompression, if appropriate.
      b. To be effective, needle decompression needs to be performed using at least a 3.25 inch, 14g needle/catheter or needle decompression thoracostomy kit.

(d) C – Circulation
   (i) In general, healthy adult trauma patients with a radial pulse and normal mentation do not need IV therapy in the Warm Zone.
   (ii) Patients with evidence of hypotension:
      a. If the patient displays signs of a closed head injury, IV fluid therapy is indicated to maintain at least a radial pulse or SBP of at least 90 mmHg.
      b. Patients in hypovolemic shock should receive a one-time 500 mL bolus of IV fluid.
   (iii) Patients in traumatic cardiac arrest should be considered deceased and no CPR should be performed in this zone.

(e) H – Hypothermia
   (i) Hypothermia in trauma patients has been associated with increased mortality. Hypothermia is easier to prevent than treat.
      a. Patients should be moved to a warmed location if possible.
      b. Efforts should be made to minimize heat loss.

(f) E – Everything else
   (i) Consider Mark I/DuoDote® for suspected organophosphate/nerve agent exposure.
   (ii) Dependent upon resource availability, burns, eye injuries, and acute pain should be managed per The Maryland Medical Protocols for Emergency Medical Services.

(g) D – Documentation
   (i) Key findings and interventions should be conveyed to the next phase of care.
(4) **Cold Zone**: (Traditional Patient Care Protocols) Area surrounding the Warm Zone. Responders can operate without concern of danger or threat to personal safety or health.

(a) Casualties are moved from the Warm Zone to the Cold Zone by way of an evacuation corridor(s).

(i) Evacuation Corridor: An area transitioning between the Warm and Cold Zone that is secured from immediate threat and allows for a mitigated risk in transporting victims from the CCP to the triage/treatment area beyond the outer perimeter.

(b) Once in the Cold Zone, casualties will require re-triage, particularly assessing for the development of a life-threatening condition and effects of Warm Zone therapy.

(i) If massive hemorrhage has not been addressed or has been ineffectively managed, it should be immediately readdressed with strategies mentioned above.

(c) Patients should be triaged and transported per standard practices.

(d) Medical care in the Cold Zone should be dictated by resource availability and, when possible, equate to the general patient care standards in *The Maryland Medical Protocols for Emergency Medical Services*.

(e) CPR may have a larger role during the evacuation phase especially for patients with electrocution, hypothermia, non-traumatic arrest, or near drowning; however, it is still casualty count/resource dependent.
1. INDICATIONS

An emerging infectious disease (EID) is an infectious disease for which incidence in humans has increased in the past two decades or threatens to increase in the near future. These diseases, which respect no national boundaries, include:

a) New infections resulting from changes or evolution of existing organisms
b) Known infections spreading to new geographic areas or populations
c) Previously unrecognized infections appearing in areas undergoing ecologic transformation
d) Old infections reemerging as a result of antimicrobial resistance in known agents or breakdowns in public health measures.

The most recent example is Ebola Viral Disease (EVD). EIDs that meet this protocol will be posted on the MIEMSS website under the Infectious Disease Tab. Seasonal influenza is not considered an EID, but some of the same principles of infection control may apply to the more common infectious diseases.

e) Signs and Symptoms of an EID are based on specific case definitions for the disease:

(1) EVD case definition includes:
   Travel history or exposure and a set of signs and symptoms that are included in the case definition, which has evolved over time.

(2) Other future EID diseases may vary in their signs and symptoms, and could include:
   (a) Respiratory congestion
   (b) Sneezing/coughing
   (c) Nausea/vomiting
   (d) Skin rashes, hives, or “poxes”
   (e) Swollen lymph nodes
   (f) General malaise
   (g) Loss of appetite
   (h) Hemorrhage from mucosal membranes
   (i) Descending neurological deficits

f) Case Definition

As EIDs become more prevalent, the Centers for Disease Control and Prevention (CDC) typically publish a description of each disease, which is utilized to determine whether to include or exclude a Patient Under Investigation (PUI) for specific testing or treatment and specific isolation or quarantine measures. These case definitions will be posted on the MIEMSS website and include specific guidance on the identification, treatment, and appropriate transport of these patients and the appropriate use of PPE.

g) Modes of transmission

(1) In direct transmission, an infectious agent is transferred from a reservoir to a susceptible host by direct contact or droplet spread.
   (a) Direct contact occurs through skin-to-skin contact, kissing, and sexual intercourse. Direct contact also refers to contact with soil or vegetation harboring infectious organisms.
   (b) Droplet spread refers to spray with relatively large, short-range aerosols produced by sneezing, coughing, or even talking. Droplet spread is classified as direct because transmission is by direct spray over a few feet, before the droplets fall to the ground.
Indirect transmission refers to the transfer of an infectious agent from a reservoir to a host by suspended air particles, inanimate objects (vehicles), or animate intermediaries (vectors).

(a) Airborne transmission occurs when infectious agents are carried by dust or droplet nuclei suspended in air. Airborne dust includes material that has settled on surfaces and become re-suspended by air currents as well as infectious particles blown from the soil by the wind. In contrast to droplets that fall to the ground within a few feet, droplet nuclei may remain suspended in the air for long periods of time and may be blown over great distances.

(b) Vehicles that may indirectly transmit an infectious agent include food, water, biologic products (blood), and fomites (inanimate objects such as handkerchiefs, bedding, or surgical scalpels).

(c) Vectors such as mosquitoes, fleas, and ticks may carry an infectious agent through purely mechanical means or may support growth or changes in the agent.

2. TREATMENT

(a) If the presence of an EID at a scene is known prior to entering, don the appropriate PPE and limit entry into the scene to essential personnel only. If an EID is discovered during assessment, immediately don the appropriate PPE, clear the scene of non-essential personnel and initiate the recommended decontamination procedures.

(b) Initiate General Patient Care.

(c) Treat the patient according to the signs and symptoms presented and according to the MIEMSS guidance for the specific EID. Procedures that increase risk of distributing fluids or secretions should be limited to those absolutely necessary to maintain life and provide the patient with a reasonable level of comfort.

(d) Contain any bodily fluids or respiratory excretions prior to transporting the patient. A SURGICAL mask may be placed on the patient to limit respiratory droplet aerosolization.

N-95 SHOULD NEVER BE PLACED ON A PATIENT AS THEY RESTRICT THE EXCHANGE OF RESPIRATORY GASES AND TYPICALLY HAVE A ONE-WAY EXPIRATORY VALVE THAT ALLOWS DROPLETS TO BE AEROSOLIZED UPON EXPIRATION DEFEATING THE PURPOSE OF PLACING A MASK ON THE PATIENT.

(e) Transport the patient to the appropriate hospital.

Hospitals have been categorized into three levels based on their capabilities to assess and treat PUIs for designated EIDs. A list of designated EIDs will be published on the MIEMSS website.

(1) Frontline Hospitals (MDH designated) – All hospitals with emergency departments must have the capability to accept, identify, and isolate a PUI for a designated EID, then follow the approved procedures to notify the local health department to arrange for transfer to an Assessment Hospital. These patients will typically be transferred within 24 hours.

(2) Assessment Hospitals (MDH designated) – A facility that has the capability to receive, isolate, and provide care for a patient while testing is completed to confirm or deny the diagnosis of the suspected EID. The patient will remain at that hospital for 4 to 5 days until the patient is discharged or transfer to an designated Treatment Hospital.
(3) Treatment Hospitals (MDH designated) – A facility assessed by the CDC to have the capability to admit and provide comprehensive care for and manage a patient with a confirmed designated EID, until the patient is no longer ill or has died.

f) Transport from the scene
PUIs at a residence should be transported directly to an Assessment Hospital unless total transport time is no longer than 45 minutes greater than transport to the nearest Frontline Hospital ED. If transport time is longer than 45 minutes greater than transport to the nearest Frontline Hospital ED, the patient must be transported to the closest appropriate Frontline Hospital. Priority 1 and Priority 2 patients with unresolved symptoms that cannot be managed outside the hospital should be taken to the closest Frontline Hospital. Receiving hospital notification of all suspected PUI patients should be done as early as possible to allow for hospital staff to prepare. Helicopter transport NOT indicated for the PUI patient.

g) Transport of a health department monitored patient
Individuals who were exposed and have some risk of contracting the disease may be monitored or even quarantined by the health department. MIEMSS will be notified by MDH if these patients become ill and require transportation by EMS to hospitals and will contact the local jurisdictional or waivered commercial EMS Operational Program to arrange that transport. DHMH will determine the destination hospital.

h) Interfacility Transfer
Transfers between hospitals will be completed by EMSOPs who have been granted a waiver from licensing to modify an ambulance specifically to transport an EID patient and have specific plans, training and quality assurance processes in place to do so. Public Safety EMSOPs may be called upon as a backup if the waivered commercial services are not available. MDH will determine the destination hospital in these cases.

i) Communication
EMS clinicians transporting PUIs for designated EIDs MUST contact the receiving hospital via EMRC prior to beginning that transport and enter the hospital through the entrance designated by the receiving hospital. The term PUI must be used to ensure the hospital understands and is prepared to receive the patient. Obtaining medical direction from the closest Frontline and Assessment Hospitals is always an option to determine the appropriate destination.

j) Refusal of transport
If a PUI for a designated EID refuses care or transport, the EMS clinician should remove him/herself from the immediate presence of the patient and contact the local health department through their dispatch center or locally defined procedures and provide as much of the following information about the patient that is available.

(1) Full name
(2) Age
(3) Gender
(4) Home address
(5) Contact phone numbers
(6) Current location
(7) Recent travel history
(8) Signs and symptoms being displayed
(9) Recent contact history with Ebola patients
The EMS clinician should expect to be involved in a discussion of the situation with health department and law enforcement officials, and if a quarantine/isolation order is issued, should be prepared to assist law enforcement in carrying out that order.

| k) Treat the patient according to the signs and symptoms presented and according to the MIEMSS guidance for the specific EID. Limit invasive procedures and any that increase risk of distributing fluids or secretions to those absolutely necessary to maintain life and provide the patient with a reasonable level of comfort. |
| l) Pediatric patients under the age of 15 discovered at the home or in a non-health care environment should be transported to a Treatment Hospital that is also a Pediatric Trauma Center if transport times are not longer than 45 minutes greater than transport to the nearest Frontline Hospital ED. If transport times are longer than 45 minutes greater than transport to the nearest Frontline Hospital ED, the patient should be taken to an Assessment Hospital (if within 45 minute transport time) or the closest Frontline Hospital. |
a) INDICATIONS

During interfacility transports, a CRT or paramedic may monitor a patient on a continuous IV amiodarone infusion provided that:

(1) Amiodarone maintenance infusion must have been started by the hospital staff prior to the interfacility transfer. An IV amiodarone maintenance infusion may NOT be started by the prehospital clinician in the prehospital setting.

b) CONTRAINDICATIONS

Patients who have not yet reached their 18th birthday.

c) PROCEDURE

(1) Follow the appropriate ALS Algorithm and maintain the infusion as directed by the sending physician.

(2) The sending physician must document the infusion to be administered on the patient’s transport record or transport note, including the concentration of the medication and the infusion rate.

(3) The infusion must be maintained on an infusion pump designed for transport, and the clinician must be trained in the appropriate use of that specific make and model infusion pump. The ambulance must have an inverter to power the pump while in the vehicle.

(4) The total volume of amiodarone infused must be recorded on the patient care report.

(5) The patient must be on a cardiac monitor and vital signs should be documented on the patient care report at least every 15 minutes.

(6) When in doubt, contact the sending physician for medical direction.
a) INDICATIONS

During interfacility transports, a CRT-(I) or paramedic may monitor a patient on a continuous IV lidocaine infusion provided that:

1) The lidocaine infusion must have been started by the hospital staff prior to an interfacility transfer. IV lidocaine infusions may NOT be started by the prehospital clinician in the prehospital setting.

b) CONTRAINDICATIONS

Patients who are clinically unstable, including but not limited to, unstable vital signs and blood pressure, current arrhythmia, and active chest pain

c) PROCEDURE

1) Follow the appropriate ALS algorithm and maintain the infusion as directed by the sending physician.
2) The sending physician must document the infusion to be administered on the patient’s transport record or transport note, including the concentration of the medication and the infusion rate.
3) The infusion must be maintained on an infusion pump designed for transport, and the clinician must be trained in the appropriate use of that specific make and model infusion pump. The ambulance must have an inverter to power the pump while in the vehicle.
4) The total volume of lidocaine infused must be recorded on the patient care report.
5) The patient must be on a cardiac monitor and vital signs should be documented on the patient care report at least every 15 minutes.
6) When in doubt, contact the sending physician for medical direction.
a) **INDICATIONS**

A CRT or a paramedic may monitor a patient on a continuous morphine sulfate infusion as long as the infusion was started by hospital staff prior to interfacility transfer (it may NOT be initiated by EMS).

b) **CONTRAINDICATIONS**

Patients who are clinically unstable, including but not limited to, unstable vital signs and blood pressure (exception: patients being transported for hospice care)

c) **PROCEDURE**

1. Maintain the infusion as directed by the sending physician.
2. The sending physician must document the infusion to be administered on the patient's transport record or transport note, including the infusion rate.
3. The infusion must be maintained on an infusion pump designed for transport, and the clinician must be trained in the appropriate use of that specific make and model infusion pump. The ambulance must have an inverter to power the pump while in the vehicle.
4. The total volume of morphine infused must be recorded on the patient care report.
5. The patient must be on a cardiac monitor and vital signs should be documented on the patient care report at least every 15 minutes.
6. When in doubt, contact the *sending* physician for medical direction.
Interfacility – PROTON PUMP INHIBITOR FOR INTERFACILITY TRANSPORT

a) INDICATIONS

A CRT or a paramedic may monitor a patient on a PPI infusion as long as the infusion was started by hospital staff prior to interfacility transfer (it may NOT be initiated by EMS).

b) CONTRAINDICATIONS

(1) Patients who have unstable vital signs or are being transferred to an intensive care environment
(2) Patients with allergic reaction to infusing agent
(3) Patients who have not yet reached their 18th birthday

c) PROCEDURE

(1) Follow the appropriate ALS algorithm and maintain the infusion as directed by the sending physician/practitioner.
(2) The paramedic will review the sending physician’s order and will review the specific PPI agent to ensure appropriate administration, indications, and absence of contraindications.
(3) The PPI infusion must be maintained on an infusion pump designed for transport, and the clinician must be trained in the appropriate use of that specific make and model infusion pump. The ambulance must be equipped to power the pump while the pump is in the vehicle.
(4) The administration of the PPI infusion will be recorded on the patient care report to include the agent’s name, dose, rate, and volume infused during transport.
(5) When in doubt, contact the sending physician/practitioner for medical direction.
1. **PURPOSE**
   When pulses are difficult to detect, or when a blood pressure cannot be measured using a stethoscope, a vascular Doppler device can be utilized.

2. **INDICATIONS**
   Inability to palpate a pulse.

3. **CONTRAINDICATIONS**
   Patients who have not yet reached their 18th birthday.

4. **POTENTIAL ADVERSE EFFECTS/COMPLICATIONS**
   None

5. **PRECAUTIONS**
   When utilizing a Doppler device, avoid applying to much pressure with the device over the artery, as this may obliterate the pulse you are attempting to detect.

6. **PROCEDURE**
   a) Identify the appropriate artery (e.g., carotid, brachial, radial, femoral, dorsalis pedis).
   b) A large dollop of gel is applied to the site and to the device.
   c) The device is held gently over the artery (too much pressure may obliterate the pulse), and the volume adjusted to hear the pulsation.
   d) If the pulse is located, the area should be wiped clean, and the exact site should be marked with a pen or marker.
   e) If blood pressure is being taken, the clinician finds the pulse and listens as the cuff is inflated. When the pulse sound disappears, you have identified the systolic pressure.
   f) If no pulse is found, then sliding the device around the appropriate area or changing the angle of the device slightly may identify the location of the pulse. Be careful not to apply to much pressure on the skin.

7. **TRAINING AND DOCUMENTATION**
   a) Clinicians must complete practical training.
   b) Description of technique
   c) Demonstration of device (features, operation, troubleshooting)
   d) Documentation requirements (eMEDS®)
   e) Scenario
1. INDICATIONS
   a) Abdominal pain, (severe; age 55 and older)
   b) Cardiac arrest
   c) Dyspnea, undifferentiated
   d) Pulseless extremity
   e) Trauma, thoraco-abdominal
   f) Tension pneumothorax, suspected

2. CONTRAINDICATIONS
   a) Prehospital FAST exam shall not delay patient transport

3. PROCEDURE
   a) Initiate appropriate medical or trauma protocol including all BLS/ALS interventions.
   b) The credentialed ALS clinician will complete the appropriate prehospital ultrasound exam recording for at least six seconds.
   c) Exam will be interpreted and relayed to the consulting hospital. The EMS base station physician may change the hospital destination based on the findings of the prehospital ultrasound exam.
   d) Assure the exam is transmitted to the receiving facility through closed, secure network, along with patient care report.
   e) Ultrasound (US) Exams:
      (1) Abdomen: for patients aged 55 and older, with a high clinical suspicion for abdominal aortic aneurysm (abdominal pain radiating to the back or flank), perform an US of the abdominal aorta.
      (2) Cardiac arrest: for patients in cardiac arrest, perform US for:
         (i) Patients who have PEA as initial rhythm: obtain a cardiac view to determine presence or absence of cardiac activity
            a) If cardiac activity is absent, consider hyperkalemia or drug toxicity and treat appropriately
            b) If cardiac activity is present, evaluate for pericardial effusion/tamponade and perform lung US to evaluate for pneumothorax
         (ii) During CPR, use color flow US to monitor for ROSC, limiting interruptions in CPR for pulse checks.
         (iii) During CPR, to evaluate for endotracheal tube placement
         (iv) Prior to termination of resuscitation, perform cardiac US to verify absence of cardiac activity
      (3) Dyspnea: for patients with undifferentiated dyspnea, perform lung US:
         (i) to evaluate for presence of abnormal “b-lines”, indicating increased interstitial fluid (CHF/pneumonia), using accepted criteria
         (ii) to evaluate for the absence of lung sliding (pneumothorax)
      (4) Pulseless extremity: for patients with high suspicion for vascular occlusion, evaluate for color flow of affected limb per doppler protocol indications.
(5) Trauma: for patients presenting with torso or abdominal pain or who present with high-impact, high-mechanism trauma, a prehospital e-FAST exam will be performed.

(i) Morison’s perihepatic view
(ii) Pelvic view
(iii) Perisplenic view
(iv) Cardiac view
(v) Lung view (to evaluate for pleural sliding artifact)
**1. Indications**

Patients eligible for entry into the Stabilization Center must be without an acute medical or traumatic complaint. If the patient is not requesting evaluation for an emergency medical condition and substance use is suspected, including suspected opioid patients who have improved with naloxone, patient must consent to be evaluated and transported to the Stabilization Center. Then the EMS clinician must complete the Stabilization Inclusion Checklist.

**2. Treatment**

Initiate patient screening. All answers must be “YES” for the referral protocol to continue. For any “NO” answers, consultation with an adult Base Station is required.

<table>
<thead>
<tr>
<th>Patient without acute medical or traumatic complaint</th>
<th>YES</th>
<th>NO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient is age 18 or older</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Patient is willing and able to cooperate with examination</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Systolic BP greater than 80 mmHg and less than 220 mmHg</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Diastolic BP greater than 50 mmHg and less than 120 mmHg</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Pulse less than 120</td>
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<td>NO</td>
</tr>
<tr>
<td>Pulse greater than 50</td>
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<td>NO</td>
</tr>
<tr>
<td>Respiratory rate less than 22</td>
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<td>NO</td>
</tr>
<tr>
<td>Respiratory rate greater than 10</td>
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<td>NO</td>
</tr>
<tr>
<td>Blood glucose less than 300 mg/dl</td>
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<td>NO</td>
</tr>
<tr>
<td>Blood glucose greater than 70 mg/dl</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Pulse oximetry greater than 92% and no supplemental oxygen required</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Patient accepts transport to Stabilization Center</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>NO evidence of significant head or truncal trauma</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>NO evidence of new head trauma (ecchymoses, hematomas)</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>NO evidence of uncontrolled bleeding</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>Patient can walk with no more than minimal assistance → No assistive devices (cane, walker permitted) → No assistance/stabilization of more than one limb required</td>
<td>YES</td>
<td>NO</td>
</tr>
</tbody>
</table>

3. Medical consultation is required for any “NO” response.

4. If all answers are “YES” or medical consultation approves if a “NO” occurs, the patient shall be transported to the Stabilization Center.
Pilot Program –
ALTERNATIVE DESTINATION PROGRAM

Alternative Destination (AD) Protocol

- Low Acuity / Priority 3 Patient
- Patient is 18 years of age or older
- Able to Communicate with EMS
- Understands Consent Form/Process
- Agrees to be transported to AD

Vital Signs Acceptable?

Yes

High Risk Conditions Absent?

Yes

Physical Exam without Time Dependent Needs

Yes

Obtain Consent from Patient

Contact & Obtain Acceptance of Patient from AD

Transport to AD

ACCEPTABLE VITAL SIGNS
- Respirations: 10-20
- Pulse: 50-120
- Pulse Ox: >92%
- Temperature: 96-102 F
- Blood Glucose: 70-300

ACCEPTABLE BLOOD PRESSURES:
- Urgent Care/PCP:
  Systolic 100-160 & Diastolic 60-100
- Stabilization/Crisis Center:
  Systolic 80-220 & Diastolic 50-120

High Risk Conditions
- Abdominal Pain, Unexplained
- Altered Mental Status
- Back Pain, Unexplained
- Chest Pain
- Dyspnea/Shortness of Breath
- Focal Neurological Deficits (Acute)
- Seizures
- Sepsis, Suspected
- Syncope
- Requires more than minimal assistance to walk
- Unable to Cooperate with History and Exam

Physical Exam/Time Dependent Needs
- Airway
- Breathing
- Circulation (Including to Extremity)
- Disability (Deficit) or Deformity
- Severe Tenderness with Palpation/Exam
- Significant Head or Truncal Trauma
- Uncontrolled Bleeding
- Require ALS Monitoring or Interventions
- Concern for Potential Deterioration in Condition

IF ANY HIGH RISK CONDITIONS OR PHYSICAL EXAM/TIME DEPENDENT NEEDS, EMS SHALL TRANSPORT TO CLOSEST APPROPRIATE ED/FEMF

IF PATIENT IS EXCLUDED BASED ON VITALS ALONE, TRANSPORT TO CLOSEST APPROPRIATE ED/FEMF UNLESS MEDICAL DIRECTION FROM APPROVED BASE STATION AUTHORIZES TRANSPORT TO ALT DESTINATION

Rev. 06/10/2019 – Approved EMS Board 06/11/2019
### Examples of Low Acuity Chief Complaints

- Allergy or hay fever
- Back pain, mild; able to walk without assistance
- Contusions or abrasions, minor
- Cough, mild; without hemoptysis or respiratory impairment
- Non-traumatic dental problems
- Diarrhea, without dizziness or other signs of dehydration
- Dizziness, chronic (recurrent or known history)
- Dysuria, mild; female
- Ear pain
- Ingrown toenails
- Itching without systemic rash
- Eye irritation without signs of active infection, minor
- Fracture, distal extremity (forearm, lower leg), isolated injury, not open, With neuro/ vascular intact
- Headache, minor without neurological impairment
- Injury follow-up (minor injury, treated previously)
- Joint pain
- Mouth blisters
- Muscle aches
- Nausea, vomiting
- Neck pain (no history of acute trauma)
- Nosebleed (resolved)
- Painless urethral discharge
- Physical exam requests (except patients with diabetes, CHF, kidney failure, cancer)
- Plantar warts
- Rectal pain/itching, minor
- Sexual disease exposure
- Simple localized rash
- Sinusitis, chronic
- Skin infection or sores, minor
- Sore throat without stridor
- Sunburn (localized without blisters)
- Vaginal discharge
- Vaginal bleeding (Hx non-pregnant, not postpartum, and requires less than one pad in 5 hours)
- Upper respiratory infection
- Work release or disability
- Wound checks
Note: This document does not contain all of the material approved by the EMS Board. For the entire text of the protocol, contact the Office of the Medical Director.

1. PURPOSE
The objective of this pilot program is to assess the impact, accuracy and safety of providing low-acuity patients, identified as Alpha patients by IAED criteria (Basic Life Support), with immediate on-scene care by a two-person team composed of a BCFD Minor Definitive Care Now (MDCN) paramedic clinician, and one of the following Advanced Level Clinicians (ALP): a UMMC Nurse Practitioner (NP), a Maryland-licensed physician affiliated with UMMC with board certification in emergency medicine ("Physician"), or UMMC Physician Assistant (PA). This will be referred to as the MDCN Team.

2. INDICATIONS
a) Low-acuity patients, identified by the IAED™ MPDS® protocol as an ‘Alpha determinant code Basic Life Support,’ who meet additional criteria outlined in the MDCN protocol below; AND
b) Patients with an incident address that falls within the geographic boundaries of the UMMC, Midtown Campus or Grace Medical Center catchment areas; AND
c) Patients who consent to participate in the MDCN Pilot Program.

3. CONTRAINDICATIONS
a) Patients who decline enrollment in MDCN Pilot Program;
b) Patients who are deemed clinically inappropriate for on-scene treatment by the MDCN Team following assessment;
c) Individuals who refuse participation by revoking written consent, verbal refusal of care at time of visit;
d) Patients who possess a language or communication barrier that inhibits the MDCN Team’s ability to appropriately address the patient’s needs at the scene;
e) Patients who are not able to or lack the capacity to understand the informed consent process; and
f) Patients who have not yet reached their 18th birthday.

4. GENERAL PROCEDURES
a) When a 9-1-1 call response for EMS service is dispatched, the MDCN Team will respond to the scene concurrently with the typical BCFD EMS response unit to Alpha-level calls within the UMMC, Midtown Campus and Grace Medical Center patient catchment areas.
b) If a patient refuses EMS care and transport, a patient refusal form and eMEDS® should be completed per The Maryland Medical Protocols for Emergency Medical Services while on scene.
c) If the patient is determined to be a low-acuity candidate for MDCN program (as defined in Section VI below), the BCFD EMS response personnel will offer the patient the option to be seen by the MDCN Team.
d) The MDCN Team will request patient consent (see MDCN Consent Form) to provide minor definitive treatment on scene.
e) Once consent is provided, patient information, including information collected by the EMS response personnel can be shared with the ALP.
f) The EMS response personnel will return to service. If the MDCN Team determines that the patient needs to be transported and the patient decides they want to be transported, or if for any reason, the patient decides they want to be transported, the MDCN Paramedic will radio PSAP for an EMS transport unit. After requesting the unit, the BCFD MDCN Paramedic will perform any advanced life support skills, as defined by the MIEMSS Protocols for EMS Clinicians, to provide all necessary care within their scope of practice, until additional EMS clinicians arrive on scene and assume patient care and transport to the closest appropriate hospital. Any care rendered under the MIEMSS Protocols will be documented in eMEDS.

g) The MDCN Team performs any additional assessment and if indicated, the ALP will render treatment. The MDCN Paramedic may assist with patient assessment (e.g., vital signs, pulse oximetry), the ALP will provide treatment associated with the MDCN Pilot Program.

h) The ALP may also offer to assist patients with setting up clinic appointments. The Operations Center, located at UMMC, may call and connect patients to appropriate care, either inside or outside of the University of Maryland Medical System (UMMS), depending on need, preference, and insurance status of the patient.

i) The MDCN Team documents the patient care encounter in the UMMC electronic health record system (“Epic”). If at any time during the encounter the patient refuses further assessment or treatment, the refusal must be documented in Epic.

j) The UMMC ALP and BCFD MDCN Paramedic clinicians will be restricted to their respective scopes of practice set by the Maryland Board of Nursing, Maryland Board of Physicians and MIEMSS.

5. ADVANCED LEVEL PRACTITIONER PROCEDURES
   a) This protocol may only be used by the Advanced Level Practitioner (ALP).
   b) MDCN Paramedics will follow The Maryland Medical Protocols for Emergency Medical Services.
   c) Under the MDCN Pilot Program, all eligible patients will be offered the choice to “opt in” to receive on-scene definitive care. Participation in this pilot program is voluntary and will require patients to provide signed, informed consent. The on-scene treatment provided by the ALP will be in accordance with the medication and procedure list as approved by MIEMSS.
Pilot Program – MINOR DEFINITIVE CARE (continued)

14.5

Pilot Program: Minor Definitive Care

d) Inclusion Criteria: the patient must provide consent and must not have any of the following exclusion criteria:

(1) A chief complaint consistent with evaluation that would indicate a need for the capabilities of a full service ED
   (a) High risk chief complaints are currently defined as dyspnea, altered mental status, syncope, chest pain, focal neurological deficits, unexplained back or abdominal pain, seizures, and sepsis (see vital sign criteria listed in 8. Medical Consultation (see below).

(2) Physical findings consistent with time-dependent needs for emergent assessment or stabilization
   (a) Signs on exams that indicate a threat to airway, breathing, circulation, circulation to an extremity, disability (deficit) or deformity, as well as severe tenderness (as indicated by an assessment of airway, breathing, circulation, disability, exposure (ABCDE), etc.).

(3) Reasonably foreseeable signs or suspicion of any deterioration of condition (e.g., airway, breathing, hemodynamic, or neurologic compromise)

(4) Any requirement for any advance life support (ALS) monitoring or ALS interventions

e) In order to include the patient in the MDCN Pilot Program, the MDCN Team will obtain a complete set of vital signs, medical history, and the ALP will obtain a signed MDCN Pilot Program Consent Form.

f) If the patient is stable and deemed by the ALP to meet the criteria of the MDCN protocol, and has an injury or disease process, which can be safely treated on-scene:

(1) The consenting patient will receive definitive on-scene care by the ALP member of the MDCN Team.

(2) If the patient refuses to participate in the MDCN Pilot Program, the patient’s condition deteriorates, or while on scene the patient changes their mind and declines to participate, the patient will be taken to the closest appropriate ED via ambulance. See 4. General Procedures above for response steps.

g) The MDCN Team will provide discharge instructions for each patient who participates in the MDCN Pilot Program.

h) In the event that the MDCN Team evaluates the consented patient and recommends ED transfer but the patient refuses, see 4. General Procedures above for appropriate actions.

6. MEDICATION MANAGEMENT

The ALP is authorized to manage drugs and devices under the following protocols:

a) The management of drugs or devices includes evaluating, initiating, altering, discontinuing, furnishing and ordering of prescriptive and over-the-counter medications.

b) Medication evaluation includes assessment of:

(1) Other medications being taken

(2) Prior medications used for current condition

(3) Medication allergies and contraindications, including appropriate labs and exams
c) The drug or device is appropriate to the condition being treated, and:
   (1) Accepted dosages per references.
   (2) Generic medications are ordered if appropriate.

d) A plan for follow-up is written in the patient's chart and provided to the patient.
e) The prescription must be written in patient's Epic chart, including name of drug, strength, instructions and quantity, and signature of the ALP.

7. DISPENSING MEDICATIONS
The ALP may dispense prescription drugs and devices, under the following protocols:

a) They have current prescriptive authority, including Maryland CDS registrations.
b) All drugs and devices ordered are limited to the Formulary OR are per the recommendations in the Resources listed in this document.
c) The drugs and devices ordered are consistent with the ALP's educational preparation or for which clinical competency has been established and maintained.
d) The drug or device ordered is appropriate to the condition being treated.
e) Patient education is given regarding the drug or device.
f) The name, title, and licensing number of the ALP is written on the transmittal order.
g) A physician affiliated with the MDCN Pilot Program is available during hours of operation for in person or telephone medical consultation.
h) The drug or device utilizes required pharmacy containers and labeling.
i) All appropriate record keeping practices of the dispensary are performed.
j) All other applicable Standardized Procedures in this document are followed during health care management.
k) All General Policies regarding Review, Approval, Setting, Education, Evaluation, Patient Records, Supervision and Consultation in these Standardized Procedures are in force.

8. MEDICAL CONSULTATION
While it is the intent of MDCN Pilot Program to respond to low-acuity calls, if immediate patient deterioration should occur, EMS transport resources shall be utilized.

MDCN Medical Director notification and/or emergent ALS transport to the closest appropriate ED with the following being examples of patients and scenarios that shall generate ALS transport:

a) Acute myocardial infarction (AMI) or symptoms consistent with AMI
b) Acute central nervous system or focal neurologic deficits
c) Severe CHF
d) Severe respiratory distress
e) $O_2$ Saturation less than 90% on room air, if acute
f) Hypotension
g) Acute altered mental status, unless intoxicated
h) Adult heart rate greater than or equal to 140
i) Emergency hypotension
j) Moderate to severe CHF
k) SBP greater than or equal to 240 or DBP greater than or equal to 140 at presentation (asymptomatic) with preexisting hypertension history
Pilot Program – MINOR DEFINITIVE CARE (continued)

l) Adult heart rate greater than or equal to 110 at time of disposition
m) The MDCN Team responds in less than 14 days for same acute complaint *Does not apply to chronic recurrent complaints unless there is a change in the complaint*

n) Elevated BP or heart rate in pregnancy or less than or equal to 6 weeks post-partum

o) Pregnancy complications

p) Chest pain (potentially consistent with angina or angina equivalent symptoms)
   (1) Nonspecific chest pain age greater than or equal to 30 with history of:
       • Hypertension
       • Diabetes
       • Smoking
       • Coronary artery disease
       • Hyperlipidemia
       • Family history of coronary artery disease by age of 60; OR
       Nonspecific chest pain age greater than or equal to 50 without risk factors
       • Abdominal pain
       • Requiring analgesic
       Nonspecific chest pain age greater than or equal to 70
       • Diabetic
       • Uncertain diagnosis

   (2) Lab Criteria:
       • D-Stick – less than 70 or greater than 300
       • Oxygen Sat 2% less than chronic levels

   (3) Vital sign and age consult criteria
       • Heart rate/minute
         □ Adult heart rate greater than or equal to 110
       • Hypertension
         □ Adult asymptomatic hypertension of SBP greater than 220 or DBP greater than 120 at time of disposition with history of hypertension
         □ Adult asymptomatic SBP greater than 195 or DBP greater than 115 at disposition without history of hypertension
1. **Indications**

Clinical suspicion for major blood loss **WITH** evidence of significant physiologic compromise

**Clinical suspicion for major blood loss, such as:**

- Penetrating trauma to the trunk
- Unstable pelvic fracture or multiple long bone fractures
- Blunt trauma mechanism consistent with major internal blood loss
- Observed major external blood loss
- Signs and symptoms of massive GI bleed, ruptured aortic aneurysm, or ruptured ectopic pregnancy

**WITH**

**Evidence of significant physiologic compromise:**

Age-defined hypotension* **PLUS** at least one of the following:

- Age-defined tachycardia**
- ETCO₂ less than 25
- Positive eFAST exam (if available)
- Lactate greater than 4 (if available)
- Capillary reperfusion greater than 3 sec
- Altered sensorium thought not secondary to intoxication or head trauma
- Witnessed PEA cardiac arrest less than 5 min duration

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<table>
<thead>
<tr>
<th>Age-Defined Hypotension</th>
<th>Age-Defined Tachycardia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ages less than 10 Yrs: systolic BP less than [70 + 2 x years)]</td>
<td>Age 1 year: greater than 190</td>
</tr>
<tr>
<td>Ages 10-65: systolic BP less than 90</td>
<td>Ages 2-4 yrs: greater than 140</td>
</tr>
<tr>
<td>Ages greater than 65: systolic BP less than 100</td>
<td>Ages 5-12 yrs: greater than 140</td>
</tr>
<tr>
<td>Any age: absent radial pulses</td>
<td>Ages greater than 12 years: 120</td>
</tr>
</tbody>
</table>

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*Age-Defined Hypotension  **Age-Defined Tachycardia

2. **Contraindications**

a) Patient indicates refusal to receive blood
b) Medic alert tag indicating patient objection to receiving blood

3. **Procedure**

a) Ensure applicable hemorrhage and shock interventions: tourniquet, wound-packing, pelvic binder, thoracic decompression
b) Assess for contraindications to administration of LTO+ WB

c) Obtain IV access (18 gauge or larger, if possible), and keep IV catheter hub accessible to allow direct connection of blood tubing. A large-bore IV extension set and large-bore stopcock may be utilized if available. Obtain pre-transfusion blood sample, if possible.

d) IV infusion is preferable to IO infusion for optimal flow rates.
e) Transfuse LTO+ WB
   (1) Patients less than 35 kg: Administer 10 mL/kg IV/IO
   (2) Patients greater than or equal to 35 kg: Administer 1 unit IV/IO
f) Apply Whole Blood identification bracelet to patient’s wrist or ankle
g) Assess for signs of transfusion reaction***
h) Assess for clinical improvement
   For patients with non-compressible hemorrhage: Look for signs of improved perfusion with presence of central pulses, but use permissive hypotension approach. Do not target systolic BP of greater than 90 mmHg unless significant TBI.
i) If inadequate clinical improvement, transfuse additional LTO+ WB
   (1) Patients less than 35 kg: Administer 10 mL/kg IV/IO
   (2) Patients greater than or equal to 35 kg: Administer 1 unit IV/IO
j) Assess for signs of transfusion reaction***
k) Upon hospital arrival, inform receiving team of patient’s receipt of whole blood, and provide empty whole blood bags and pre-transfusion blood sample (if obtained) for hospital blood bank evaluation.

***Possible transfusion reaction signs and symptoms: Hives, wheezing, rigors, fevers, abdominal pain, vomiting, sudden worsening of hypotension or tachycardia that is not consistent with patient’s underlying condition

If transfusion reaction is suspected:
   • Immediately discontinue the transfusion
   • Administer dexamethasone IV/IO AND diphenhydramine IV/IO in age appropriate doses

A new transfusion from a different unit of LTO+ WB may be initiated if patient reassessment indicates continued need for blood.
1. PURPOSE
To define the indications for the use of BiPAP by a paramedic for the continuation of BiPAP airway management which has been initiated prior to EMS arrival.

a) The level of care required for the interfacility transport of a “Chronic BiPAP Patient” is within the scope of a practice of a paramedic who has been credentialed, is competent, and received adequate training specific to the patient’s condition and equipment necessary to provide care.

b) Exception: A CRT-I or EMT may transport a chronically ventilated patient who is going for routine medical care and has in attendance a patient provided attendant who can manage the patient’s specific BiPAP equipment.

2. INDICATIONS

a) Patient has established BiPAP settings that have no changes within 6 hours or changes reflecting improvement in the patient’s condition.

b) Patients who are 13 years of age or older. If less than 13 years of age, patients shall be screened by SCT service (medical director or nurse) to determine the crew composition.

c) Exception: A CRT-I or EMT may transport patients of any age who are chronically on BiPAP if they are being transported for routine appointments, which are not related to a respiratory complaint. A patient-provided attendant who can manage the patient’s specific BiPAP equipment must be present during the transport.

3. CONTRAINDICATIONS
Circumstances in which endotracheal intubation or a surgical airway is indicated

4. PROCEDURE

a) Assure patent airway.

b) Perform appropriate patient assessment, including obtaining vital signs, pulse oximeter (SpO₂) reading, and cardiac rhythm.

c) Apply BiPAP device per manufacturer’s instructions.

d) Program the device to match the settings of the BiPAP machine that the patient is currently using.

e) Assess the patient after placing the BiPAP device selected for transfer. If respiratory distress occurs, support the patient with a BVM and supplemental oxygen until facility personnel reestablish therapy with original BiPAP device.

f) Continuously reassess the patient.

g) Monitor continuous pulse oximetry.

h) Monitor continuous ETCO₂ with nasal prongs, if available. (Adults only)

i) Follow the appropriate set of standing orders for continued transport.

j) Confirm the availability of a BiPAP device at the destination facility.

FOR CIRCUMSTANCES IN WHICH THE PATIENT DOES NOT IMPROVE OR CONTINUES TO DETERIORATE DESPITE BIPAP AND/OR MEDICAL THERAPY, TERMINATE BIPAP ADMINISTRATION AND PERFORM BVM VENTILATION AND ENDOTRACHEAL INTUBATION IF NECESSARY.
1. PURPOSE
   To define the indications for the initiation of BiPAP by a paramedic

2. INDICATIONS
   Patients who are 13 years of age or older, presenting with respiratory distress or failure,
   due to respiratory infection, pulmonary edema, COPD or asthma. The patient must have a
   patent, self-maintained airway and spontaneous respirations.

3. CONTRAINDICATIONS
   a) Circumstances in which endotracheal intubation or a surgical airway is indicated
   b) Patients under 13 years of age

4. PROCEDURE
   a) Ensure emergency equipment is immediately available and an alternate airway manage-
      ment plan has been established.
   b) Assure patient airway.
   c) Perform appropriate patient assessment, including obtaining vital signs, pulse oximeter
      (SpO₂) reading, and cardiac rhythm.
   d) Apply BiPAP device per manufacturer’s instructions.
   e) Set initial inspiratory positive airway pressure (IPAP) and expiratory positive airway
      pressure (EPAP) to decrease patient respiratory effort and adjust as needed.
      (1) Start with EPAP at 5 cm of water (max 8 cm of water).
      (2) Start with IPAP at 10 cm of water (max 15 cm of water).
   f) Reassess the patient after placing the BiPAP device.
   g) Monitor continuous pulse oximetry.
   h) Monitor continuous ETCO₂ with nasal prongs, if available. (Adults only)
   i) Notify the receiving facility that the patient is on a BiPAP device.
   j) If the patient is located within a healthcare facility, observe the patient on
      BiPAP at the referring facility. If the patient is not tolerating BiPAP, consult with the
      sending or receiving physician prior to transport.

FOR CIRCUMSTANCES IN WHICH THE PATIENT DOES NOT IMPROVE OR CONTINUES TO
DETERIORATE DESPITE BIPAP, TERMINATE BIPAP ADMINISTRATION AND PERFORM BVM
VENTILATION AND ENDOTRACHEAL INTUBATION IF NECESSARY.
1. **PURPOSE**
   To define the indications for the initiation of HHFNC by a Paramedic (for patients 13 years or older) or SCT Paramedic (for patients less than 13 years old).

2. **INDICATIONS**
   Patients with hypoxic respiratory failure, which may be due to the following causes: pulmonary edema, pneumonia, pulmonary embolism, pulmonary hypertension, or interstitial lung disease.

3. **CONTRAINDICATIONS**
   a) Inability to provide continuous, heated humidification using an approved delivery device.
   b) Inability to provide therapy through an appropriate sized nasal prongs
   c) Insufficient supply of oxygen to complete the transport
   d) Basilar skull fracture or severe facial trauma
   e) Circumstances in which endotracheal intubation or a surgical airway is indicated
   f) Circumstances in which the patient does not improve or continues to deteriorate despite CPAP administration

4. **PROCEDURE**
   a) Ensure that an adequate supply of oxygen is available for the transport.
      (1) Calculate the amount of oxygen needed prior to departure.
      (2) Ensure that you have at least two times the amount of oxygen anticipated.
   b) Perform appropriate patient assessment, including obtaining vital signs, pulse oximeter (SpO₂) reading, and cardiac rhythm.
   c) Set FiO₂ to maintain SpO₂ at or above 94% (or to patient’s baseline oxygen saturation, if known).
   d) Set flow rate in liters per minute (L/min) to decrease work of breathing.
      (1) Flow calculation: 2 L/kg/min up to the first 12 kg, plus 0.5 L/kg/min for each kg thereafter, up to a maximum flow rate of 60 L/min.
   e) Reassess vitals, work of breathing, mental status and breath sounds.
   f) Consider the need for escalation of respiratory support if patient remains in respiratory failure on more than 2 L/kg/min of flow or maximum settings for the delivery device.

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**FOR CIRCUMSTANCES IN WHICH THE PATIENT DOES NOT IMPROVE OR CONTINUES TO DETERIORATE ON HHFNC, TERMINATE HHFNC ADMINISTRATION AND ESCALATE RESPIRATORY SUPPORT TO PROVIDE POSITIVE PRESSURE VENTILATION VIA CPAP, BIPAP, BVM OR ENDOTRACHEAL INTUBATION, IF NECESSARY.**
Pediatric Section

g) Pediatric patients (under 13 years of age) who require HHFNC must be transported by a nurse, nurse practitioner, respiratory therapist or physician, who is credentialed, educated, and competent in dealing with the ventilator and the ventilated patient.

h) Exception: If a critical interfacility transfer is needed and a credentialed, educated, and competent healthcare clinician (as listed in “g” above) is genuinely unavailable, then a credentialed, educated and competent SCT paramedic may attend the patient with the addition of a second ALS clinician.

(1) SCT service (medical director or nurse) will screen the call prior to transport.
(2) Both the sending and receiving physicians must be in agreement with the team configuration prior to transport.
1. **PURPOSE**
   Endotracheal intubation using video laryngoscopy involves visualizing the glottic opening using specialized technology to view “around the corner” and pass the endotracheal tube, under optimal visualization, into the trachea. The purpose is to provide airway and ventilatory support for apnea, hypoxia, hypoventilatory respiratory failure, or respiratory insufficiency. The video laryngoscope device must have the following features:
   a) Color monitor
   b) Anti-fog mechanism
   c) Video recording device
   d) Appropriately-sized blade for the patient being intubated

2. **INDICATIONS**
   Video laryngoscopy and orotracheal intubation is indicated for patients who meet one or more of the following criteria and for whom appropriately-sized equipment is available:
   a) Apnea or agonal respirations
   b) Airway reflex compromised
   c) Ventilatory effort compromised
   d) Injury or illness involving the airway
   e) Potential for airway or ventilatory compromise

3. **CONTRAINDICATIONS**
   Lack of an appropriately-sized laryngoscope blade for the patient being intubated

4. **POTENTIAL ADVERSE EFFECTS/COMPPLICATIONS**
   a) Trauma to the mouth, pharynx, larynx, trachea, esophagus
   b) Right mainstem bronchus intubation
   c) Vomiting
   d) Secondary brain injury resulting from hypoxia and/or hypotension
   e) Displacement of a properly placed endotracheal tube
   f) Esophageal intubation

5. **PRECAUTIONS**
   a) Attempt visualization and endotracheal intubation up to two times. If additional attempts are indicated, consult medical direction and consider what changes would result in improved visualization and success at endotracheal placement of the ET tube.
   b) Confirm placement of the endotracheal tube in the trachea as described in Airway Management: Orotracheal Intubation protocol.

6. **PROCEDURE**
   a) Insert the Video Laryngoscope Device midline into the pharynx.
   b) Advance the Video Laryngoscope Device midline to center the vocal cords on the video screen.
   c) Pass the endotracheal tube between the vocal cords, remove the stylet, and advance the tube to the desired depth.
   d) Secure the endotracheal tube and verify correct placement.
Optional Supplemental Protocol – ANTIMICROBIAL INFUSION
FOR INTERFACILITY TRANSPORT

1. PURPOSE
During interfacility transports, a paramedic may monitor a patient on a continuous IV antimicrobial medication infusion as long as the following criteria have been met.

2. INDICATIONS
The antibiotics infusion must have been started by the hospital staff prior to an interfacility transfer. IV antimicrobial infusions may NOT be initiated by the prehospital clinician.

3. CONTRAINDICATIONS
a) Patients who have unstable vital signs or are being transferred to an intensive care environment
b) Patients with allergic reaction to infusing antibiotic agent or class
c) Pediatric patients

4. PROCEDURE
a) Follow the appropriate ALS algorithm and maintain the infusion as directed by the sending physician/practitioner.
b) The paramedic will review the sending physician’s antibiotics order and will review the specific antibiotic agent to ensure appropriate administration, indications, and absence of contraindications.
c) Unless not indicated per the medication profile, the antimicrobial infusion must be maintained on an infusion pump designed for transport, and the clinician must be trained in the appropriate use of that specific infusion pump. The ambulance must have an inverter to power the pump while in the vehicle.
d) The administration of the antibiotics infusion will be recorded on the patient care report to include the antibiotic agent’s name, dose, rate, and volume infused during transport.
e) When in doubt, contact the sending physician/practitioner for medical direction.

5. SPECIAL CONSIDERATIONS
a) The ALS service or jurisdiction must provide and document training of the ALS clinicians on the operation of infusion pump(s) being used.
b) The ALS service or jurisdiction must provide and document training of the ALS clinicians on the general administration of antimicrobials. However, due to the vast array of antimicrobials, the paramedic must utilize a practice of evaluating each patient care situation with the use of current medication reference materials to ensure appropriate administration of the infusion.
c) The ALS service or jurisdiction must also have a quality improvement (QI) program monitoring the appropriateness and quality of care provided. The QI program should be directed or coordinated by, at minimum, an ALS clinician.
1. PHARMACOLOGY
   Antimicrobials are agents that kill microorganisms or suppress their multiplication or growth.

2. PHARMACOKINETICS
   Antimicrobial agents are classified functionally according to the manner in which they adversely affect a microorganism.

3. INDICATIONS
   Treatment of known or suspected infectious disease, or as prophylaxis for an infectious process

4. CONTRAINDICATIONS
   a) Patients who have unstable vital signs or are being transferred to an intensive care environment
   b) Patients with allergic reaction to specific antibiotic agent or class
   c) Pediatric patients

5. ADVERSE EFFECTS AND PRECAUTIONS
   Antimicrobials have various adverse effects depending on the specific agent’s mechanism of action. Current medication reference materials should be consulted for specific patient situation.

6. DOSAGE
   a) Adult: Administer per practitioner order.
   b) Pediatric: Not indicated.
1. INDICATIONS

Depending on its form, cyanide can enter the body through inhalation, ingestion, or absorption through the skin. Cyanide should be suspected in occupational or smoke exposures (e.g., firefighting), industrial accidents, natural catastrophes, suicide and murder attempts, chemical warfare, and terrorism (whenever there are multiple casualties of an unclear etiology).

Non-specific and early signs of cyanide exposure (inhalation, ingestion, or absorption) include the following signs and symptoms: anxiety, vertigo, weakness, headache, tachypnea, nausea, dyspnea, vomiting, and tachycardia.

“High concentrations of cyanide” will produce:

- Markedly altered level of consciousness
- Seizure
- Respiratory depression or respiratory arrest or
- Cardiac dysrhythmia (other than sinus tachycardia)

The rapidity of onset is related to the severity of exposure (inhalation or ingestion) and may have dramatic, immediate effects causing early hypertension with subsequent hypotension, sudden cardiovascular collapse, or seizure/coma.

**Alert**

**PATIENTS WHO HAVE SUSTAINED A BURN AND/OR TRAUMATIC INJURY SHOULD BE GIVEN TREATMENT SPECIFIC TO THOSE INJURIES, INCLUDING APPLYING SPINAL PROTECTION, IF INDICATED. THE SMELL OF (BITTER) ALMONDS IS NOT A RELIABLE SIGN AND THE CLINICIAN SHOULD NOT ATTEMPT TO INHALE LOCAL AIR NOR PATIENT BREATH TO DETERMINE IF THE ALMOND SMELL IS PRESENT.**

BE SURE TO ASSESS FOR EVIDENCE OF TRAUMATIC OR MEDICAL CAUSES FOR PATIENT’S ALTERED MENTAL STATUS.

2. TREATMENT

a) Remove the patient from the source of exposure. (In the smoke inhalation victim, maintain appropriate clinician respiratory protection, SCBA.)

b) Restore or maintain airway patency.

c) Administer 100% oxygen via non-rebreather mask or bag-valve-mask.

d) Provide aggressive advanced airway management.
Optional Supplemental Protocol – CYANIDE POISONING (continued)

15.5

- e) Establish IV access with LR.
- f) Use glucometer and treat patient accordingly.
- g) There is no widely available, rapid, confirmatory cyanide blood test. Treatment decisions must be made on the basis of clinical history and signs and symptoms of cyanide intoxication. For the patient with an appropriate history and manifesting one or more of “high concentrations of cyanide” signs or symptoms:

  1. Collect a pre-treatment blood sample in the appropriate tube for lactate and cyanide levels.
  2. If patient history is suggestive of CO inhalation, follow Carbon Monoxide/Smoke Inhalation protocol
  3. If patient (adult or pediatric) has a reported oral cyanide ingestion and does not manifest signs and symptoms meeting administration criteria, medical consultation is required for administration of hydroxocobalamin (consider simultaneous consultation with poison control and medical consultation).
  4. ADULT: Administer hydroxocobalamin. Initial dose is 5 grams administered over 15 minutes SLOW IV. Each 2.5 gram vial of hydroxocobalamin for injection is to be reconstituted with 100 mL of LR and administered at 10–15 mL/minute. An additional 5 gram dose may be administered with medical consultation.
  5. PEDIATRIC: Administer hydroxocobalamin 70 mg/kg (reconstitute concentration is 25 mg/mL). Each 2.5 gram vial of hydroxocobalamin for injection is to be reconstituted with 100 mL of LR and administered at 10–15 mL/minute. Maximum single dose is 5 grams.

HYDROXOCOBALAMIN MAY CAUSE TEMPORARY RED DISCOLORATION OF THE SKIN, URINE, AND MUCOUS MEMBRANES (WHICH IS NOT TO BE CONFUSED WITH THE RARE SIGN OF CARBON MONOXIDE POISONING). THE DEVICES THAT RELY ON COLORIMETRY (E.G., PULSE OXIMETER AND CO LEVEL) WILL BE INTERFERED WITH BY THE COLOR CHANGE AND ARE NOT RELIABLE FOR PATIENT ASSESSMENT.

NOTIFY HOSPITAL OF ADMINISTRATION OF HYDROXOCOBALAMIN AND DO NOT ADMINISTER SODIUM THIOSULFATE THROUGH THE SAME IV, AS THIS MAY CAUSE CRYSTALLINE PRECIPITATION.
HYDROXOCOBALAMIN

1. PHARMACOLOGY
   Hydroxocobalamin is a form of Vitamin B-12.

2. PHARMACOKINETICS
   Hydroxocobalamin binds to the cyanide ion, forming cyanocobalamin, which is
   excreted in the urine.

3. INDICATIONS
   Signs and symptoms of high concentrations of cyanide exposure with an
   appropriate clinical history are indications for treatment as there is no widely avail-
   able, rapid, confirmatory cyanide blood test.

   “High concentrations of cyanide” will produce:
   • Markedly altered level of consciousness
   • Seizure
   • Respiratory depression or respiratory arrest or
   • Cardiac dysrhythmia (other than sinus tachycardia)

   Mechanism of action of cyanide in the body
   Effects begin within seconds of inhalation and within 30 minutes of inges-
   tion. The rapidity of onset is related to the severity of exposure (inhalation or
   ingestion) and may have dramatic, immediate effects causing sudden car-
   diovascular collapse or seizure/coma.

   Initial effects of poisoning include headache, faintness, vertigo, excitement,
   anxiety, a burning sensation in the mouth and throat, breathing difficulty,
   increased heart rate, and hypertension. Nausea, vomiting, and sweating are
   common.

   Smell of almonds is not a reliable sign and the clinician should not attempt to
   inhale local air nor patient breath to determine if the almond smell is present.

4. CONTRAINDICATIONS
   Patients with known anaphylactic reactions to hydroxocobalamin or cyanocobalamin

5. ADVERSE EFFECTS
   a) Reddish discoloration of the skin and urine (which is not to be confused with the rare
      sign of carbon monoxide poisoning). The devices that rely on colorimetry (e.g., pulse
      oximeter and CO level) will be interfered with by the color change and are not reliable
      for patient assessment.
   b) Rash
   c) Increased blood pressure
   d) Nausea
   e) Headache
   f) Decreased white cell count
6. PRECAUTIONS
   a) Notify hospital of administration of hydroxocobalamin and do not administer sodium thiosulfate through the same IV, as this may cause crystalline precipitation.
   b) Administer slowly over 15 minutes.
   c) Watch for administration site reactions.
   d) Monitor for hypertensive response to administration.

BE SURE TO ASSESS FOR EVIDENCE OF TRAUMATIC OR MEDICAL CAUSES FOR PATIENT’S ALTERED MENTAL STATUS.

7. DOSAGE
   a) Collect a pre-treatment blood sample in the appropriate tube to assess cyanide level.
   b) ADULT: Administer hydroxocobalamin. Initial dose is 5 grams administered over 15 minutes SLOW IV. (Each 2.5 gram vial of hydroxocobalamin for injection is to be reconstituted with 100 mL of LR and administered at 10–15 mL/minute.).

   An additional 5 gram dose may be administered with medical consultation.
   c) PEDIATRIC: Administer hydroxocobalamin 70 mg/kg (reconstitute concentration is 25 mg/mL). Each 2.5 gram vial of hydroxocobalamin for injection is to be reconstituted with 100 mL of LR and administered at 10–15 mL/minute. Maximum single dose 5 grams.
   d) If patient (adult or pediatric) has a reported oral cyanide ingestion and does not manifest signs and symptoms meeting administration criteria, consider medical consultation for administration of hydroxocobalamin.
Optional Supplemental Protocol – DIRECT TO TRIAGE PROTOCOL

VITAL SIGNS ACCEPTABLE? (SEE CHART – 1)
IF YES, MOVE ON

HIGHL RISK CONDITIONS? (SEE CHART – 2)
IF ANY PRESENT, STOP
IF NONE, MOVE ON

TIME DEPENDENT NEEDS? (SEE CHART – 3)
IF ANY PRESENT, STOP
IF NONE, MOVE ON

END OF CALL
• SHORT FORM COPIED AND GIVEN TO APPROPRIATE NURSE FOR RN SIGNATURES
• PATIENT TRANSFERRED OFF STRETCHER

PATIENT PLACED DIRECTLY IN WAITING ROOM VIA WHEELCHAIR, AT REGISTRATION, SIGNATURES OBTAINED AND PATIENT IS LEFT WITH MIEMSS APPROVED SHORT FORM

DISCUSSION TAKES PLACE WITH PATIENT ABOUT PLACEMENT IN Triage

ACCEPtable VITAL SIGNS:
• RESPIRATIONS: 10-20
• PULSE: 60-100
• PULSE OX: >92% (room air)
• TEMPERATURE: 96-101°F
• BLOOD GLUCOSE (if indicated): 71-299 MG/DL
• BLOOD PRESSURES:
  • BETWEEN 110 AND 180 - SYSTOLIC
  • BETWEEN 60 AND 100 – DIASTOLIC

HIGh RISK CONDITIONS
• UNCREASED ABDOMINAL PAIN
• ALTERED MENTAL STATUS
• UNCREASED BACK PAIN
• CHEST PAIN
• DYSPEA / SHORTNESS OF BREATH
• (ACUTE) FOCAL NEUROLOGICAL DEFICITS
• SEIZURES
• SIPHOS (SUSPECTED)
• SYNCOPE
• SUICIDAL / HOMICIDAL IDEATIONS
• REQUIRES MORE THAN MINIMAL ASSISTANCE TO WALK
• UNABLE TO COOPERATE WITH HISTORY AND EXAM

TIME DEPENDENT NEEDS
• AIRWAY
• BREATHING
• CIRCULATION (INCLUDING TO EXTREMITY)
• DISABILITY (DEFICIT) OR DEFORMITY
• SEVERE TENDERNESS WITH PALPATION / EXAM
• SIGNIFICANT HEAD OR TRUNCAL TRAUMA
• UNCONTROLLABLE BLEEDING
• REQUIRES ALS MONITORING OR INTERVENTIONS
• CONCERN FOR POTENTIAL DETERIORATION

Approved 10.12.2021
1. PURPOSE
The goal of this program is to allow an EMT to acquire and transmit a 12-lead (15-lead if trained to perform) electrocardiogram (EKG) to the receiving facility and possibly reduce the door to reperfusion time for the AMI patient.

2. PRESENTATION
Chest discomfort that may radiate to the arm, shoulders, jaw, or back. Generally described as a crushing pain or toothache. May be accompanied by shortness of breath, sweating, nausea, or vomiting.

OR

a) Chest discomfort. Some heart attacks involve discomfort in the center of the chest that lasts for more than a few minutes or that goes away and comes back. This discomfort can feel like uncomfortable pressure, squeezing, or fullness.

b) Discomfort in other areas of the upper body. Symptoms can include discomfort in one or both arms or in the back, neck, jaw, or stomach.

c) Shortness of breath. This symptom often accompanies chest discomfort. However, it can also occur prior to the chest discomfort.

d) Other signs. These may include breaking out in a cold sweat, nausea, light-headedness, or a sense of impending doom.

e) Post-cardiac arrest with ROSC.

f) Medical history and contributing factors.

(1) A previous heart attack or procedure to open up coronary arteries
(2) Family history of heart disease
(3) Diabetes mellitus
(4) High blood pressure
(5) High blood cholesterol
(6) Overweight
(7) Physical inactivity
(8) Cigarette smoking

3. INDICATIONS
Any patient complaining of chest discomfort or exhibiting signs, symptoms, or medical history as outlined in Section 2 (Presentation)

4. CONTRAINDICATIONS
Acquisition of a 12-lead EKG should not take precedence over required life-saving measures (e.g., CPR, assisting respirations, clearing or maintaining a patient’s airway, checking blood glucose, extrication, or removing a patient from a dangerous scene).
5. PROCEDURE
   a) Initiate General Patient Care.
   
   b) Initiate *Cardiac Emergencies: Chest Pain* protocol.
   
   c) Position patient (1) (2).
   
   d) Place chest and limb leads (3) (4).
   
   e) Turn on monitor.
   
   f) Set patient age and a patient identifier.
   
   g) Acquire 12-lead (5).
   
   h) Consult with receiving facility.
   
   i) Transmit 12-lead (6).
   
   j) Continue patient care.

(1) Unrestricted access to the skin in the chest area, arms, and lower legs is required to allow for correct placement of electrodes. Do your best to protect the patient’s privacy. Once the electrodes are positioned and connecting leads are appropriately attached, the patient should be covered with a sheet to preserve their dignity during the procedure.

(2) If unable to place patient in the recumbent position, include this information in your hospital consult and note it in the written narrative of your patient care report.

(3) Remove electrodes from a sealed package immediately before use. Using previously unpacked electrodes or electrodes with expired date codes may impair EKG signal quality.

(4) When placing electrodes on female patients, always place the leads V3-V6 under the breast rather than on the breast.

(5) Acquisition of a 12-lead EKG should take no more than 5 minutes.

(6) Transmission of the 12-lead EKG to the receiving facility should be done en route to the receiving facility. There is no need to delay transport to transmit a 12-lead EKG.
6. **INDIVIDUAL EMT APPROVAL FOR PARTICIPATION**
   a) The EMT 12-Lead EKG Program is open to all Maryland EMTs that have been providing direct patient care for a minimum of one year.
   
   b) Clinicians must be members of an ALS company that currently owns a local system compatible 12-lead device.

7. **ONGOING DEMONSTRATION OF PROFICIENCY**
   After the initial training program is completed, the EMT will participate in an annual refresh-er training program.

8. **REVIEW OF EACH CALL**
   The clinician will submit copies of each 12-lead EKG and patient care report to their jurisdictional Quality Review Committee.
Optional Supplemental Protocol – GLYCOPROTEIN IIb/IIIa ANTAGONIST INFUSION

(Paramedic only)

1. PURPOSE

During interfacility transports, a paramedic may monitor a patient on a continuous IV glycoprotein IIb/IIIa infusion as long as the following criteria have been met.

2. INDICATIONS

The glycoprotein IIb/IIIa infusion must have been started by the hospital staff prior to an interfacility transfer. IV glycoprotein IIb/IIIa transports may NOT be started by the prehospital clinician in the prehospital setting.

3. CONTRAINDICATIONS

a) Patients who are clinically unstable, including but not limited to unstable vital signs and blood pressure, or current arrhythmia

b) Pediatric patients

4. PROCEDURE

a) Maintain the infusion as directed by the sending physician.

b) The sending physician must document the infusion to be administered on the patient’s transport record or transport note. This includes the concentration of the medication and the infusion rate.

c) The infusion must be maintained on an infusion pump designed for transport. The clinician must be trained in the appropriate use of the specific infusion pump. The ambulance must have an inverter to power the pump while in the vehicle.

d) The total volume of glycoprotein IIb/IIIa infused must be recorded on the patient care report.

e) The patient must be on a cardiac monitor and vital signs should be documented on the patient care report at least every 15 minutes.

f) When in doubt, contact the sending physician for medical direction.

5. SPECIAL CONSIDERATIONS

The ALS service or jurisdiction must provide and document training of the ALS clinicians on the operation of the infusion pump(s) being used. They must also have a quality improvement (QI) program monitoring the appropriateness and quality of care provided. The QI program should be directed or coordinated by, at minimum, a paramedic.
Optional Supplemental Protocol –
GLYCOPROTEIN IIb/IIIa ANTAGONIST INFUSION (continued)

(Paramedic only)

1. PHARMACOLOGY
   Platelet glycoprotein antagonist. This agent reversibly prevents fibrinogen and von Willenbrand’s factor from binding to the glycoprotein IIb/IIIa receptor, inhibiting platelet aggregation.

2. PHARMACOKINETICS
   Glycoprotein IIb/IIIa has a half-life of 2.5 hours. Metabolism of this drug is limited and is excreted via the kidneys.

3. INDICATIONS
   Patients with acute coronary syndrome including those with Percutaneous Coronary Intervention (PCI)

4. CONTRAINDICATIONS
   a) Hypersensitivity, active internal bleeding, history of bleeding, stroke within one month, major surgery with severe trauma, severe hypotension, history of intracranial bleeding, intracranial neoplasm, arteriovenous malformation/aneurysm, aortic dissection, or dependence on renal dialysis
   b) Pediatric patients

5. SIDE EFFECTS/ADVERSE REACTIONS
   a) Cardiovascular: Stroke, hypotension
   b) Systemic: Bleeding, anaphylaxis
   c) Other: Hematuria, thrombocytopenia

6. PRECAUTIONS
   Glycoprotein IIb/IIIa is a medication designed to inhibit the clotting factor in blood. Patients on this medication should be protected from further injuries that may cause bleeding. Attempts to start IVs should not be made without a doctor’s orders.

7. DOSAGE
   a) INITIAL BOLUS: Given at sending facility and should be documented.
   b) MAINTENANCE IV DRIP: Follow Standard Dosing. Maintain drip based on patient weight and sending physician’s orders.

IF CHEST PAIN OR HYPOTENSION DEVELOPS DURING TRANSPORT, THE PARAMEDIC MUST CONSULT WITH EITHER THE SENDING OR RECEIVING PHYSICIAN FOR FURTHER INSTRUCTIONS.
Optional Supplemental Protocol –
HEPARIN INFUSION FOR INTERFACILITY TRANSPORT

(Paramedic only)

1. PURPOSE

During interfacility transports, a paramedic may monitor a patient on a continuous IV heparin infusion as long as the following criteria have been met.

2. INDICATIONS

The heparin infusion must have been started by the hospital staff prior to an interfacility transfer. IV heparin infusions may NOT be started by the prehospital clinician in the prehospital setting.

3. CONTRAINDICATIONS

a) Patients who have had trauma or surgery to the brain, eye, spinal cord, urinary tract, joints, or retroperitoneum within the last 7 days
b) Patients with active bleeding
c) Third trimester pregnancy

4. PROCEDURE

a) Follow the appropriate ALS algorithm and maintain the infusion as directed by the sending physician.
b) The sending physician must document the infusion to be administered on the patient’s record or transport note, including the concentration of the units per hour.
c) The infusion must be maintained on an infusion pump designed for transport, and the clinician must be trained in the appropriate use of that specific infusion pump. The ambulance must have an inverter to power the pump while in the vehicle.
d) The total volume of heparin infused must be recorded on the patient care report.
e) The patient must be on a cardiac monitor and vital signs should be documented on the patient care report every 15 minutes.
f) When in doubt, contact the sending physician for medical direction.

5. SPECIAL CONSIDERATIONS

The ALS service or jurisdiction must provide and document the training of ALS clinicians on the operation of the infusion pump(s) being used. They must also have a quality improvement (QI) program monitoring the appropriateness and quality of care provided. The QI program should be directed or coordinated by, at minimum, an ALS clinician.
1. PHARMACOLOGY
   Heparin is an anticoagulant that works by neutralizing several of the clotting factors
   (XIII, XII, XI, X, IX, and II).

2. PHARMACOKINETICS
   a) Heparin inhibits the coagulation mechanism in 3 sites:
      (1) activation of factor X
      (2) formation of thrombin from prothrombin
      (3) conversion of fibrinogen to fibrin
   b) Heparin’s effect, which is to retard or prevent blood clotting, is immediate. The half-life
      of intravenous heparin is 1–1.5 hours.

3. INDICATIONS
   a) Thromboembolic disease, such as pulmonary embolism deep vein thrombophlebitis,
      and arterial embolization
   b) Acute myocardial infarction. (Heparin may be given alone or in conjunction with throm-
      bolytic therapy.)

4. CONTRAINDICATIONS
   a) Patients who have had trauma or surgery to the brain, eye, spinal cord, urinary tract,
      joints, or retroperitoneum within the last 7 days
   b) Patients with active bleeding
   c) Third trimester pregnancy

5. ADVERSE EFFECTS
   Increased potential for bleeding

6. PRECAUTIONS
   a) Inadvertent infusion of too much heparin can result in over-anticoagulation and the
      potential for bleeding complications.
   b) If it is necessary to draw blood or start an IV while a patient is receiving heparin, extra
      time to hold pressure over the puncture site will be necessary to stop the bleeding.
   c) Use with caution for patients with extreme hypertension.

7. DOSAGE
   a) Adult: Follow the written order from the sending physician. Paramedic may
      transport patients at a maximum heparin drip rate of 18 units/kg per hour or
      2,000 per units per hour, whichever is higher. For doses exceeding the maxi-
      mum, a SCT Paramedic or nurse is required for transport.
   b) Pediatric: Not indicated.
1. INDICATIONS

An exposure to hydrofluoric acid presenting with any of the following:
   a) Severe throbbing pain at sites of exposure
   b) Redness, edema, blistering burns to the skin or blanched white areas of skin
   c) Cough, stridor, bronchospasm, pulmonary edema
   d) Hypotension
   e) Dyrsrhythmias or electrolyte imbalances (hypocalcemia, hypomagnesemia, hyperkalemia)

2. BLS
   a) Maintain scene safety.
   b) Remove patient from toxic environment using appropriate PPE.
   c) Decontaminate and remove patient's clothing, if contaminated.
   d) Thoroughly irrigate all affected areas with copious water, normal saline, or Lactated Ringer's for a minimum of 15 minutes.
   e) Evaluate for trauma.
   f) Check blood glucose and treat per Hypoglycemia protocol.

3. EYE EXPOSURE

   Continuously flush eyes with Lactated Ringer's, normal saline, or sterile water for a minimum of 15 minutes (or 1 liter) via nasal cannula tubing or manual irrigation.

4. TOPICAL EXPOSURE
   a) Apply 2.5% USP calcium gluconate gel (Calgonate®) to affected area(s) until cessation of pain, if available. Reapply to affected area of skin every 15 minutes.
   b) If hands or fingers are involved, apply the calcium gluconate gel to the hand. Place additional calcium gluconate gel into a medical glove, and then insert the affected hand into the glove and massage.

5. ALS
   a) Establish IV access, if possible, in an unaffected area of the body.
   b) Assess for and treat hypotension and hypoxia.
   c) Perform continuous cardiac monitoring and treat any dysrhythmias associated with electrolyte imbalance (hypocalcemia, hypomagnesemia, hyperkalemia).
   d) Obtain 12-lead EKG, if situation allows.
   e) Consult with poison control and burn center (or trauma center, if traumatic injury is also present).

6. EYE EXPOSURE
   Irrigate with Lactated Ringer’s, normal saline, or sterile water via nasal cannula tubing or manual irrigation.
7. **RESPIRATORY EXPOSURE**
   If respiratory distress, cough and/or wheezing, administer 4mL of nebulized 2.5% calcium gluconate solution every 5 minutes as needed (2.5mL 10% calcium gluconate solution in 7.5mL normal saline), if available.

8. **SYSTEMIC TOXICITY**
   If signs of systemic toxicity, including hypotension, altered mental status, severe respiratory distress, ventricular dysrhythmia, or signs of hypocalcemia or hyperkalemia:

   **Calcium gluconate**
   a) Adult dosing: Calcium gluconate 3 grams IV/IO of a 10% solution over 5 minutes. If symptoms persist, may repeat a dose of calcium gluconate once in 10 minutes.
   b) Pediatric dosing: Calcium gluconate 60 mg/kg IV/IO of a 10% solution (max dose: 2,000 mg) over 5 minutes. If symptoms persist, may repeat a dose of calcium gluconate once in 10 minutes.

   May repeat dose of calcium gluconate once in 10 minutes, if symptoms persist.

   and

   **Magnesium sulfate**
   a) Adult dosing: 2 grams IV/IO over 10 minutes
   b) Pediatric dosing: 50 mg/kg/dose IV/IO (max dose: 2,000 mg) over 10 minutes

9. **CLINICAL PEARLS**
   a) Pain may be disproportionate to physical exam findings.
   b) Effects from more dilute HF acids or its vapors may be delayed. Relief of pain is an excellent indication of the success of treatment.
   c) Calcium chloride and calcium gluconate are NOT the same medication and are NOT interchangeable.
   d) HF exposure may result from industrial incidents, fires involving lithium ion batteries, glass etching, metal cleaning and electronics manufacturing, or exposure to rust remover solutions or brick cleaning solutions.
   e) HF differs from other acids because the fluoride ion quickly penetrates the skin, causing destruction of deep tissue layers.
   f) Burns larger than 25 square inches (160 square cm) may result in serious systemic toxicity.
   g) The Immediately Dangerous to Life or Health level of hydrofluoric acid is 30 ppm.
15.10

Optional Supplemental Protocol –
HYDROFLUORIC (HF) ACID EXPOSURE (continued)

Calcium Gluconate

1. Pharmacology
   a) Increases serum calcium
   b) Combines with hydrofluoric acid (HF) to prevent extraction of calcium from tissues and bones
   c) Reduces risk of cardiac arrhythmias due to hyperkalemia

2. Pharmacokinetics
   Rapid onset of action with IV administration

3. Indications
   Hydrofluoric acid exposure; see specific indications under “dosing” below

4. Contraindications
   Hypercalcemia

5. Adverse Effects
   a) Bradycardia, hypotension and cardiac arrhythmias, leading to syncope or cardiac arrest, may occur with rapid administration
   b) Tissue necrosis and calcinosis, particularly with extravasation from intravenous catheter

6. Precautions
   a) Use with caution on patients taking digitalis, as calcium may increase ventricular irritability and precipitate digitalis toxicity.
   b) If given with sodium bicarbonate, calcium will precipitate; flush IV between administering these medications.

7. Dosage
   a) Respiratory exposure / distress
      If respiratory distress, cough and/or wheezing: administer 4 mL of nebulized 2.5% calcium gluconate solution every 5 minutes as needed up to a maximum of 3 doses (2.5 mL 10% calcium gluconate solution in 7.5 mL normal saline), if available.

   b) Systemic toxicity
      If signs of systemic toxicity, including hypotension, altered mental status, severe respiratory distress, ventricular dysrhythmia, or signs of hypocalcemia or hyperkalemia:
      (1) Adult dosing: Calcium gluconate 3 grams IV/IO of a 10% solution over 5 minutes.
         If symptoms persist, may repeat a dose of calcium gluconate once in 10 minutes.
Optional Supplemental Protocol –
HYDROFLUORIC (HF) ACID EXPOSURE (continued)

15.10

(2) Pediatric dosing: Calcium gluconate 60 mg/kg IV/IO of a 10% solution (max
dose: 2,000 mg) over 5 minutes. If symptoms persist, may repeat a dose of
calcium gluconate once in 10 minutes.

c) Topical exposure

(1) Apply 2.5% USP calcium gluconate topical (Calgonate®) to affected area(s) until
cessation of pain, if available. Reapply to affected area of skin every 15 minutes.

(2) If hands or fingers are involved, apply the calcium gluconate gel to the hand. Place
additional calcium gluconate gel into a medical glove, and then insert the affected
hand into the glove and massage.
Optional Supplemental Protocol – INTRANASAL NALOXONE FOR COMMERCIAL SERVICES BLS CLINICIANS

July 2018: Naloxone is required for Public Safety EMT and EMR (October ’17) and remains Optional Supplemental Program for BLS Commercial Services (initially implemented September ’13).

1. PURPOSE
When encountered with a patient exhibiting respiratory depression with a confirmed or suspected opioid/narcotic overdose, an EMT and EMR may administer intranasal naloxone provided the following criteria have been met.

2. INDICATIONS
A patient suffering respiratory depression caused by a known or suspected opioid/narcotic overdose

3. CONTRAINDICATIONS
a) None clinically significant in the adult patient
b) Patients less than 28 days old

4. PROCEDURE
a) Ensure that naloxone is indicated and the medication is not expired.
b) Inject volume of air into vial that is equal to desired volume of medication to be removed using a needle (blunt tip preferred) and 2 mL or 3 mL syringe.
c) Pull back on syringe plunger to remove desired volume of medication.
d) Use gradations on syringe to measure volume of medication to nearest 0.10 mL.
e) Safely remove needle from syringe and dispose of in sharps container.
f) Attach mucosal atomization device to luer-lock of syringe.
g) Place tip of mucosal atomization device in the nare and briskly push the plunger forward, administering half of the total volume of medication (up to a MAXIMUM of 1 mL per nare).
h) Repeat previous step in the other nare, delivering the remaining half of the medication.
i) Monitor patient for response and continue supportive care.

IF EMS OPERATIONAL PROGRAM USES A DIFFERENT FORMULARY/CONCENTRATION OR MEDICATION PACKAGING (E.G., PRE-FILLED SYRINGE OR AMPULE), CLINICIANS MUST RECEIVE PROPER TRAINING REGARDING SAFETY, PREPARATION, AND CONVERSION TO INTRANASAL ATOMIZATION OF THE MEDICATION.
Naloxone (Narcan®)

1. PHARMACOLOGY
   Reverses all effects due to opioid (morphine-like) agents. This drug will reverse the respiratory depression and all central and peripheral nervous system effects.

2. PHARMACOKINETICS
   a) Onset of action is within a few minutes with intranasal (IN) administration.
   b) Patients responding to naloxone may require additional doses and transportation to the hospital since most opioids/narcotics last longer than naloxone.
   c) Has no effect in the absence of opioid/narcotic.

3. INDICATIONS
   To reverse respiratory depression induced by opioid/narcotic agent

4. CONTRAINDICATIONS
   Patients under 28 days of age

5. ADVERSE EFFECTS
   Opioid withdrawal

6. PRECAUTIONS
   a) Naloxone may induce opiate withdrawal in patients who are physically dependent on opioids.
   b) Certain drugs may require much higher doses of naloxone for reversal than are currently used.
   c) Should be administered and titrated so respiratory efforts return, but not intended to restore full consciousness.
   d) Intranasal naloxone must be administered via nasal atomizer.
   e) Naloxone has a duration of action of 40 minutes; the effect of the opioid/narcotic may last longer than naloxone and patients should be encouraged to be transported.

CLINICIANS MUST CONTACT A BASE STATION PHYSICIAN FOR PATIENTS WISHING TO REFUSE TRANSPORT AFTER BLS ADMINISTRATION OF NALOXONE.

7. DOSAGE
   a) Adult: Administer naloxone 2 mg IN, dividing administration of the dose equally between the nares to a maximum of 1 mL per nare, OR administer 4 mg/0.1 mL IN in one nare.
   b) Pediatric:
      (1) Child aged 28 days to adult: Administer 2 mg IN, dividing administration of the dose equally between the nares to a maximum of 1 mL per nare, OR administer 4 mg/0.1 mL IN in one nare.
      (2) Child less than 28 days: Not Indicated

Repeat as necessary to maintain respiratory activity.
1. PURPOSE
In order to provide precise medication dosing, a paramedic with jurisdictionally approved training may administer select medications and crystalloid fluids via an IV infusion pump.

2. INDICATIONS
a) An IV infusion pump, when available, should be used for infusion of weight-based medications in The Maryland Medical Protocols for Emergency Medical Services, including but not limited to vasopressors and antiarrhythmics.
b) It may also be used to administer specific volumes of crystalloid fluids.

3. CONTRAINDICATIONS
The clinician has not completed appropriate device-specific training.

4. PROCEDURE
a) Verify IV infusion pump has sufficient battery charge and that a charging cable is readily accessible.
b) Select medication to be administered from IV infusion pump drug library.
c) If medication is weight-based, ensure that accurate weight has been entered.
d) Confirm appropriate drug concentration.
e) If medication is not in drug library, enter required fields manually; this should be confirmed by a second EMS clinician whenever possible.
f) Check that IV tubing is primed, either manually or via IV infusion pump.
g) Confirm that there is no air in the line prior to attaching tubing to patient.
h) Attach tubing to patient.
i) Infuse medication or crystalloid fluid.

5. SPECIAL CONSIDERATIONS
a) The clinician should be familiar and comfortable with how to troubleshoot all alarms.
b) Slow IV push medications may be administered via syringe attached to pump tubing.
1. INDICATIONS
a) Nerve agents are a group of highly toxic chemicals that may be released in a WMD event. These agents act to inhibit cholinesterase, and therefore prolong the effects of acetylcholine. These agents are potent, long-acting, and all bind to acetylcholine irreversibly unless an oxime is given.
b) Nerve agents include Tabun (GA), Sarin (GB), Soman (GD) and GF. There are also V agents, such as VX.
c) The G-type agents evaporate (become vapor) or may be dispersed in the air by weapons. When a person inhales this vapor, effects begin within seconds to minutes.
d) The V-type agents are oily and evaporate very slowly. They persist on the ground, foliage, etc., for long periods. Exposure to this liquid on the skin causes effects to start as soon as 10 minutes or as long as 18 hours after contact. The vapor hazard from these is not as great as from the G-type agents.
e) Many insecticides currently in use are organophosphates and are chemically related to nerve agents. The organophosphate insecticides may have a slower onset and a longer lasting effect compared with nerve agents.
f) Characteristic signs and symptoms may identify nerve agent poisoning. After vapor exposure, early manifestations of poisoning occur in the eyes, nose, and airway. With liquid/dermal contact exposure, early manifestations occur in the skin and the GI tract. Thus, when looking at the chart below, consider the mechanism of release and the associated signs and symptoms (refer to the chart below with the mnemonic P-SLUDGE-MC). (NOTE: This mnemonic is used for all organophosphate toxicity. Pupillary response occurs only with vapor exposure and will not be seen unless there is direct liquid contact with the eye. Urinary incontinence is also very rare.)

<table>
<thead>
<tr>
<th>Nerve Agents</th>
<th>Signs and Symptoms of Chemical Agents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vapor Exposure</td>
<td>Liquid Exposure</td>
</tr>
<tr>
<td>P - Pinpointing pupils</td>
<td>✓</td>
</tr>
<tr>
<td>S - Salivation</td>
<td>✓</td>
</tr>
<tr>
<td>L - Lacrimation (tearing)</td>
<td>✓</td>
</tr>
<tr>
<td>U - Urination</td>
<td>✓</td>
</tr>
<tr>
<td>D - Defecation</td>
<td>✓</td>
</tr>
<tr>
<td>G - Gastrointestinal; pain/gas</td>
<td>✓</td>
</tr>
<tr>
<td>E - Emesis (vomiting)</td>
<td>✓</td>
</tr>
<tr>
<td>M - Muscle twitching</td>
<td>✓</td>
</tr>
<tr>
<td>C - Convulsions</td>
<td>✓</td>
</tr>
<tr>
<td>B - Bradycardia</td>
<td>✓</td>
</tr>
<tr>
<td>B - Bronchospasm</td>
<td>✓</td>
</tr>
<tr>
<td>B - Bronchorrhea</td>
<td>✓</td>
</tr>
</tbody>
</table>
g) EMS clinicians must know the following MILD, MODERATE, and SEVERE signs and symptoms of nerve agent poisoning. When clinicians recognize most or all of the symptoms listed below, they must IMMEDIATELY receive treatment (first aid or buddy aid).

(2) MILD poisoning (self-aid). Casualties with mild symptoms may experience most or all of the following:
   (a) Unexplained runny nose
   (b) Unexplained sudden headache
   (c) Sudden drooling
   (d) Difficulty in seeing (dimness of vision, constricted pupil)
   (e) Tightness in the chest or difficulty in breathing
   (f) Wheezing and coughing
   (g) Localized sweating and muscular twitching in the area of the contaminated skin
   (h) Stomach cramps
   (i) Nausea without vomiting

(2) MODERATE effects would be the above, but also include more severe effects such as diarrhea, moderate to severe difficulty breathing, and some skeletal-muscular twitching/fasciculations. The progression of symptoms from mild to moderate indicates either inadequate treatment or continuing exposure to the nerve agent.

(3) SEVERE symptoms. Clinicians with severe symptoms will not be able to treat themselves and must receive prompt buddy aid and medical treatment. Casualties with severe symptoms may experience most or all of the MILD symptoms plus most or all of the following:
   (a) Impaired thinking
   (b) Increasing wheezing and increased difficulty breathing
   (c) Severe pinpoint pupils
   (d) Red eyes with tearing
   (e) Vomiting
   (f) Severe muscular twitching and general weakness
   (g) Involuntary defecation
   (h) Convulsions
   (i) Unconsciousness
   (j) Respiratory Failure
   (k) Bradycardia

h) Prevention of poisoning

(1) In the setting of an exposure to a nerve agent, the most rapid absorption occurs through the respiratory tract. When it is suddenly determined that clinicians are in the “hot zone”, do not look for the invisible vapor cloud. Clinicians should hold their breath until they don and clear their breathing apparatus or protective masks. Once masked, a clinician will then give the alarm to other clinicians. This may be done with hand signals or through the mask. If a fellow clinician is severely poisoned with altered consciousness in the hot zone, the initial, less-poisoned masked clinician should mask the casualty.
Optional Supplemental Protocol – MARK I/DuoDote Kits (ATROPINE and 2-PAM AUTO-INJECTORS) (continued)

(2) When the masked casualty is severely poisoned after exposure to vapor and liquid, they should be decontaminated by removing clothing, blotting the agent (if a liquid exposure), and diluting the agent by using a flush with large amounts of water. Decontamination should be done as soon as possible, but it will usually occur in the warm zone or a safe area.

(3) When treating a severely poisoned casualty, the treating clinician should take care to avoid exposure to the liquid agent (which could occur when kneeling next to the casualty). Squatting next to the casualty while masking or treating him/her will help the caregiver to avoid exposure to liquid nerve agent.

(4) Do not administer nerve agent antidotes before actual exposure to nerve agents or development of clinical symptoms occurs. Nerve agent antidotes may degrade performance in the hot zone (creating a heat-stressed clinician) and should be administered only when symptoms and signs of nerve agent poisoning are present.

2. TREATMENT

a) The ABC priorities of prehospital treatment require modification to AABCs (Antidote then ABCs). The antidote (Atropine and 2-PAM) should be given as soon as possible, because toxic exposure to the nerve agent will make ventilation difficult. If the antidote is not immediately available, prevent further exposure to the nerve agent, provide ABC support, and evacuate the patient to an area where the antidote is available.

b) Based on signs and symptoms in a mass casualty incident (MCI) or on-site chemical testing that confirms nerve or organophosphate agent presence in a mass casualty incident, a certified EMR or EMT may administer MARK I/DuoDote kits (up to total of three kits) as buddy care to public safety personnel or when directed to do so by an ALS clinician. The midazolam 5 mg or diazepam 10 mg auto-injector (CANA) can only be administered by an ALS clinician when three MARK I/DuoDote kits are administered in a severe exposure. Medical consultation is not required in these situations.
Optional Supplemental Protocol –
MARK I/DuoDote Kits (ATROPINE and 2-PAM AUTO-INJECTORS) (continued)

**CHEMPACK**
Adult Nerve Agent Exposure Treatment

**Mild Exposure**
Patients who can walk and talk who may present with miosis, rhinorhea, increased salivation, nausea
- Atropine 2mg IM + Pralidoxime 600mg IM

**Moderate Exposure**
Patients with mild dyspnea, ataxia, miosis, or muscle cramping
- Atropine 4mg IM + Pralidoxime 1200mg IM

**Severe Exposure**
Patients who may have severe respiratory distress, seizures, extreme SLUDGEM
(See below)
- Atropine 6mg IM + Pralidoxime 1800mg IM + Diazepam 10mg IM OR Midazolam 10mg IM

May repeat 3-5 minutes until symptoms resolve

***AUTO-INJECTORS SHOULD BE USED FOR ALL ADULT EMS PATIENTS***

Medications may come packaged as either DuoDotes, Mark I Kits, ATNAA kits, individual Atropen + 600mg Pralidoxime Autoinjectors, or in individual medication vials
2mg Atropens are not available in all CHEMPACK caches

**Adult Vial Medication Directions:**
- **Atropine** (0.4mg/ml in 20mL): Draw up medication in 5mL syringe (5mL)
- **Pralidoxime** (300mg/mL): For Intramuscular (IM) injection: Add 3.3mL of sterile water into a single 1-gram vial, which results in a 300mg/mL concentration. Do not exceed 2mL per IM injection
- **Diazepam & Midazolam** (5mg/mL in 10mL): draw up 2mL in 3mL syringe for IM administration for initial dose of 10mg IM

S= Salivation
L= Lacrimation (tear production)
U= Urination
D= Defecation
G= Gastrointestinal distress
E= Emesis
M= Muscle Twitching & Miosis
(constricted pupils)
Optional Supplemental Protocol – MARK I/DuoDote Kits (ATROPINE and 2-PAM AUTO-INJECTORS) (continued)

15.13

CHEMPACK
Pediatric Nerve Agent Exposure Treatment

Mild Exposure
Patients who can walk and talk who may present with miosis, rhinorrhea, increased salivation, and nausea

<table>
<thead>
<tr>
<th>Weight (Kg)</th>
<th>Atropine Dose (IM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 10 kg</td>
<td>0.5mg</td>
</tr>
<tr>
<td>10 kg - 25 kg</td>
<td>1 mg</td>
</tr>
<tr>
<td>26 kg - 50 kg</td>
<td>2 mg</td>
</tr>
<tr>
<td>Over 50 kg</td>
<td>2 mg</td>
</tr>
</tbody>
</table>

Moderate Exposure
Patients with mild dyspnea, ataxia, miosis, or muscle cramping

<table>
<thead>
<tr>
<th>Weight (Kg)</th>
<th>Dose (IM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 10 kg</td>
<td>Atropine 1mg + Pralidoxime 600mg</td>
</tr>
<tr>
<td>10 kg - 25 kg</td>
<td>Atropine 2mg + Pralidoxime 600mg</td>
</tr>
<tr>
<td>26 kg - 50 kg</td>
<td>Atropine 2mg + Pralidoxime 1200mg</td>
</tr>
<tr>
<td>Above 50 kg</td>
<td>Atropine 2mg + Pralidoxime 1200mg</td>
</tr>
</tbody>
</table>

May repeat 3-5 minutes until symptoms resolve

Severe Exposure
Patients who may have severe respiratory distress, seizures, extreme SLUDGEM
(See below)

<table>
<thead>
<tr>
<th>Weight (Kg)</th>
<th>Dose (IM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 25 kg</td>
<td>Atropine 2mg + Peralidoxime 600mg + Diazepam 2.5mg OR Midazolam 2.5mg</td>
</tr>
<tr>
<td>26 kg - 50 kg</td>
<td>Atropine 4mg + Peralidoxime 1200mg + Diazepam 5mg OR Midazolam 5mg</td>
</tr>
<tr>
<td>Above 50 kg</td>
<td>Atropine 6mg + Peralidoxime 1800mg + Diazepam 10mg OR Midazolam 10mg</td>
</tr>
</tbody>
</table>

May repeat 3-5 minutes until symptoms resolve

Atropine/Pralidoxime may come packaged as either DuoDotes, Mark I Kits, ATNAA kits, individual Atropine + Peralidoxime autoinjectors, or in individual medication vials.

Treatment via Atropine & Peralidoxime Autoinjectors is preferred

CANA autoinjectors are not indicated for pediatric patients less than 50kg

Pediatric Vial Medication Instructions:
Atropine (0.4mg/mL, 20mL): Draw-up medication in 3mL, 5mL, or 10mL syringe as indicated
Pralidoxime (300mg/mL) Add 3.3mL of sterile water into a single 1 gram vial, which results in a 300mg/mL concentration. Do not exceed 2mL per IM injection
Diazepam & Midazolam (5mg/mL, 10mL) Draw 0.2mg/kg IM to a maximum 10mg

Color Coding and unit amount for Atropens

- 0.5 mg auto-injector (blue)
- 1 mg auto-injector (red)
- 2 mg auto-injector (green)

(S: Salivation
L: Lacrimation (tear production)
U: Uridnation
D: Defecation
G: Gastrointestinal distress
E: Emesis
M: Muscle Twitching & Miosis (constricted pupils))
Scope of practice for paramedic personnel has been expanded to allow select immunization and Purified Protein Derivative (PPD) testing by paramedic personnel. The immunizations that are allowed to be performed include Hepatitis B, Influenza, COVID-19, and PPD. This program is an optional supplemental protocol requiring the EMSOP medical director and the EMSOP to authorize select trained paramedic personnel to perform these functions. There are program requirements below. Please note that you must have a written memorandum of understanding between your EMS service and the local health department or hospital before this program can be instituted.

In order to become recognized and authorized to implement the immunization and testing program for paramedics, you must complete the application and submit a copy of the health department memorandum of understanding to the Office of the State EMS Medical Director. At that time, you will receive a copy of all of the pertinent documents and instructional material. Your jurisdiction will then be recognized as an authorized optional immunization and testing jurisdiction.

When you are implementing this program, we strongly encourage you to advise EMS personnel at risk to seek vaccination where possible.

**REQUIREMENTS**

1. Medical Director: Must have a jurisdictional medical director who is willing to take responsibility for the program.
2. Must be under the Infection Control Program for the Jurisdiction.
3. Immunization record form with documentation of all pertinent information about vaccination or test, including the patient’s primary care practitioner.
4. Direct linkage with occupational medicine/employee health and a memorandum of understanding (MOU) with local public health service/department.
5. Statewide protocol approved by the EMS Board.
6. ALS resuscitation equipment (refer to *The Maryland Medical Protocols for Emergency Medical Services*) must be available on-site during vaccinations.
7. Must use the comprehensive training curriculum developed by MIEMSS Infection Control Committee.
8. Physician does not have to be physically present for the administration of vaccinations or tests by the trained paramedic (Vaccination and Testing Officer (VTO)).
9. Program instruction must be directed by and have participation by the jurisdictional medical director to select paramedics who will become the VTOs.
10. This is not for post-exposure prophylaxis (patient must be seen by occupational medicine/physician for consent and treatment).
11. Mechanism for meeting FDA storage and refrigeration standards for vaccines and testing with the use of the Maryland Inventory Control Sheet.
12. Mechanism for follow-up
   a) For additional vaccinations for completion of series
   b) For potential complications of vaccinations or symptoms noted on adverse event form (meeting federal reporting requirements)
   c) Patient contact phone number for complications (e.g., bad vaccine “lot”)
13. Must have a standardized informed consent form and standardized vaccine pre-screening questionnaire form.

14. Vaccinations allowable are:
   a) Influenza - Commercial and 9-1-1
   b) Hepatitis B
   c) COVID-19 - Commercial and 9-1-1

15. Testing
   PPD Screening (Intradermal) - 9-1-1 only

16. Recommend 30-minute observation period (to be determined by the jurisdictional medical director) post-immunization administration with ALS personnel and equipment available.
HEPATITIS B VACCINATION

1. INDICATIONS
   Pre-exposure: preventive

2. CONTRAINDICATIONS
   History of anaphylactic reaction to baker’s yeast

3. ADVERSE EFFECTS
   Not clinically significant

4. PRECAUTIONS
   (1) Recipients must read and sign consent form.
   (2) CDC recommends antibody testing 1–2 months after the third dose to determine immunity.

5. DOSE
   (three total, using a 3 mL syringe with 1” 25 gauge needle)
   Initial 1 mL IM (deltoid)
   2nd dose 4 weeks after initial; 1 mL IM (deltoid)
   3rd dose 5–6 months after 2nd dose; 1 mL IM (deltoid)

INFLUENZA VACCINATION

1. INDICATIONS
   (1) Persons who attend to patients at high risk for complications (e.g., the elderly)
   (2) Persons with chronic medical conditions
   (3) Pregnant women who will be in the second or third trimester of pregnancy during influenza season
   (4) Clinicians of essential community services

2. CONTRAINDICATIONS
   History of anaphylactic hypersensitivity to eggs

3. ADVERSE EFFECTS
   (1) More common: soreness at the injection site that lasts up to 2 days
   (2) Less common: fever, malaise, myalgia beginning 6–12 hours after vaccination and persisting for 1 to 2 days.

4. PRECAUTIONS
   (1) Vaccine should be delayed in the presence of acute febrile illness; administer after symptoms have abated.
   (2) It takes two weeks to develop adequate antibodies against the vaccine virus strain.
   (3) Optimal time for organized vaccination campaigns is usually the period from October through mid-November.
   (4) Because influenza vaccine contains only noninfectious viruses, it cannot cause influenza.
   (5) Recipients must read and sign consent or refusal form.
5. **DOSE**
   (using a 3 mL syringe with 1" 25 gauge needle)
   0.5–1 mL IM (deltoid)

**PURIFIED PROTEIN DERIVATIVE (PPD) TEST**

1. **INDICATIONS**
   Yearly administration for health care clinicians

2. **CONTRAINDICATIONS**
   (1) Previous positive reaction to PPD
   (2) History of TB

3. **ADVERSE EFFECTS**
   Not clinically significant

4. **PRECAUTIONS**
   Recipients must read and sign consent form.

5. **PROCEDURE**
   (1) Injection is given intradermally and should be read 48–72 hours post injection.
   (2) Feel the induration with your fingertips.
   (3) Measure with approved device in millimeters (mm).
      (a) Less than 5 mm is negative.
      (b) Equal to or greater than 5 mm requires clinical correlation and evaluation by jurisdictional medical director or other appropriate physician.

**NOTE**
Do not use erythema as margins; measure only the induration.
1. **PURPOSE**
   Mechanical CPR (mCPR) devices perform chest compressions at a consistent and reliable rate and depth, never fatigue, and are not susceptible to other human factors that degrade resuscitation quality. Additionally, the use of an mCPR device while transporting an in-progress resuscitation allows for effective CPR and increases safety by allowing clinicians to be restrained during transport.

2. **PRESENTATION**
   Patients in cardiac arrest who have an established resuscitation in progress

3. **INDICATIONS**
   a) Active cardiac arrest resuscitation
   b) Applied in a standby mode for transport to any patient
      (1) who achieves ROSC, OR
      (2) who clinicians believe will progress to cardiac arrest

4. **CONTRAINDICATIONS**
   Patients who have not yet reached their 13th birthday

5. **PROCEDURE:**
   a) Application of an mCPR device may not begin until after two 2-minute cycles of manual chest compressions.
   b) Any mCPR device must be applied in a manner that limits any break in compressions to less than 10 seconds.
   c) The 10-second breaks for device application must only occur around a normal 2-minute compression interval and simultaneously while performing rhythm interpretation and defibrillation.
   d) Apply the mCPR device according to manufacturer instructions, keeping in mind that minimizing breaks in compressions to less than 10 seconds may require that an mCPR device be applied over two or more 2-minute cycles of chest compressions.
   e) Once applied, devices must be used in accordance with manufacturer recommendations, but the goal should be to limit breaks in compressions as little as possible. This goal can be accomplished by:
      (1) Only pausing the mCPR device for rhythm interpretation
      (2) Pausing only long enough to identify the rhythm, and then starting again
      (3) Delivering defibrillations while chest compressions are in progress
   f) An mCPR device (if available) should be applied in a standby mode for transport to any patient who achieves ROSC or patients who clinicians believe will progress to cardiac arrest.

6. **PRECAUTIONS**
   Application of an mCPR device shall not cause delays in assessing for a shockable rhythm or the initiation of manual CPR.
7. **INITIAL TRAINING**
   The jurisdictional medical director must certify that personnel have received a locally-approved training program prior to implementation.

8. **ONGOING DEMONSTRATION OF PROFICIENCY**
   The jurisdictional medical director must reaffirm that EMSOP clinicians have received annual training with the mCPR device.
1. **INDICATIONS**

Patients who may qualify for a home visit by the MIH team include:

a) Patients who have called 9-1-1 for any medically-related reason five times in any six-month interval (patient’s consent required) or

b) Patients who are referred to the MIH team by other allied health professionals or EMS clinicians (patient’s consent required)

2. **PRECAUTIONS**

a) Upon initiation of the home visit, if any patient exhibits any signs or symptoms that would require transport to an emergency department, the MIH team will contact the county dispatch center and request an emergent response.

(1) MIH paramedic will perform all assessments and care based upon current *Maryland Medical Protocols for Emergency Medical Services* until the appropriate 9-1-1 EMS resource’s arrival. Patient care may then be transferred to that EMS unit.

(2) MIH paramedic will not perform blood draws (exception: blood glucose and hemoglobin A1c), medication administration, or ALS interventions unless an immediate life-threatening condition has been identified and the local 9-1-1 Center has been notified of need for an emergent response.

b) MIH home visits may be conducted by a paramedic (MIH service-credentialed) or a paramedic/RN team or a paramedic/NP team.

3. **CONTRAINDICATIONS**

Patients who will not qualify for this program include:

a) Patients already receiving care from a patient-centered medical home (PCMH) or who have already established individual home health care or use a visiting nurse agency (unless the MIH program is partnered with a home health agency)

b) Patients who refuse participation by revoking written consent, verbal refusal of care at time of visit, or are already enrolled in another home health care program

c) Patients who have not reached their 18th birthday

4. **PROCEDURE**

After an individual has consented to be included in this program, the MIH paramedic will:

a) Provide a recognized uniformed presence for individual reassurance and familiarity

b) Assess the individual's home environment

   (1) Assess for signs of neglect or abuse

   (2) Assess for safety issues (e.g., slip/fall risk, smoke detector, fire hazards, exposed electrical wiring)

   c) Obtain vital signs and patient’s weight

d) Obtain and review the patient’s past medical, family, and social history

e) Conduct a basic physical assessment

f) Review medication list (with NP/RN team or telemedicine support)

   g) Review behavioral health needs (with NP/RN team or telemedicine support)

h) Assess the patient’s overall health status using an approved survey tool (e.g., EQ-5D)

   i) Make appropriate health professional contacts and provide medication education (with NP/RN team or telemedicine support) and referrals
5. MEDICAL CONSULTATION
   a) Online medical consultation may be obtained from a jurisdictional/commercial EMS medical director or through a designated base station.
   b) Paramedics cannot accept orders from primary care physicians on the phone or on-scene unless individual has an immediate life-threatening condition and the physician is going to the hospital with the patient on EMS unit.
   c) Nurse practitioners (NP) and registered nurses (RN) may not direct an EMS clinician to perform any skill or medical intervention that is not within his or her scope of practice nor provide “Medical Consultation” as referenced in The Maryland Medical Protocols for Emergency Medical Services.

6. DOCUMENTATION
   a) MIH paramedic will complete an eMEDS® report using the MIH module.
   b) Other MIH team members (NP/RN) will document in an electronic health record approved by their home agency.
   c) MIH teams will establish policies and procedures for sharing of protected health information across allied health, social services, and community organizations, with resources available for patients.
   d) In the event that an immediate life-threatening condition is identified and the MIH paramedic initiated urgent/emergent EMS care:
      (1) MIH paramedic will complete an eMEDS® report in the standard (emergency response) eMEDS® system or Commercial EMSOP equivalent.
Optional Supplemental Protocol – MOBILE INTEGRATED HEALTH: COVID-19 Monoclonal Antibody Administration

1. INDICATIONS
   a) Treatment of COVID-19 Infection: Patients 12 years of age or older and weighing at least 40 kg who meet all of the following criteria:
      (1) COVID-19 positive
      (2) Mild-moderate symptoms, not requiring hospitalization
      (3) Within 10 days of symptom onset
      (4) No supplemental oxygen requirement (or no increase in chronic home oxygen therapy)
      (5) At high risk for progression to severe disease or death as evidenced by one of the following:
         (a) Age 65 years or older
         (b) Obesity or overweight (BMI greater than or equal to 25)
         (c) Diabetes
         (d) Cardiovascular disease, including hypertension
         (e) Chronic lung disease (e.g., COPD, asthma)
         (f) Chronic kidney disease
         (g) Immunocompromising condition or immunosuppressant medication
         (h) Sickle cell disease
         (i) Pregnancy
   b) COVID-19 Post-Exposure Prophylaxis: Patients 12 years of age or older and weighing at least 40 kg who:
      (1) Are not fully vaccinated or who are immunocompromised, and
      (2) Have been exposed to a COVID-19 positive person via close contact (within 6 feet for 15 minutes or more, direct physical contact)
   c) Obtain vital signs prior to administering monoclonal antibodies
   d) Intravenous infusion (preferred)
      (1) Establish intravenous access.
      (2) Prepare one of the following infusions by diluting monoclonal antibodies in appropriate volume of normal saline, per manufacturer package insert or FDA EUA instructions:
         (a) Bamlanivimab 700 mg / etesevimab 1,400 mg
         (b) Casirivimab 600 mg / indevimab 600 mg
         (c) Sotrovimab 500 mg
      (3) Infuse monoclonal antibodies with an infusion pump, appropriate filter, and infusion set using the rate recommended by package insert. Do not administer as IV push or bolus.
      (4) Monitor patient during the infusion for any sign of allergic reaction or hypersensitivity.
         (a) If any adverse or allergic reaction occurs, stop the infusion immediately and treat per Allergic Reaction or Anaphylaxis protocols.
      (5) After infusion is complete, flush the line to ensure delivery of all of the medication.
      (6) Monitor the patient for adverse effects and obtain vital signs every 15 minutes, for 60 minutes after completion of the infusion.
e) Subcutaneous administration (casirivimab/indevimab only, if IV unavailable)
   (1) Prepare two syringes of casirivimab and two syringes of indevimab, per package insert
   (2) Administer four subcutaneous injections, each at a different injection site, into the proximal thighs and back of the proximal arms, per package insert.
   (3) Do not administer into skin that is tender, bruised, scarred, or damaged.
   (4) Monitor the patient for adverse effects and obtain vital signs every 15 minutes, for 60 minutes after medication administration.
   (5) If any adverse or allergic reaction occurs, treat per Allergic Reaction or Anaphylaxis protocols.

2. Clinical Pearls
   a) Ensure storage, handling, and preparation per FDA and manufacturer recommendations:
      (1) Bamlanivimab/etesevimab: https://www.fda.gov/media/145802/download
      (2) Casirivimab/imdevimab: https://www.fda.gov/media/145611/download
      (3) Sotrovimab: https://www.fda.gov/media/149534/download
   b) Report any serious adverse events (www.fda.gov/medwatch/report.htm).
1. **PURPOSE**
   Naloxone is a prescription medication indicated for the reversal of respiratory depression or unresponsiveness due to opioid overdose. Increasing the accessibility and availability of naloxone to family members, close friends, or the public, specifically those at risk for an opioid overdose, may reduce the chance of a prolonged hypoxic event or eventual cardiac arrest.


2. **INDICATIONS**
   a) Following an administration of naloxone prior to arrival of EMS or as described by The Maryland Medical Protocols for Emergency Medical Services or
   b) Following evaluation by a crisis intervention team (e.g., Safe Station for opioid treatment referral) that has identified an opioid dependent individual when immediate placement cannot occur and the individual is released.

3. **CONTRAINDICATIONS**
   “Leave Behind” naloxone shall not be dispensed to anyone who has not yet reached their 18th birthday.

4. **PROCEDURE**
   a) Following completion of all general patient care, which may include a patient-initiated refusal of care, naloxone hydrochloride(s) and necessary paraphernalia that has been approved by the EMS Operational Program in accordance with Maryland Department of Health Guidelines may be issued.
   b) Document the distribution of naloxone in the patient care report as required by the EMS Operational Program.
Optional Supplemental Protocol – PELVIC STABILIZATION BINDER DEVICE

All levels of EMS clinicians, if appropriately trained in the device

1. INDICATIONS
   All of the following blunt trauma patients with physical findings indicative of pelvic fracture should have a Pelvic Stabilization Binder Device applied.
   a) Evidence of pelvic instability on examination of the pelvis
   b) Patients complaining of pelvic pain on examination of the pelvis
   c) Pain on iliac compression
   d) Pain on compression of the pubic symphysis
   e) Blood at the urethral meatus
   f) Vaginal bleeding
   g) Perineal or scrotal hematoma
   h) All blunt trauma patients with an unreliable physical exam and significant mechanism of injury may be considered for application of a Pelvic Stabilization Binder Device.

   PREGNANCY IS NOT A CONTRAINDICATION TO THE APPLICATION OF THE PELVIC STABILIZATION BINDER DEVICE WHEN INDICATED.

2. CONTRAINDICATIONS
   a) Patient for whom the smallest available pelvic stabilization binder is too wide and places pressure on abdomen or chest
   b) Children under 144 cm (4’8”) will generally NOT fit small-size adult pelvic stabilizing devices.

3. PROCEDURE
   a) Assess for pelvic instability.
      In order to not increase bleeding, only one exam should be performed to evaluate for pelvic fracture. Multiple exams will disrupt clot formation.
   b) Identify the greater trochanter of each femur.
      The greater trochanter is the bony prominence of the lateral upper thigh.
   c) Identify the anterior superior iliac spine.
   d) Check size with estimating stabilization device and center at the greater trochanter.
      Ensure the top of the binder does not go above the anterior superior iliac spine.
   e) The patient should be placed in a supine position prior to application of the pelvic stabilization binder device.
   f) Place pelvic binder around the patient, centered at the level of the greater trochanter.
   g) If a backboard is used, place the binder on the backboard prior to placing the patient on the backboard.
   h) Ensure patient has been undressed and adequate exposure is provided.
   i) Tighten the binder as directed by the manufacturer’s instructions for the specific stabilization binder.
   j) Once pelvic stabilization binder device is applied, do not remove until directed to do so by a physician.
4. PRECAUTIONS
   a) Incorrectly placing the pelvic stabilization binder device at the level of the iliac wing could cause harm by widening the pelvic fracture. Assess after application of the pelvic stabilization binder device.
   b) Continue with patient care.
   c) EMS clinicians should also assess distal pulses before and after the application of the pelvic stabilization binder device.
   d) For EMS units with long transport times and with patients requiring large volumes of fluid resuscitation, the patient will need to be periodically monitored to make sure that the device is not becoming too tight due to expansion of the pelvic area from accumulation of fluids that have third spaced to the pelvic area.
   e) If clinicians feel the device is becoming too tight, it should be slowly loosened and then reapplied.
Optional Supplemental Protocol –
RSI – ADULT: RSI OPTIONAL SUPPLEMENTAL PROGRAM

1. INDICATIONS
   a) Inability to tolerate laryngoscopy, and:
      (1) GCS less than or equal to 8 with respiratory rate less than or equal to 8 or greater than or equal to 35 or
      (2) GCS less than or equal to 8 with oxygen saturation less than or equal to 90% on non-rebreather face mask
   b) On-line medical direction for RSI may be requested in the following situations:
      (1) GCS less than or equal to 8 with clenched jaw or inability to adequately suction airway, and without above respiratory parameters
      (2) Respiratory extremis with contraindication to noninvasive ventilation (CPAP)
      (3) Burns: inhalational burn with objective signs of developing airway compromise
      (4) Critically ill or severely injured patient with imminent airway compromise

2. CONTRAINDICATIONS
   Patients who have not yet reached their 15th birthday
   Caution – interior STEMI patients who may be preload-dependent

3. PREPARATION
   a) Pre-oxygenate with nasal cannula oxygen 15 lpm and bag valve mask attached to high-flow oxygen
   b) Monitor vital signs q5min, continuous pulse oximetry, ETCO₂, and cardiac rhythm
   c) Ensure functioning IV/IO and fluid therapy as per protocol.
   d) Evaluate for difficult airway.
   e) Perform focused RSI neurologic exam.
   f) Prepare equipment
      (1) Intubation equipment
      (2) Bag-Valve-Mask (BVM)
      (3) ET tube introducer (bougie)
      (4) Suction
      (5) RSI medications
      (6) Alternative airway device
      (7) Cricothyroidotomy equipment
      (8) Video laryngoscopy equipment

4. RSI PROCEDURE
   a) Sedation
      Adequate sedation must be provided to prevent awareness during paralysis from neuromuscular blockade.
      **Etomidate** may be used for sedation in patients who are aware of their surroundings.
      Initial dose: Administer 0.3 mg/kg IVP/IO over 30–60 seconds. If the patient is hypotensive or the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP/IO over 30–60 seconds.
      Repeat dose: May repeat 0.15 mg/kg IVP/IO in 2–3 minutes if inadequate sedation.
      OR
**Ketamine** is preferred for patients who are aware of their surroundings or have hypotension or possible hypovolemia.

Initial dose: Administer 2 mg/kg IVP/IO over 60 seconds.
Repeat dose: May repeat dose of 2 mg/kg IVP/IO over 60 seconds in 2-3 minutes if inadequate sedation.

**OR**

**Midazolam** can be considered for patients with isolated head injury and elevated blood pressure, especially with possible seizure activity. Midazolam should not be used for patients with hypotension, and should be avoided with possible hypovolemia.

Dose: Administer 0.1 mg/kg IVP/IO over 1–2 minutes.
Maximum single dose is 5 mg.

Only one sedative agent should be administered prior to succinylcholine unless otherwise directed by medical consultation.

b) In-line cervical spine stabilization shall be maintained by second caregiver for trauma patients.

c) Administer paralytic medication.

**Succinylcholine**: Administer 1.5 mg/kg rapid IVP/IO. Maximum single dose is 200 mg. If inadequate paralysis after 2-3 minutes, verify IV/IO patency. Repeat succinylcholine 1 mg/kg IVP/IO. Maximum single dose is 200 mg.

Contraindications for succinylcholine:
1. Burns greater than 24 hours old
2. Spinal cord injury greater than 24 hours old
3. Known neuromuscular disease (Guillain-Barre syndrome, myasthenia gravis, amyotrophic lateral sclerosis, muscular dystrophy)
4. Chronic renal failure on hemodialysis/Presence of hemodialysis access
5. Malignant hyperthermia

**OR**

**Vecuronium**: 0.1 mg/kg IVP/IO; if inadequate paralysis after 2-3 minutes, verify IV/IO patency. Repeat vecuronium 0.05 mg/kg IVP/IO. Vecuronium is the preferred paralytic for patients with a history of malignant hyperthermia or contraindications for succinylcholine.

WHEN VECURONIUM IS USED, MAINTENANCE OF SEDATION MUST BE ASSURED. THE PATIENT MAY NOT SHOW TRADITIONAL SIGNS OF VENTILATORY BUCKING.

d) Intubate trachea and verify ET placement.

### 5. SUCCESSFUL ENDOTRACHEAL TUBE PLACEMENT

(1) Secure ET.

(2) Ventilate the patient at a rate that maintains an ETCO$_2$ of 35-40 mmHg. For the head-injured patient with signs/symptoms of herniation, target ETCO$_2$ of 30 mmHg.

(3) If significant resistance to ventilation occurs as succinylcholine wears off (4–5 minutes), refer to *Ventilatory Difficulty Secondary to Bucking* protocol.
6. **UNSUCCESSFUL ENDOTRACHEAL TUBE PLACEMENT**
   a) Place an airway adjunct (OPA or NPA) and apply high-flow oxygen by nasal cannula (no desat).
   b) Resume BVM ventilation for at least 30 seconds and attempt to correct hypoxia, if present. If unable to ventilate, see “If Unable to Ventilate” below.
   c) Reevaluate airway strategies and consider additional oral ET intubation attempt.
   d) If unsuccessful, resume BVM ventilation.
   e) Insert an approved alternative airway device (refer to Extralglottic Airway protocol).
   f) Attach capnography and ventilate to ETCO$_2$ level of 35-40 mmHg. For head-injured patients with signs/symptoms of herniation, target ETCO$_2$ of 30 mmHg.
   g) If significant resistance to ventilation occurs as succinylcholine wears off (4–5 minutes), or if patient exhibits difficulty in tolerating an approved alternative airway device as succinylcholine wears off, refer to Ventilatory Difficulty Secondary to Bucking protocol.

7. **IF UNABLE TO VENTILATE**
   Insert an approved alternative airway device (refer to Extralglottic Airway protocol).

8. If still unable to ventilate using an approved alternative airway device, remove it and perform cricothyroidotomy (refer to Cricothyroidotomy protocol).
1. INDICATIONS
Patients successfully intubated with an endotracheal tube, an approved alternative airway device, or cricothyroidotomy, for whom the ability to provide manual or mechanical ventilation is impaired secondary to bucking or combativeness.

2. CONTRAINDICATIONS
Unsecured airway.

3. PROCEDURE
a) Consider additional sedation
   - **Etomidate**
     Dose: Administer 0.3 mg/kg IVP/IO over 30–60 seconds. If the patient is hypotensive or the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP over 30–60 seconds. May repeat 0.15 mg/kg IVP/IO every 15 minutes to a total of 3 doses.
   
   OR
   - **Ketamine** may be preferred for patients who have hypotension or possible hypovolemia, or if ventilatory difficulty is thought to be the result of pain response.
     Dose: Administer 2 mg/kg IVP/IO over 60 seconds. May repeat 2 additional doses of 1 mg/kg for IVP/IO every 10–15 minutes to a total of 3 doses as needed.

   Additional doses require medical consultation.

   OR
   - **Midazolam** can be considered for patients with isolated head injury and elevated blood pressure, especially with possible seizure activity. Midazolam should not be used for patients with hypotension, and should be avoided with possible hypovolemia.
     Dose: Administer 0.1 mg/kg IVP/IO over 1–2 minutes, titrated to abate bucking and relax ventilation while maintaining systolic BP greater than 90 mmHg (110 mmHg if injuries include a suspected head injury). Maximum single dose is 5 mg.

   Additional doses require medical consultation.

b) If ventilatory difficulty is thought to be the result of pain response,
   - **Ketamine may be used as above.**
   
   OR
   Opioid may be used per Pain Management protocol in addition to, or instead of, midazolam, ketamine, or etomidate. Titrate to abate bucking and relax ventilation while maintaining systolic BP greater than 90 mmHg.

SEDATION MUST BE PROVIDED PRIOR TO VECURONIUM FOR A PATIENT WHO IS EITHER RESPONSIVE TO STIMULUS, OR WHO MAY BECOME RESPONSIVE TO STIMULUS DURING NEUROMUSCULAR BLOCKADE. USE OF VECURONIUM REQUIRES FUNCTIONING ETCO₂ MONITORING. VECURONIUM MAY ONLY BE USED IF CONTINUOUS, BREATHE TO BREATHE ETCO₂ MONITORING CAN BE PROVIDED.
c) If significant resistance to ventilation continues despite adequate sedation and analgesia, the paramedic may administer:
   (1) Vecuronium 0.05 mg/kg IVP/IO. Maximum single dose is 10 mg.
   (2) Dose may be repeated in 2-3 minutes, if necessary.
   (3) **Maintenance of amnesia**
       Follow above dosing of either *etomidate* or *ketamine* with required repeat dosing every 10–15 minutes.

d) Continue to monitor oxygen saturation and ventilate to desired ETCO₂.
e) Obtain on-line medical direction if further problems present.
Optional Supplemental Protocol –
RSI – ADULT: PROTOCOL FOR CRICOTHYROIDOTOMY (SURGICAL AND NEEDLE) 15.22

1. INDICATIONS
   a) Inability to ventilate despite having tried BVM with oropharyngeal/nasopharyngeal airway, ET placement, and an alternative airway device (if not contraindicated)
   b) Inability to place ET in the setting of life-threatening upper airway hemorrhage
   c) Foreign body completely obstructing upper airway that cannot be removed via BLS maneuvers or Magill forceps with direct visualization

2. CONTRAINDICATIONS
   Patients under the age of 8 should not receive surgical cricothyroidotomy. They may receive only needle cricothyroidotomy.

3. PREPARATION
   a) Prepare suction and cricothyroidotomy kit.
   b) Begin at sternal notch and locate cricoid cartilage.
   c) Palpate cricothyroid membrane anteriorly between cricoid cartilage and thyroid cartilage.
   d) Prepare skin with betadine or alcohol swabs.

4. SURGICAL CRICOTHYROIDOTOMY
   a) Clinicians must use a designated technique and procedure for establishing the airway through the cricothyroid membrane that has been approved by the program medical director.
   b) Insert a 6.0 mm cuffed ET tube, using the natural curve of tube.
   c) Insert ET tube to just beyond cuff.
   d) Inflate cuff and ventilate patient.
   e) Monitor oxygen saturation and ETCO₂ level.
   f) Secure ET tube. (Do not cut or trim ET tube.)
   g) If significant resistance to ventilation develops, or if patient develops difficulty in tolerating successful cricothyroidotomy, refer to Ventilatory Difficulty Secondary to Bucking or Combativeness protocol.

5. NEEDLE CRICOTHYROIDOTOMY
   a) Insert 12- or 14-gauge over-the-needle catheter through the cricothyroid membrane at a 45-degree angle toward the feet. Aspiration of air with a syringe indicates tracheal entry.
   b) Hold needle in place and advance catheter, then remove needle.
   c) Attach catheter hub to intermittent jet oxygen insufflator valve.
   d) Manually secure catheter at hub at all times to prevent kinking or displacement.
   e) Monitor oxygen saturation.
   f) If significant resistance to ventilation develops, or if patient develops difficulty in tolerating cricothyroidotomy, refer to Ventilatory Difficulty Secondary to Bucking or Combativeness protocol.
1. INDICATIONS
   a) Inability to tolerate laryngoscopy and have the following:
      (1) GCS less than or equal to 8, indicated by a patient that will not: open eyes, cry, say
          words, or show purposeful movement in response to painful stimulus.
      AND
      (2) Respiratory insufficiency, demonstrated by oxygen saturation less than or equal to
          90% on non-rebreather face mask, respiratory rate less than or equal to 8, or respira-
          tory rate greater than or equal to 45 (age less than 1 yr), greater than or equal to
          40 (age 1–5 yrs), greater than or equal to 35 (age 6–9 yrs) with signs of air hunger
          and accessory muscle use.

   PATIENTS WITH AN IDENTIFIED DIFFICULT AIRWAY WHO CAN BE BAGGED TO AN OXYGEN
   SATURATION GREATER THAN 90% REQUIRE ON-LINE MEDICAL DIRECTION FOR RSI,
   PREFERABLY FROM A PEDIATRIC BASE STATION.

   b) On-line medical direction for RSI may be requested (preferably from a Pediatric
      Base Station), in the following situations:
      (1) GCS less than or equal to 8 with clenched jaw, inability to adequately suction air-
          way, and without above respiratory parameters
      (2) Respiratory extremis with contraindications to nasotracheal intubation (respiratory
          rate greater than or equal to 35 with air hunger, use of accessory muscles, and oxy-
          gen saturation less than or equal to 90% on non-rebreather face mask)
      (3) Identified difficult airway patient with a GCS less than or equal to 8 and signs of
          respiratory insufficiency who cannot tolerate laryngoscopy but is able to be bagged
          to an oxygen saturation greater than 90%

2. CONTRAINDICATIONS
   a) Conditions that may cause hyperkalemia:
      (1) Burns greater than 24 hours old
      (2) Spinal cord injury greater than 24 hours old
      (3) Known neuromuscular disease (Guillain-Barré Syndrome, myasthenia gravis, amyotro-
          phic lateral sclerosis, muscular dystrophy)
      (4) Chronic renal failure on hemodialysis/presence of hemodialysis access

   b) History of malignant hyperthermia

3. PREPARATION
   a) Pre-oxygenate with 90–100% oxygen.
   b) Monitor oxygen saturation with pulse oximetry and EKG.
   c) Ensure functioning IV and fluid therapy as per protocol.
   d) Evaluate for difficult airway.
   e) Perform focused RSI neurologic exam.
Optional Supplemental Protocol – RSI – PEDIATRIC: RSI OPTIONAL SUPPLEMENTAL PROGRAM (continued)

f) Prepare equipment
   (1) Intubation kit: Recommended to carry both cuffed and uncuffed ET tubes for patients less than 8 years of age or 25 kg.
   (2) Bag-Valve-Mask (BVM) with manometer. (Manometer may be part of the BVM or separate.)
   (3) Suction
   (4) RSI kit
     (a) Prepare medications
     (b) Alternative airway device, cricothyroidotomy equipment
   (5) Capnograph

4. RSI PROCEDURE
a) Adequate sedation must be provided to prevent awareness during paralysis from neuromuscular blockade.

*Etomidate*, if available, will be the preferred agent for patients who are aware of their surroundings and do not have hypotension or possible hypovolemia.

Dose: Administer 0.3 mg/kg IVP/IO over 30–60 seconds. If the patient is hypotensive or the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP/IO over 30–60 seconds. May repeat 0.15 mg/kg IVP/IO in 2–3 minutes if inadequate sedation.

*Ketamine* may be used if etomidate is unavailable, and may be preferred for patients who have hypotension or possible hypovolemia.

Dose: Administer 2 mg/kg IVP/IO over 60 seconds.

*Midazolam* should be considered for patients with isolated head injury and elevated blood pressure, especially with possible seizure activity. Midazolam should not be used for patients with hypotension, and should be avoided with possible hypovolemia.

Dose: Administer 0.05 mg/kg IVP/IO over 1–2 minutes. Maximum single dose is 5 mg. **Hold for** BP less than 60 in neonates (patients less than 28 days old), less than 70 in infants (patients less than 1 year of age), less than [70 + (2 x years) = systolic BP] for patients greater than 1 year of age.

b) For patients with head injury or suspected increased intracranial pressure, administer *lidocaine* 1 mg/kg IVP/IO over 1–2 minutes.

c) If patient is less than 8 years of (or if age unknown and using ET tube smaller than 6.0), pretreat patient with atropine 0.02 mg/kg IVP/IO.

d) In-line cervical spine stabilization by second caregiver (in trauma setting)

e) Apply cricoid pressure (by third caregiver).

f) *Succinylcholine*: Administer 1.5 mg/kg rapid IVP/IO.

g) Intubate trachea and verify ET placement.

h) If inadequate relaxation after 2–3 minutes, repeat succinylcholine 1.0 mg/kg IVP/IO.
5. **SUCCESSFUL ENDOTRACHEAL TUBE PLACEMENT**
   a) Release cricoid pressure and secure ET.
   b) Ventililate to ETCO$_2$ of 30–32 mmHg.
   c) If significant resistance to ventilation occurs as succinylcholine wears off (4–5 minutes), refer to *Ventilatory Difficulty Secondary to Bucking* protocol.

6. **UNSUCCESSFUL ENDOTRACHEAL TUBE PLACEMENT**
   a) Maintain cricoid pressure and resume BVM ventilation for 30 seconds.
   b) If unable to ventilate, see “If Unable to Ventilate” below.
   c) Reattempt oral ET intubation.
   d) If unsuccessful, resume BVM ventilation for 30 seconds.
   e) Insert an extraglottic airway (see *Airway Management: Extraglottic Airway* protocol).

7. **IF UNABLE TO VENTILATE**
   If unable to ventilate, verify appropriate oropharyngeal airway placement and reposition BVM for optimal mask seal. If still unable to ventilate, refer to *Needle Cricothyroidotomy* protocol.
1. **INDICATIONS**
   Patients successfully intubated with an endotracheal tube, or needle cricothyroidotomy, for whom the ability to provide manual or mechanical ventilation is impaired secondary to bucking or combativeness.

2. **CONTRAINDICATIONS**
   Unsecured airway

3. **PROCEDURE**
   a) **Etomidate**, if available, will be the preferred agent for patients who are aware of their surroundings and do not have hypotension or possible hypovolemia.
      
      Dose: Administer 0.3 mg/kg IVP/IO over 30–60 seconds. If the patient is hypotensive or the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP/IO over 30–60 seconds. May repeat 0.15 mg/kg IVP/IO in 2–3 minutes if inadequate sedation.

   OR
   Ketamine may be used if etomidate is unavailable, and may be preferred for patients who have hypotension or possible hypovolemia, or if ventilatory difficulty is thought to be the result of pain response.
   
   Dose: **Ketamine**: 2 mg/kg IVP/IO over 60 seconds. May repeat 1 mg/kg for IVP/IO every 10–15 minutes to a total of three doses as necessary.
   
   OR
   Midazolam should be considered for patients with isolated head injury and elevated blood pressure, especially with possible seizure activity. Midazolam should not be used for patients with hypotension, and should be avoided with possible hypovolemia.
   
   Dose: Administer 0.05 mg/kg IVP/IO over 1–2 minutes, titrated to abate bucking and relax ventilation while maintaining systolic BP: greater than 60 in neonates, 70 in infants, [70 + (2 x years) = systolic BP] for patients greater than 1 year of age. Maximum single dose is 5 mg.

   b) If ventilatory difficulty is thought to be the result of pain response,
   Ketamine: Dose: 2 mg/kg IVP over 60 seconds. May repeat 1 mg/kg IVP/IO every 10–15 minutes as necessary to a total of three doses as necessary.

   OR
   Opioid may be used per Pain Management protocol in addition to, or instead of, midazolam, ketamine, or etomidate. Titrate to abate bucking and relax ventilation while maintaining systolic BP greater than 60 in neonates, 70 in infants, [70 + (2 x years) = systolic BP] for patients greater than 1 year of age.
Optional Supplemental Protocol –
RSI – PEDIATRIC: VENTILATORY DIFFICULTY SECONDARY TO BUCKING OR
COMBATIVENESS IN INTUBATED PATIENTS (continued)

c) If significant resistance to ventilation continues, the paramedic may administer:

   (1) Vecuronium 0.05 mg/kg IVP/IO (may not be used for patients with needle cricothyroidotomy because of inability to monitor breath to breath ETCO₂). Maximum single dose is 10 mg.

**PRE-SEDATION MUST BE PROVIDED WHEN VECURONIUM IS ADMINISTERED TO A PATIENT WHO IS EITHER RESPONSIVE TO STIMULUS OR MAY BECOME RESPONSIVE TO STIMULUS DURING NEUROMUSCULAR BLOCKADE. VECURONIUM MAY ONLY BE USED IF CONTINUOUS, BREATH TO BREATH ETCO₂ MONITORING CAN BE PROVIDED.**

(2) Dose may be repeated in 2-3 minutes if necessary.
(3) **Maintenance of Amnesia**
   Follow above dosing of either *etomidate* or *ketamine* with required repeat dosing every 10–15 minutes.

d) Continue to monitor oxygen saturation and ventilate to desired ETCO₂.
e) **Obtain on-line medical direction (preferably from a Pediatric Base Station),** if further problems present.
Optional Supplemental Protocol –
RSI – PEDIATRIC: PROTOCOL FOR CRICOTHYROIDOTOMY (SURGICAL AND NEEDLE) 15.25

Surgical (for 8 years old or greater) and Needle

1. INDICATIONS
   a) Inability to ventilate despite having tried BVM with oropharyngeal/nasopharyngeal airway, ET placement, and alternative airway device (if not contraindicated)
   b) Inability to place ET in the setting of life-threatening upper airway hemorrhage
   c) Completely obstructing upper airway foreign body that cannot be removed via BLS maneuvers or Magill forceps with direct visualization

2. PREPARATION
   a) Prepare suction and cricothyroidotomy kit.
   b) Begin at sternal notch and locate cricoid cartilage.
   c) Palpate cricothyroid membrane anteriorly between cricoid cartilage and thyroid cartilage.
   d) Prepare skin with betadine or alcohol swabs.

3. SURGICAL CRICOTHYROIDOTOMY FOR 8 YEARS OLD OR GREATER
   a) Stabilize thyroid cartilage and make vertical incision (1–1 1/2 inches) over cricothyroid membrane. Alternatively, a needle puncture dilator device may be utilized.
   b) Palpate cricothyroid membrane with gloved finger and carefully make transverse incision through membrane. Insert scalpel handle and rotate 90 degrees.
   c) Insert a 5 to 6.0 mm cuffed ET tube, using the natural curve of tube.
   d) Insert ET tube to just beyond cuff.
   e) Inflate cuff and ventilate patient.
   f) Monitor oxygen saturation and ETCO₂ carbon dioxide level.
   g) Secure ET tube. (Do not cut or trim ET tube.)
   h) If significant resistance to ventilation develops, or if patient develops difficulty in tolerating successful cricothyroidotomy, refer to Ventilatory Difficulty Secondary to Bucking or Combativeness protocol.

4. NEEDLE CRICOTHYROIDOTOMY
   a) Insert 12- or 14-gauge over-the-needle catheter through the cricothyroid membrane at a 45-degree angle toward the feet. Aspiration of air with a syringe indicates tracheal entry.
   b) Hold needle in place and advance catheter, then remove needle.
   c) Attach catheter hub to intermittent jet oxygen insufflator valve.
   d) Manually secure catheter at hub at all times to prevent kinking or displacement.
   e) Monitor oxygen saturation.
   f) If significant resistance to ventilation develops, or if patient develops difficulty in tolerating cricothyroidotomy, refer to Ventilatory Difficulty Secondary to Bucking or Combativeness protocol.

**ONLY NEEDLE CRICOTHYROIDOTOMY SHOULD BE PERFORMED FOR PATIENTS LESS THAN AGE 8 WHO MAY REQUIRE CRICOTHYROIDOTOMY.**
ETOMIDATE (AMIDATE)

1. PHARMACOLOGY
   Hypnotic

2. PHARMACOKINETICS
   A short-acting nonbarbiturate hypnotic agent without analgesic properties

3. INDICATIONS
   Pre-sedation of responsive patients prior to administration of neuro-muscular blocking agents

4. CONTRAINDICATIONS
   Known hypersensitivity to etomidate

5. ADVERSE EFFECTS
   a) Respiratory depression or apnea
   b) Hypotension (infrequent)
   c) Involuntary myoclonus
   d) Adrenal suppression (possible with repeated dosing)

6. PRECAUTIONS
   a) The effects of etomidate can be accentuated by CNS depressants such as opioids and alcohol.
   b) Myoclonic movements are common and should not be confused for fasciculations due to a depolarizing neuromuscular blocking agent or seizure activity.

7. DOSAGE
   a) Adult:
      Administer 0.3 mg/kg IVP/IO over 30–60 seconds.
      If the patient is hypotensive or the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP/IO over 30–60 seconds.
      **Ventilatory Difficulty Secondary to Bucking or Combative in Intubated Patients:**
      Administer 0.3 mg/kg IVP/IO over 30–60 seconds. If the patient is hypotensive or the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP/IO over 30–60 seconds.
      May repeat 0.15 mg/kg IVP/IO every 15 minutes to a total of three doses.
   b) Pediatric:
      Administer 0.3 mg/kg IVP/IO over 30–60 seconds.
      If the clinician suspects hypovolemia, the initial dose will be 0.15 mg/kg IVP/IO over 30–60 seconds. May repeat 0.15 mg/kg IVP/IO after succinylcholine effects resolve and patient is bucking or combative. May repeat 0.15 mg/kg IVP/IO every 15 minutes to a total of three doses.
      Additional doses require medical consultation.
KETAMINE (KETANEST®, KETASET®, KETALAR®)

1. **PHARMACOLOGY**
   Hypnotic Analgesic

2. **PHARMACOKINETICS**
   A rapid-acting hypnotic analgesic agent characterized by normal pharyngeal-laryngeal reflexes, normal or enhanced skeletal muscle tone, and possible cardiovascular and respiratory stimulation.

3. **INDICATIONS**
   a) Pre-sedation of responsive patients prior to administration of neuromuscular blocking agents
   b) Sedation of intubated patients with ventilatory difficulty secondary to bucking or combative

4. **CONTRAINDICATIONS**
   Known hypersensitivity to ketamine

5. **ADVERSE EFFECTS**
   a) Although respiration is frequently stimulated, respiratory depression may occur with rapid IV administration. Laryngospasm has been known to occur.
   b) Although hypotension may occur, blood pressure and heart rate are frequently stimulated.
   c) Involuntary myoclonus that may mimic seizure activity
   d) Possible enhanced secretions
   e) Possible unpleasant dreams and delirium upon emergence from sedation

6. **PRECAUTIONS**
   a) The likelihood of respiratory depression and undesired pressor effects is increased by too rapid IV administration.
   b) Myoclonic movements are possible and should not be confused for fasciculations due to a depolarizing neuromuscular blocking agent, seizure activity, or emergence from sedation.

7. **DOSAGE**
   a) Adult:
      Administer 2 mg/kg IVP/IO over 60 seconds.
      May repeat 2 mg/kg IVP/IO after succinylcholine effects resolve if patient is bucking or combative.
      May repeat 1 mg/kg for IVP/IO every 10–15 minutes to a total of 3 doses, as necessary.
      Additional doses require medical consultation.

   b) Pediatric:
      Administer 2 mg/kg IVP/IO over 60 seconds.
      May repeat 2 mg/kg IVP/IO after succinylcholine effects resolve if patient is bucking or combative.
      May repeat 1 mg/kg for IVP/IO every 10–15 minutes to a total of 3 doses as necessary.
      Additional doses require medical consultation.
MIDAZOLAM (VERSED®)

1. PHARMACOLOGY
   a) Sedative
   b) Hypnotic

2. PHARMACOKINETICS
   A short-acting benzodiazepine with strong hypnotic and amnestic properties

3. INDICATIONS
   a) Pre-sedation of responsive patients prior to administration of neuro-muscular blocking agents
   b) Sedation of intubated patients with ventilatory difficulty secondary to bucking or combativeness

4. CONTRAINDICATIONS
   a) Hypotension
   b) Acute narrow-angle glaucoma
   c) Known hypersensitivity to midazolam

5. ADVERSE EFFECTS
   a) Respiratory depression or apnea
   b) Hypotension
   c) Amnesia

6. PRECAUTIONS
   The effects of midazolam can be accentuated by CNS depressants such as opioids and alcohol

7. DOSAGE
   a) Adult:
      Administer 0.1 mg/kg, SLOW IVP/IO over 1–2 minutes, while maintaining systolic BP greater than 90 mmHg (110 mmHg if injuries include a suspected head injury).
      Maximum single dose is 5 mg.
   b) Pediatric:
      Administer 0.05 mg/kg SLOW IVP/IO over 1–2 minutes, while maintaining systolic BP greater than 60 in neonates, 70 in infants,
      \[70 + (2 \times \text{years}) = \text{systolic BP}\] for patients greater than 1 year of age. Maximum single dose is 5 mg.

   **WARNING:** ADMINISTER UP TO 0.05 MG/KG IV WHEN TREATING ENDOTRACHEAL TUBE BUCKING, STOPPING ONCE BUCKING HAS RESOLVED AND VENTILATION IS RELAXED.
SUCCINYLCHOLINE (ANECTINE®)

1. PHARMACOLOGY
   Neuromuscular blocking agent (depolarizing)

2. PHARMACOKINETICS
   Paralyzes skeletal muscles, including respiratory muscles, and removes gag reflex

3. INDICATIONS
   To achieve paralysis to facilitate endotracheal intubation in patients as per Rapid Sequence Intubation protocol

4. CONTRAINDICATIONS
   a) Conditions that may cause hyperkalemia:
      (1) Burns greater than 24 hours old
      (2) Spinal cord injury greater than 24 hours old
      (3) Known neuromuscular disease (Guillain-Barré syndrome, myasthenia gravis, amyotrophic lateral sclerosis, muscular dystrophy)
      (4) Chronic renal failure on hemodialysis or presence of hemodialysis access
   b) History of malignant hyperthermia
   c) Patients with known hypersensitivity to the drug

5. ADVERSE EFFECTS
   a) Bradycardia
   b) Prolonged paralysis

6. PRECAUTIONS
   Paralysis occurs in 1–2 minutes and generally lasts 4–6 minutes.

7. DOSAGE/ROUTE
   a) Adult:
      Administer 1.5 mg/kg rapid IVP/IO to a maximum single dose of 200 mg.
      If relaxation is inadequate after 2–3 minutes, a repeat dose of 1 mg/kg rapid IVP/IO may be given to a maximum single dose of 200 mg.
   b) Pediatric:
      Administer 1.5 mg/kg rapid IVP/IO to a maximum dose of 200 mg.
      If relaxation is inadequate after 2–3 minutes, a repeat dose of 1 mg/kg rapid IVP/IO may be given to a maximum dose of 200 mg.
VECURONIUM (NORCURON®)

1. PHARMACOLOGY
   Neuromuscular blocking agent (non-depolarizing)

2. PHARMACOKINETICS
   Paralyzes skeletal muscles, including respiratory muscles

3. INDICATIONS
   a) For treatment of ventilatory difficulty secondary to bucking or combativeness in intubated patients
   b) Patients with a history of malignant hyperthermia or contraindications to succinylcholine

4. CONTRAINDICATIONS
   Patients with known hypersensitivity to the drug

5. ADVERSE EFFECTS
   a) Bradycardia
   b) Prolonged paralysis

6. PRECAUTIONS
   a) Sedation must be provided prior to administering vecuronium when administered to a patient who is either responsive to stimulus or who may become responsive to stimulus during neuromuscular blockade.
   b) Paralysis occurs within 2–4 minutes and generally lasts 25–40 minutes.

7. DOSAGE/ROUTE
   a) RSI procedure
      Adult:
      0.1 mg/kg IVP/IO; if inadequate paralysis after 2-3 minutes, verify IV/IO patency.
      Repeat vecuronium 0.05 mg/kg IVP/IO.
   b) Ventilatory bucking or combativeness
      Adult:
      (1) Administer vecuronium 0.05 mg/kg IVP/IO. Maximum single dose is 10 mg.
      (2) Dose may be repeated once in 2-3 minutes, if necessary.
   c) Pediatric:
      (1) Administer vecuronium 0.05 mg/kg IVP/IO (may not be used for patients with needle cricothyroidotomy because of inability to monitor breath to breath ETCO₂).
      Maximum single dose is 10 mg.
      (2) Dose may be repeated once in 2-3 minutes, if necessary.
SPECIALTY CARE PARAMEDIC (Paramedic only)

The scope of practice for the specialty care paramedic (SP) is defined by a floor and a ceiling of care. The entry level for this program is Maryland licensed paramedic. The floor of the SP is the existing *The Maryland Medical Protocols for Emergency Medical Services*, including the Optional Supplemental Protocols: CPAP, Glycoprotein II/III Antagonist, Heparin, Scene/Chronic Ventilator, and Mark I/DuoDote. (The Optional Supplemental protocols Wilderness and Transport of Acute Ventilator Interfacility Patient are not included as part of ALS transports.) The medications and procedures listed within *The Maryland Medical Protocols for Emergency Medical Services* may be administered by the SP based on the written interfacility transfer orders of the sending medical director of the commercial specialty care service (without manipulation of *The Maryland Medical Protocols for Emergency Medical Services*) or receiving physician without having to request online base station medical consultation.

The ceiling for the SP is defined by the medications and procedures that are defined as “RN” or are not listed within the tables below. Those medications or skills that are listed as “RN” require familiarization by the SP but are the responsibility of the transport nurse or physician constituting the patient care team.

If a medication or procedure is listed within the scope of practice for the SP, it applies to both adult and pediatric patients unless otherwise noted.

The practice environment for these medications and procedures will be strictly for the interfacility transfer of patients and not extended into the realm of the 9-1-1 response unless otherwise noted.

<table>
<thead>
<tr>
<th>Classification of Drugs and Procedures</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SP</strong></td>
</tr>
<tr>
<td>A specialty care paramedic (SP) may initiate, monitor, and maintain without a transport nurse if they have successfully completed an EMS Board-approved specialty care program. (The commercial ambulance must still meet the requirement of an additional ALS clinician and EMT driver to complete the specialty care transport.)</td>
</tr>
<tr>
<td><strong>RN</strong></td>
</tr>
<tr>
<td>A transport nurse or physician is onboard – SP needs familiarity with the medication or procedure but SP may not perform or administer.</td>
</tr>
</tbody>
</table>
## Optional Supplemental Protocol – SPECIALTY CARE PARAMEDIC (continued)

<table>
<thead>
<tr>
<th>Medication - Procedure</th>
<th>Specialty Care Paramedic (SP)</th>
<th>Team with Nurse (RN)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A. Medications</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Sedatives</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Etomidate (Amidate®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>b. Lorazepam (Ativan®)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>c. Midazolam (Aersed®)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>d. Propofol (Diprivan®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>2. Analgesics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Fentanyl (Sublimaze®)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>b. Hydromorphone (Dilaudid®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>c. Meperidine (Demerol®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>d. Non-narcotic analgesics (e.g., Ketorolac)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>3. Paralytics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. All types</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>4. Antihypertensives</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. All types</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>5. Volume Expanders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Albumin</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>b. Blood products (except Pilot-Whole Blood)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>c. Dextran</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>d. Hespan</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>e. Plasmanate</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>6. Vasopressors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Dobutamine (Dobutrex®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>b. Epinephrine – drip</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>c. Norepinephrine (Levaphed®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>d. Phenylylephrine</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>7. Bronchodilators</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Metaproterenol (Alupent®)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>b. Theophylline – IV</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>c. Terbutaline (Brethine®) - Inhaled</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>d. L-Albuterol (inhaled)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>8. Anti-Anginals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Atenolol (Tenormin®)</td>
<td>SP (adults only)</td>
<td>RN</td>
</tr>
<tr>
<td>b. Metoprolol (Lopressor®)</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>c. Nitroglycerin (Tridil®) – IV</td>
<td>SP (adults only)</td>
<td></td>
</tr>
<tr>
<td>d. Propranolol (Inderal®)</td>
<td>SP</td>
<td>RN</td>
</tr>
</tbody>
</table>
### Medication - Procedure (Continued)

<table>
<thead>
<tr>
<th>Medications (Continued)</th>
<th>Specialty Care Paramedic (SP)</th>
<th>Team with Nurse (RN)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>9. Fibrinolytics/Thrombolitics</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. All types</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td><strong>10. Anti-Coagulants/Anti-Platelets</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. All Types</td>
<td>SP (adults only)</td>
<td></td>
</tr>
<tr>
<td><strong>11. Anti-Emetic</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. All types anti-emetic</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td><strong>12. Miscellaneous</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Flumazenil AD (Romazicon®)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>b. Insulin – IV</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>c. Insulin in TPN</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>d. Mannitol (Osmitrol®)</td>
<td></td>
<td>SP</td>
</tr>
<tr>
<td>e. Magnesium Sulfate (added to mixed drip – e.g., with vitamins)</td>
<td></td>
<td>SP</td>
</tr>
<tr>
<td>f. Potassium Chloride (only maintenance infusions; not bolusing)</td>
<td></td>
<td>SP</td>
</tr>
<tr>
<td>g. Sodium Bicarbonate Drip</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>h. Steroids – IV (not initiated)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>i. Tocolytics (including Magnesium Sulfate)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>j. Uterine stimulants (e.g., oxytocin)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td><strong>13. Anti-Arrhythmic</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Bretylium (Bretylol®)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>b. Digoxin (Lanoxin®)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>c. Diltiazem Drip</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>d. Esmolol (Brevibloc®)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>e. Metoprolol (Lopressor®)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>f. Procainamide (Pronestyl®)</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>g. Quinidine Sulfate &amp; Gluconate</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td><strong>14. Anti-Convulsants (also see sedatives)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a. Barbiturates</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>b. Phenytoin (Dilantin®)/Fosphenytoin</td>
<td></td>
<td>SP</td>
</tr>
<tr>
<td>c. Other non-benzodiazepine anti-convulsants</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td><strong>15. Diuretics</strong></td>
<td>SP</td>
<td></td>
</tr>
</tbody>
</table>
### Optional Supplemental Protocol – SPECIALTY CARE PARAMEDIC (continued)

#### B. Invasive Procedures

<table>
<thead>
<tr>
<th>Medication - Procedure (Continued)</th>
<th>Specialty Care Paramedic (SP)</th>
<th>Team with Nurse (RN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Chest Escharotomies</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>2. Chest Tubes Insertion</td>
<td></td>
<td>RN</td>
</tr>
<tr>
<td>3. Chest Tube or Surgical Drain with vacuum system/placed to suction</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>4. Laryngeal Mask Airway</td>
<td>SP (adult only)</td>
<td></td>
</tr>
<tr>
<td>5. Needle Cricothyroidotomy</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>6. Rapid Sequence Intubation</td>
<td>SP</td>
<td>RN</td>
</tr>
<tr>
<td>7. Surgical Cricothyroidotomy</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>8. Urinary catheter insertion</td>
<td>SP</td>
<td></td>
</tr>
</tbody>
</table>

#### C. Non-Invasive Procedures

<table>
<thead>
<tr>
<th>Medication - Procedure (Continued)</th>
<th>Specialty Care Paramedic (SP)</th>
<th>Team with Nurse (RN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. IV Pumps</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>2. Ostomy care</td>
<td>SP</td>
<td></td>
</tr>
</tbody>
</table>

#### D. System Monitoring

<table>
<thead>
<tr>
<th>Medication - Procedure (Continued)</th>
<th>Specialty Care Paramedic (SP)</th>
<th>Team with Nurse (RN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Arterial Line/Cardiac Sheath</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>2. CVP line (monitor but not performing measures)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>3. Intracranial Pressure Monitor/Line</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>4. Swan-Ganz</td>
<td>RN</td>
<td></td>
</tr>
</tbody>
</table>

#### E. Specialized Equipment

<table>
<thead>
<tr>
<th>Medication - Procedure (Continued)</th>
<th>Specialty Care Paramedic (SP)</th>
<th>Team with Nurse (RN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Acute Ventilated Interfacility Patient – Transport Service’s Ventilator (except as in E6)</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>2. Internal Pacer with external control</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>3. Intra-Aortic Balloon Pump</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>4. Peritoneal Dialysis Systems</td>
<td>SP</td>
<td></td>
</tr>
<tr>
<td>5. Specialty Ventilator (e.g., pediatric or when hospital ventilator must accompany patient)</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>6. Transport Isolette/Incubator</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td>7. Ventricular Assist Devices</td>
<td>*</td>
<td></td>
</tr>
</tbody>
</table>

*If being discharged home or to rehab, may go by ALS.
1. **INDICATIONS**
   Patients must have reached their 15th birthday and may present with any of the following conditions:
   a) Inability to oxygenate despite having tried BVM with oropharyngeal/nasopharyngeal airway, ET placement, and supraglottic airway (if not contraindicated)
   b) Inability to place ET in the setting of life-threatening upper airway hemorrhage
   c) Completely obstructing upper airway foreign body that cannot be removed via BLS maneuvers or Magill forceps with direct visualization

2. **EQUIPMENT**
   CLINICIANS MAY USE PRE-ASSEMBLED EQUIPMENT OR AN FDA-APPROVED KIT, AS PRESCRIBED BY THE PROGRAM MEDICAL DIRECTOR.

3. **PROCEDURE**
   a) Clinicians must use a designated technique and procedure for establishing the airway through the cricothyroid membrane that has been approved by the program medical director as part of this pilot.
   b) Upon completion of the skill (or at an appropriate time during the sequence of patient care) the clinician will obtain medical direction and also notify the receiving physician/emergency department with the following information:
      (1) Patient condition
      (2) Reason for surgical cricothyroidotomy
      (3) Complications arising from procedure (if any)
      (4) Patient response to treatment
1. INTRODUCTION

a) Scope and Applicability

1. These protocols are intended for use during high-risk, large-scale, and extended law enforcement or homeland security operations.

2. The Tactical Emergency Medical Services (TEMS) clinician is not directly responsible for any person(s) outside the direct field of operations, whose care may safely be provided by the local EMS Operational Program.

3. These protocols supplement the current version of *Maryland Medical Protocols for Emergency Medical Services* and, at the Tactical Physician’s discretion, may incorporate other EMS protocol components such as: Wilderness, Interfacility, Pilot/Optional, and WMD sections.

4. The Tactical Emergency Medical Services Protocols shall be used only by Tactical EMS clinicians sponsored by a law enforcement agency and operating under law enforcement command.

5. To be approved, there must be a written, integrated relationship between the EMS Operational Program and the TEMS program, with both the EMS Operational Program Medical Director and the TEMS Medical Director having signed off on the agreement.

6. Tactical EMS Clinicians at the EMT or ALS levels may administer the medications and perform the procedures listed in these protocols only after receiving specific training on their use and only under the medical direction of a Tactical Physician.

7. The primary function of the Tactical EMS Clinician is to support law enforcement or homeland security operations by facilitating the health and safety of critical public safety personnel inside the perimeter of high-risk, large-scale, and extended operations.

8. Once the patient is removed from the law enforcement perimeter of operations, the TEMS Protocol will end, *The Maryland Medical Protocols for Emergency Medical Services* will be implemented, and the transition of care will be made to the local EMS agency.

9. An exception may be made when the Tactical EMS Clinician’s specialized training is needed to manage a specific illness/injury.

   a) If the Tactical EMS Clinician’s specialized training is needed to manage the patient’s illness/injury, then the highest-trained Tactical EMS Clinician shall ride to the hospital with the patient to maintain medications that are not allowed by *The Maryland Medical Protocols for Emergency Medical Services*.

   b) If, during transport, Tactical EMS personnel encounter a significant conflict between TEMS Protocols and those of the transporting EMS agency, they should attempt to contact their own Tactical Physician and request a dual consult with the local Base Station Physician.

   c) If they cannot reach a Tactical Physician, they should contact the local EMS Base Station for on-line medical consultation.
b) Definition of Tactical Environment
   (1) Any law enforcement or homeland security operation where deployed personnel are in a large-scale operation or where the risk of injury is sufficiently high as to warrant the presence of on-scene emergency medical services clinicians.
   (2) Types of operations may include: high-risk warrant service, hostage-barricade situations, emergency ordinance disposal, executive protection details, civil demonstration or protest, dynamic training operations, aquatic operations, high-angle, search and rescue missions, and acts of terrorism.
   (3) Any prolonged law enforcement deployment, where performance decrement or environmental issues may arise and the safety of the public and deployed law enforcement personnel would benefit from the presence of a Tactical EMS Clinician to monitor these circumstances.

c) Demonstration of Need
   (1) Jurisdictions that seek approval for a Tactical EMS Program shall submit a demonstration-of-need letter outlining the necessity for the program.
   (2) The letter shall be submitted to the State EMS Medical Director for approval and include the following:
      (a) Name of organization and scope of the proposed Tactical EMS Team
      (b) Name and qualifications of the Tactical Medical Director and other Tactical Physicians
      (c) Name and qualifications of the Tactical EMS Coordinator and other Tactical EMS Clinicians

d) Sponsoring Law Enforcement Agency Requirements
   (1) Sponsoring Law Enforcement Agencies shall be responsible for
      (a) Completing background investigations appropriate for medical clinicians working in and around law enforcement operations
      (b) Providing appropriate personal protective equipment, to accommodate conditions that the team may reasonably encounter, to the Tactical EMS Clinicians and Tactical Physician(s) and ensure adequate training in the equipment's use and capabilities
      (c) Providing written documentation to MIEMSS that addresses the medical liability and personal injury considerations of the Tactical EMS Clinicians/Physician(s)

e) Tactical EMS Clinician/Tactical Physician Minimum Training Requirements
   (1) The Tactical EMS Clinician shall be a Maryland-certified EMT or Maryland-licensed ALS clinician and have successfully completed a nationally-recognized Counter-Narcotic Tactical Operation Medical Support/Integrated Force Health Clinician Program (CONTOMS/IFHP) or equivalent Tactical Clinician course that includes instruction and training in
      (a) Team wellness and health management, including preventive medicine
      (b) Providing care under fire/basic weapons safety
      (c) Officer rescue
      (d) Planning medical operations and medical intelligence
      (e) Response to the active shooter
Optional Supplemental Protocol – TACTICAL EMS (continued)

(f) Orientation to specialized medical gear personal protective equipment used in tactical medical operations
(g) Remote medical assessment ("medicine across the barricade")
(h) Response and management of WMD events, including field-expedient decontamination ("hasty decon") procedures
(i) Operational security, light and sound discipline, helicopter operations, pyrotechnic and other chemical agents, as utilized by law enforcement teams
(j) Less-than-lethal weaponry, the injuries they may cause, and any specific interventions required

(2) The Tactical EMS Clinician shall have responsibilities for part or all of these protocols, as summarized as follows, based on either EMT or ALS (CRT-I or paramedic) level certification.

<table>
<thead>
<tr>
<th>INTERVENTION</th>
<th>EMT</th>
<th>ALS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Provision of access to medications: ibuprofen, naproxen, fexofenadine, cetirizine, pseudoephedrine, oxymetazoline nasal spray, Mylanta®, cimetidine, loperamide, clove oil, acetaminophen, tramadol, caffeine, modafinil, ondansetron OD, scopolamine patch, ophthalmologic proparacaine/tetracaine and fluorescein, prednisone PO, dexamethasone PO, albuterol MDI, aspirin, epinephrine 1 mg/mL IM, naloxone IN, glucose PO</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>Administration of medications in Protocol, not listed above</td>
<td></td>
<td>•</td>
</tr>
<tr>
<td>Cyanoacrylate tissue adhesive</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>Field expedient wound closure (stapling)</td>
<td></td>
<td>•</td>
</tr>
<tr>
<td>Conducted electrical weapon (CEW) dart removal</td>
<td>•</td>
<td>•</td>
</tr>
</tbody>
</table>

(3) The Tactical EMS Clinician shall document each patient contact utilizing a patient care report (PCR) (eMedS®). The documentation must be consistent with current MIEMSS regulations for interventions, as summarized in the above table.

(4) The Tactical Physician shall possess an unrestricted Maryland License (preferred Emergency Medicine, General/Orthopedic/Trauma Surgery, or Critical Care), have experience in on-line medical direction, and have completed a nationally-recognized (CONTOMS/IFHP or equivalent) tactical medical director's course that includes instruction and training in the following topics:
(a) History of/need for tactical EMS provision
(b) Administrative/command concerns and responsibilities
(c) Care under fire
(d) Special equipment/hazards in the tactical environment
(e) Forensic examination
(f) Medicine "across the barricade”
(g) Medical threat assessment
Optional Supplemental Protocol –
TACTICAL EMS (continued)

f) Quality Assurance Properties
   (1) Individual Tactical EMS Clinicians must be approved for TEMS Program Participation by the TEMS Medical Director.
   (2) Classroom lecture
   (3) Mannequin instruction
   (4) Must demonstrate proficiency through skills testing and written test
   (5) Ongoing demonstration of proficiency
   (6) A verification of all TEMS skills and review of TEMS principles of safety will be performed on an annual basis by the Medical Director, or the clinician may document utilization of skills in the field
   (7) Review of each call
      (a) Upon completion of the tactical incident, notification of any implementation of the TEMS Protocol will be made to your jurisdictional TEMS supervisor, who will ensure notification to TEMS Medical Director.
      (b) TEMS Medical Director will review and evaluate all TEMS interventions within 48 hours of resolution of the tactical incident and provide feedback.
   (8) The TEMS program will maintain a detailed TEMS database and will provide an annual report to the State EMS Medical Director.

2. GENERAL PROTOCOLS
   a) Medical Direction
      (1) Tactical EMS Clinicians may provide medical care using Tactical Medical Protocols only under the medical direction of a Tactical Physician.
      (2) Immediately available telephone or radio contact during an operation shall be considered a reasonable substitute for in-person supervision of Tactical EMS Clinicians.
      (3) In the absence of medical direction by a Tactical Physician, jurisdictional trained and designated Tactical EMS Clinicians should defer to their usual EMS protocols.
   b) Operational Command
      (1) Operational command within a law enforcement perimeter of operation lies with the law enforcement commander. At times, the safety and success of the law enforcement objectives may override the need to care for casualties. The law enforcement commander is responsible for the care and movement of casualties within a law enforcement operation.

3. SPECIAL CONSIDERATION FOR TACTICAL EMS
   a) The execution of some law enforcement operations may require that minor illness or injury in essential public safety personnel be treated and, to the extent that it is medically safe to do so, that those treated personnel return to duty. Fitness for duty of public safety personnel with minor injuries or illnesses shall be determined by the law enforcement commander in consultation with a Tactical Physician.
   b) Prescription and over-the-counter (OTC) medications may be used for the treatment (or “symptomatic relief”) of constitutional symptoms as required to promote the health, safety, and functionality of persons necessary to the operation.
Optional Supplemental Protocol –
TACTICAL EMS (continued)

The Tactical EMS Clinician(s) under the Tactical Physician will know the indications/contraindications for the medications available to them (as will be delineated under “Additional Medications for Tactical EMS,” to follow). At the EMT level, medications will be made available to those persons under the Tactical Clinician’s care to self-select and self-medicate at the individual requesting person’s own discretion regarding appropriateness of use.

c) The Tactical EMS Clinician may provide care to all persons associated with the operation, and shall be responsible for initial access, assessment, and stabilization (within the scope of The Maryland Medical Protocols for Emergency Medical Services) of those victims, bystanders, and suspects within the “warm” or “hot” zones until they may be extracted to local EMS clinicians. The Tactical EMS clinician is not directly responsible for any person(s) outside the direct field of operations, whose care may safely be provided by the local EMS Operational Program.

4. SPECIFIC PROCEDURES

a) Cyanoacrylate tissue adhesive
   (1) Purpose: To limit blood loss, pain, and risk of secondary contamination/injury to a minor open wound
   (2) Indications
      (a) Clean wounds
      (b) Minor bleeding wounds difficult to control with other interventions
      (c) Wounds in personnel who must remain operational
   (3) Contraindications
      (a) Grossly contaminated wounds
      (b) Greater than two hours since infliction of wound
      (c) Macerated/ crushed surrounding tissue
      (d) Wounds near the eyes
   (4) Potential adverse effects/ complications
      (a) This is not intended to constitute definitive wound closure; however, if properly cleaned prior to procedure, may be reviewed by physician without further intervention.
      (b) Transient local pain at application site may be reported.
   (5) Precautions
      (a) Ask regarding previous reaction/exposure to agent.
      (b) Advise patient of requirement for further evaluation by physician.

b) “Field expedient” wound closure (stapling)
   (1) Purpose: To limit blood loss and risk of secondary contamination injury to an open wound.
   (2) Indications
      (a) Clean wounds
      (b) Delay in transportation to definitive care will be or is anticipated to be several hours
      (c) Bleeding wounds difficult to control with other interventions
      (d) Wounds in personnel who must remain operational
Optional Supplemental Protocol – TACTICAL EMS (continued)

(3) Contraindications
   (a) Grossly contaminated wounds
   (b) Greater than six hours since infliction of wound
   (c) Macerated/crushed surrounding tissue
   (d) Situations with less than two hours anticipated time to transportation to definitive care
   (e) Facial wounds

(4) Potential adverse effects/complications
   (a) This is not intended to constitute definitive wound closure—this will minimize the risk for increased infection and increased foreign body retention.

(5) Precautions
   (a) Ask regarding local anesthetic allergies.
   (b) Advise patient of requirement for further evaluation by physician.
   c) Impaled conducted electrical weapon dart removal
      (1) ANY conducted electrical weapon dart impalement to the head, neck, hands, feet, or genitalia must be stabilized in place and evaluated by a physician.
      (2) In order to safely transport the patient, attempted extraction may be made one time by a Tactical EMS Clinician as long as the dart is not lodged in a location listed in (1) above and is not fully embedded up to the hub in tissue.
      (3) All patients receiving conducted electrical weapon intervention will need to be transported to the emergency department for assessment.

5. SUPPLEMENTAL FORMULARY FOR TACTICAL EMS
   a) Tactical EMS clinicians may administer the following medications to support and maintain Tactical personnel in the operation environment. Bolded medications are required as part of the standardized TEMS load-out at the EMT or ALS level; the others are optional.
      (1) Antihistamines/Decongestants
          (a) Pseudoephedrine (Sudafed®)
          (b) Cetirizine (Zyrtec®)
          (c) Diphenhydramine (Benadryl®)
          (d) Fexofenadine (Allegra®)
          (e) Oxymetazoline nasal spray (Afrin®)
      (2) Gastrointestinal
          (a) Antacid (Mylanta® or other equivalent antacid)
          (b) Cimetidine (Tagamet®—or other equivalent H2 blocker)
          (c) Loperamide (Imodium®)
          (d) 5-HT3 Antagonist (Zofran® ODT/Ondansetron, 5-HT3 antagonist)
          (e) Metoclopramide (Reglan®) (injectable)
          (f) Dimenhydrinate (Dramamine®)
          (g) Meclizine (Antivert®) (for motion sickness)
          (h) Scopolamine transdermal
Optional Supplemental Protocol – TACTICAL EMS (continued)

(3) Ophthalmologicals
   (a) Proparacaine or tetracaine (Alcaine®) ophthetic
   (b) Fluorescein stain (and blue light)
   (c) Eye irrigation solution
   (d) Erythromycin ophthalmic ointment
   (e) pH paper

(4) Antimicrobials/antiviral (agent-specific training)
   (a) Ciprofloxacin (following exposure or prophylaxis)
   (b) Triple Antibiotic Ointment (Bacitracin®/Polymyxin®/Neomycin®)
   (c) Amoxicillin/clavulanic acid (Augmentin®)
   (d) Cefazolin (Ancef®) (PO or IV) (for trauma applications when transport delayed)
   (e) Clindamycin (Cleocin®)
   (f) Trimethoprim/sulfamethoxazole (Bactrim®)
   (g) Azithromycin (Zithromax®)
   (h) Doxycycline
   (i) Mupirocin topical ointment (Bactroban®)
   (j) Emtricitabine and tenofovir (Truvada®) (high-risk post-exposure management)

(5) Steroids
   (a) Prednisone (PO)
   (b) Dexamethasone (Decadron®) (IV/IM and/or PO)

(6) Analgesics/anesthetics
   (a) Acetaminophen (PO)
   (b) Ibuprofen (Motrin®/Advil®)
   (c) Naproxen (Aleve®/Naprosyn®) (PO)
   (d) Tramadol (Ultram®) (PO)
   (e) Ketamine
   (f) Naloxone (Narcan®) (IN and/or IV)
   (g) Lidocaine (transdermal for muscular relief, or IM/SQ for stapling as temporizing measure only, alternate dosing regimen)
   (h) Fentanyl transmucosal (PO)
   (i) Clove oil (for topical dental analgesia)
   (j) Ketorolac (Toradol®) (Injectable)

(7) Sleep/wake
   (a) Caffeine (No-Doz®)
   (b) Zaleplon (Sonata®) (sleeper)
   (c) Modafinil (Provigil®)

(8) Wound management
   (a) Cyanoacrylate tissue adhesive (Dermabond®)
   (b) Topical hemostatic agent
   (c) Steri-strips
   (d) Staples

(9) ACLS/resuscitation
   (a) Albuterol MDI
Optional Supplemental Protocol – TACTICAL EMS (continued)

(10) Anti-hypoglycemics
   (a) Oral glucose

(11) Additional Medications for Tactical EMS: The following is a list of medications from the Maryland Medical Protocols that is strongly encouraged to be readily accessible to complement the Tactical Medic's Formulary.
   Aspirin (EMT, ALS) ................................................. Non-Operational
   Atropine multi-dose (ALS) .......................................... Non-Operational
   Dexamethasone (ALS) ............................................. Operational
   Dextrose (ALS) ....................................................... Non-Operational
   Epinephrine (1 mg/mL) (EMT, ALS) ......................... Non-Operational
   Haldol (ALS) ........................................................... Non-Operational
   Morphine or fentanyl for injection (ALS) .................... Non-Operational
   Midazolam (ALS) ..................................................... Non-Operational
   Nitroglycerin (ALS) ................................................ Non-Operational

OPERATIONAL: THE MEDICATION MAY BE GIVEN TO A LAW ENFORCEMENT MEMBER WHO MAY CONTINUE TO PERFORM THEIR ASSIGNED DUTIES.

NON-OPERATIONAL: ONCE THE MEDICATION HAS BEEN ADMINISTERED, THE LAW ENFORCEMENT MEMBER IS REMOVED FROM THEIR ASSIGNED DUTIES SINCE THE MEDICATION OR THE ASSOCIATED MEDICAL/TRAUMATIC COMPLAINT FOR WHICH THE MEDICATION IS INDICATED MAY IMPAIR THEIR ABILITY TO PERFORM CRITICAL LAW ENFORCEMENT TASKS AND DUTIES.

b) Tactical EMS Medical Formulary
   (1) Antihistamines/Decongestants
      (a) Pseudoephedrine (Sudafed®)
         (i) AVAILABILITY.................................30 mg or 60 mg tablets (OTC)
         (ii) ACTION........................................Decongestant
         (iii) INDICATIONS..............................Nasal congestion; rhinorrhea
         (iv) CONTRAINDICATIONS..............Known hypersensitivity; hypertension
         (v) PRECAUTIONS.........................N/A
         (vi) OPERATIONAL STATUS..............Operational
         (vii) SIDE EFFECTS.........................Insomnia
         (viii) INTERACTIONS.....................N/A
         (ix) DOSAGE..............................30–60 mg, every 4–6 hours, as needed
      (b) Cetirizine (Zyrtec®)
         (i) AVAILABILITY...............................10 mg tablet
         (ii) ACTION......................................Non-sedating antihistamine
         (iii) INDICATIONS..............................Allergic symptoms
         (iv) CONTRAINDICATIONS..............Known hypersensitivity
         (v) PRECAUTIONS.........................Hypertension; liver/kidney dx
         (vi) OPERATIONAL STATUS..............Operational
         (vii) SIDE EFFECTS.........................Dry mouth, urinary retention
         (viii) INTERACTIONS.....................N/A
         (ix) DOSAGE..............................10 mg/once daily
Optional Supplemental Protocol –
TACTICAL EMS (continued)

(c) Diphenhydramine (Benadryl®)
(i) AVAILABILITY..........................25 mg or 50 mg tablets
(ii) ACTION................................Sedating antihistamine
(iii) INDICATIONS............................Allergic symptoms
(iv) CONTRAINDICATIONS...........Known hypersensitivity
(v) PRECAUTIONS..........................Hypertension; liver/kidney dx
(vi) OPERATIONAL STATUS..............NON-OPERATIONAL
(vii) SIDE EFFECTS........................Dry mouth, urinary retention, somnolence
(viii) INTERACTIONS......................N/A
(ix) DOSAGE..................................25–50 mg every 4–6 hours, as needed; per MD/DO

(d) Fexofenadine (Allegra®)
(i) AVAILABILITY..........................60 mg tablet
(ii) ACTION................................Non-sedating antihistamine
(iii) INDICATIONS............................Allergic symptoms
(iv) CONTRAINDICATIONS...........Known hypersensitivity
(v) PRECAUTIONS..........................Hypertension history
(vi) OPERATIONAL STATUS..............Operational
(vii) SIDE EFFECTS........................Dry mouth, urinary retention
(viii) INTERACTIONS......................N/A
(ix) DOSAGE..................................60mg/once or twice daily

(e) Oxymetazoline nasal spray (Afrin®)
(i) AVAILABILITY..............................Nasal spray 0.05%
(ii) ACTION................................Nasal vasoconstriction; decongestant
(iii) INDICATIONS............................Rhinorrhea; sinus congestion and pain
(iv) CONTRAINDICATIONS...........Known hypersensitivity
(v) PRECAUTIONS..........................Hypertension
(vi) OPERATIONAL STATUS..............Operational
(vii) SIDE EFFECTS........................Nosebleed (minor) possible; often used in treatment of nosebleed
(viii) INTERACTIONS......................N/A
(ix) DOSAGE..................................Two sprays per nare, 2–3 times per day

(2) Gastrointestinal
(a) Antacid (Mylanta® or other equivalent antacid)
(i) AVAILABILITY...........................Liquid (OTC)
(ii) ACTION................................Antacid
(iii) INDICATIONS.............................GI upset, GERD, PUD, gastritis, esophagitis
(iv) CONTRAINDICATIONS...........Known hypersensitivity
Optional Supplemental Protocol –
TACTICAL EMS (continued)

(v) PRECAUTIONS.........................Some medications require acidic pH and should not be taken at same time with this medication

(vi) OPERATIONAL STATUS.........Operational

(vii) SIDE EFFECTS.......................N/A

(viii) INTERACTIONS....................Loose stools possible

(ix) DOSAGE.................................15–45 mL every 4–8 hours

(b) Cimetidine (Tagamet®—or other equivalent H2 blocker)

(i) AVAILABILITY..........................200/300/400 mg tablets; 300 mg IV/IM

(ii) ACTION.....................................H2 blocker

(iii) INDICATIONS.........................PUD, GERD, esophagitis, gastritis

(iv) CONTRAINDICATIONS..............Known hypersensitivity; concomitant Proton Pump Inhibitor (PPI) use

(v) PRECAUTIONS.........................N/A

(vi) OPERATIONAL STATUS.........Operational

(vii) SIDE EFFECTS.......................N/A

(viii) INTERACTIONS....................N/A

(ix) DOSAGE.................................300 mg IV/IM/PO every 6–8 hours; 400 mg twice daily

(c) Loperamide (Imodium®)

(i) AVAILABILITY..........................2 mg tablet (OTC) and 1mg/5mL suspension

(ii) ACTION.....................................Anti-diarrheal

(iii) INDICATIONS.........................Diarrhea

(iv) CONTRAINDICATIONS..............Known hypersensitivity; hypertension; bloody diarrhea

(v) PRECAUTIONS.........................N/A

(vi) OPERATIONAL STATUS.........Operational

(vii) SIDE EFFECTS.......................ENT dryness

(viii) INTERACTIONS....................N/A

(ix) DOSAGE.................................4 mg first dose; 2 mg each subsequent episode until stool formed; maximum 16 mg per day

(d) 5-HT3 Antagonist (Zofran® ODT/Ondansetron, 5-HT3 antagonist)

(i) AVAILABILITY..........................IM/IV injectable; tablets

(ii) ACTION.....................................Anti-emetic; anti-motion sickness

(iii) INDICATIONS.........................Nausea/vomiting

(iv) CONTRAINDICATIONS..............Known hypersensitivity

(v) PRECAUTIONS.........................Long QT history

(vi) OPERATIONAL STATUS.........Operational

(vii) SIDE EFFECTS.......................N/A

(viii) INTERACTIONS....................N/A

(ix) DOSAGE.................................Per MD/DO
Optional Supplemental Protocol – TACTICAL EMS (continued)

(e) Metoclopramide (Reglan®) (injectable)
   (i) AVAILABILITY.................................IM/IV injectable; 10 mg
   (ii) ACTION......................................Anti-emetic; promotes
        GI motility
   (iii) INDICATIONS.........................Nausea/vomiting
   (iv) CONTRAINDICATIONS................Known hypersensitivity
   (v) PRECAUTIONS
       Dystonic reaction risk (treat with diphenhydramine);
       may see sedation
   (vi) OPERATIONAL STATUS................NON-OPERATIONAL
   (vii) SIDE EFFECTS...........................Sedation; dystonia
   (viii) INTERACTIONS..........................N/A
   (ix) DOSAGE..................................10–20 mg IM/IV/PO every 4 hours,
       as needed; per MD/DO

(f) Dimenhydrinate (Dramamine®)
   (i) AVAILABILITY.................................IM/IV injectable; 50 mg tablet
   (ii) ACTION......................................Anti-emetic; anti-motion sickness
   (iii) INDICATIONS.........................Nausea/vomiting
   (iv) CONTRAINDICATIONS................Known hypersensitivity
   (v) PRECAUTIONS
       May see sedation
   (vi) OPERATIONAL STATUS................NON-OPERATIONAL
   (vii) SIDE EFFECTS...........................Sedation
   (viii) INTERACTIONS..........................N/A
   (ix) DOSAGE..................................50–100 mg IM/IV/PO every 4 hours,
       as needed; per MD/DO

(g) Meclizine (Antivert®) (for motion sickness)
   (i) AVAILABILITY.................................25–50 mg tablet
   (ii) ACTION......................................Anti-emetic; anti-motion sickness
   (iii) INDICATIONS.........................Nausea/vomiting
   (iv) CONTRAINDICATIONS................Known hypersensitivity
   (v) PRECAUTIONS
       May see sedation
   (vi) OPERATIONAL STATUS................NON-OPERATIONAL
   (vii) SIDE EFFECTS...........................Sedation
   (viii) INTERACTIONS..........................N/A
   (ix) DOSAGE..................................25–50 mg PO every 4 hours,
       as needed; per MD/DO

(h) Scopolamine transdermal
   (i) AVAILABILITY.................................1 mg patch
   (ii) ACTION......................................Anti-emetic; anti-motion sickness
   (iii) INDICATIONS..............................Nausea/vomiting/motion sickness prevention
   (iv) CONTRAINDICATIONS................Known hypersensitivity, hx angle closure glaucoma; hypersensitivity to belladonna alkaloids, seizures, urinary retention
   (v) PRECAUTIONS
       May cause sedation, disorientation underwater
Optional Supplemental Protocol – TACTICAL EMS (continued)

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(vi) OPERATIONAL STATUS: Operational (if previously tolerated scopolamine)

(vii) SIDE EFFECTS: Sedation

(viii) INTERACTIONS: Use with caution when taking other potentially sedative drugs or anticholinergics

(ix) DOSAGE: 1 mg patch every 3 days, as needed; per MD/DO

(3) Ophthalmologicals

(a) Proparacaine or Tetracaine (Alcaine®) opthalmic

(i) AVAILABILITY: Ocular anesthetic solution

(ii) ACTION: Topical anesthetic

(iii) INDICATIONS: To facilitate eye exam; relieve eye pain; per MD/DO

(iv) CONTRAINDICATIONS: Known hypersensitivity

(v) PRECAUTIONS: Ensure eye protection from foreign objects after exam

(vi) OPERATIONAL STATUS: Operational

(vii) SIDE EFFECTS: N/A

(viii) INTERACTIONS: Eye pain

(ix) DOSAGE: 1–2 drops per eye; per MD/DO

(b) Fluorescein stain (and blue light)

(i) AVAILABILITY: Single application strips

(ii) ACTION: Dye to facilitate eye exam

(iii) INDICATIONS: Suspected eye injury (foreign body/corneal abrasion)

(iv) CONTRAINDICATIONS: Known hypersensitivity

(v) PRECAUTIONS: N/A

(vi) OPERATIONAL STATUS: Operational

(vii) SIDE EFFECTS: N/A

(viii) INTERACTIONS: N/A

(ix) DOSAGE: One drop per eye

(c) Eye irrigation solution

(i) AVAILABILITY: 100 mL, 200 mL bottles (other sizes may also be available)

(ii) ACTION: To facilitate irrigation of contaminants from the eye

(iii) INDICATIONS: Following exposure of foreign body or chemical to eye

(iv) CONTRAINDICATIONS: Known hypersensitivity

(v) PRECAUTIONS: Not be used in penetrating eye trauma

(vi) OPERATIONAL STATUS: Operational

(vii) SIDE EFFECTS: N/A

(viii) INTERACTIONS: N/A

(ix) DOSAGE: Irrigate until an eye pH of 7.4 is achieved
Optional Supplemental Protocol – TACTICAL EMS (continued)

(d) Erythromycin ophthalmic ointment
   (i) AVAILABILITY.................................0.5% ointment
   (ii) ACTION........................................Macrolide antibiotic
   (iii) INDICATIONS..............................Per MD/DO—infected exposures
   (iv) CONTRAINDICATIONS....................Known hypersensitivity to penicillins
   (v) PRECAUTIONS...............................Topical use only
   (vi) OPERATIONAL STATUS.....................Operational
   (vii) SIDE EFFECTS..............................GI upset; nausea/vomiting; diarrhea
   (viii) INTERACTIONS.............................N/A
   (ix) DOSAGE........................................Per MD/DO

(e) pH paper
   (i) AVAILABILITY.................................Rolls or precut pieces of paper (other sizes may also be available)
   (ii) ACTION........................................To measure baseline and repeat pH during decontamination/irrigation
   (iii) INDICATIONS..............................Following exposure to foreign body or chemical to eye or skin
   (iv) CONTRAINDICATIONS....................Known hypersensitivity
   (v) PRECAUTIONS..............................Not be used in penetrating eye trauma
   (vi) OPERATIONAL STATUS....................Operational
   (vii) SIDE EFFECTS.............................N/A
   (viii) INTERACTIONS.............................N/A
   (ix) DOSAGE........................................One strip approximately 1–2 inches; per MD/DO

(4) Antimicrobials/antiviral (agent-specific training)
   (a) Ciprofloxacin (following exposure or prophylaxis)
      (i) AVAILABILITY...............................250/500/750 mg tablets; 400 mg IVPB; 250 or 500/5 suspension
      (ii) ACTION......................................2nd generation quinolone antimicrobial agent
      (iii) INDICATIONS..............................Per MD/DO—infected exposures
      (iv) CONTRAINDICATIONS....................Known hypersensitivity
      (v) PRECAUTIONS..............................N/A
      (vi) OPERATIONAL STATUS....................Operational
      (vii) SIDE EFFECTS................................GI upset, nausea/vomiting, diarrhea, yeast infection
      (viii) INTERACTIONS............................N/A
      (ix) DOSAGE......................................Per MD/DO
Optional Supplemental Protocol – TACTICAL EMS (continued)

(b) Triple antibiotic ointment or equivalent
(Bacitracin®/Polymyxin®/Neomycin®)
(i) AVAILABILITY..............................Topical ointment
(ii) ACTION..................................Polypeptide antibiotic
(iii) INDICATIONS..........................Per MD/DO—infectious exposures
(iv) CONTRAINDICATIONS..............Known hypersensitivity
(v) PRECAUTIONS........................Topical use only
(vi) OPERATIONAL STATUS.............Operational
(vii) SIDE EFFECTS........................Local irritation, GI upset
(viii) INTERACTIONS........................N/A
(ix) DOSAGE..................................Apply to superficial scrapes, burns, wounds, prior to dry sterile dressing.

(c) Amoxicillin/clavulanate (Augmentin®)
(i) AVAILABILITY..........................875 or 125 mg tablets
(ii) ACTION................................Beta-lactamase inhibitors
(iii) INDICATIONS.........................Per MD/DO—infectious exposures
(iv) CONTRAINDICATIONS..............Known hypersensitivity to penicillins
(v) PRECAUTIONS.........................Liver/Kidney dx
(vi) OPERATIONAL STATUS.............Operational
(vii) SIDE EFFECTS.......................GI upset; nausea/vomiting; diarrhea
(viii) INTERACTIONS........................N/A
(ix) DOSAGE................................Per MD/DO

(d) Cefazolin (Ancef®) (PO or IV) (for trauma applications when transport delayed)
(i) AVAILABILITY..........................0.5–2 grams IM/IV
(ii) ACTION..................................1st generation Cephalosporin antimicrobial agent
(iii) INDICATIONS.........................Per MD/DO—infectious exposures/trauma
(iv) CONTRAINDICATIONS..............Known hypersensitivity to PCN or Cephalosporins
(v) PRECAUTIONS........................N/A
(vi) OPERATIONAL STATUS.............NON-OPERATIONAL
(vii) SIDE EFFECTS.......................GI upset, nausea/vomiting, diarrhea, yeast infection
(viii) INTERACTIONS........................N/A
(ix) DOSAGE................................Per MD/DO

(e) Clindamycin (Cleocin®)
(i) AVAILABILITY..........................150 or 300 mg tablets; reconstituted liquid 75mg/5mL
(ii) ACTION..................................Antibiotic
(iii) INDICATIONS.........................Suspected pharyngitis or respiratory infection, cellulitis
(iv) CONTRAINDICATIONS..............Hypersensitivity to clindamycin
(v) PRECAUTIONS........................N/A
(vi) OPERATIONAL STATUS.............Operational
Optional Supplemental Protocol –
TACTICAL EMS (continued)

(vii) SIDE EFFECTS......................Diarrhea
(viii) INTERACTIONS.....................N/A
(ix) DOSAGE.................................Pediatrics – 10 mg/kg every 8 hours
Adult – 300 mg every 8 hours

(f) Trimethoprim/Sulfadiazine (Bactrim®)
(i) AVAILABILITY.........................DS tablet
(ii) ACTION...............................Sulfonamide antibiotic
(iii) INDICATIONS........................Per MD/DO—infectious exposures
(iv) CONTRAINDICATIONS..............Known hypersensitivity
(v) PRECAUTIONS.........................Liver/kidney dx, anemia, thrombocytopenia
(vi) OPERATIONAL STATUS..............Operational
(vii) SIDE EFFECTS......................GI upset, nausea/vomiting, diarrhea
(viii) INTERACTIONS.....................N/A
(ix) DOSAGE.................................Per MD/DO

(g) Azithromycin (Zithromax®)
(i) AVAILABILITY.........................250 mg tablet
(ii) ACTION...............................Macrolide antibiotic
(iii) INDICATIONS........................Per MD/DO—infectious exposures
(iv) CONTRAINDICATIONS..............Known hypersensitivity to penicillins
(v) PRECAUTIONS.........................Liver/kidney dx
(vi) OPERATIONAL STATUS..............Operational
(vii) SIDE EFFECTS......................GI upset, nausea/vomiting, diarrhea
(viii) INTERACTIONS.....................N/A
(ix) DOSAGE.................................Per MD/DO

(h) Doxycycline
(i) AVAILABILITY.........................100 mg tablet
(ii) ACTION...............................Tetracycline antibiotic
(iii) INDICATIONS........................Per MD/DO—infectious exposures
(iv) CONTRAINDICATIONS..............Known hypersensitivity to tetracyclines, pregnancy
(v) PRECAUTIONS.........................Liver/kidney dx, photoreactivity rash
(vi) OPERATIONAL STATUS..............Operational
(vii) SIDE EFFECTS......................GI upset, nausea/vomiting, diarrhea
(viii) INTERACTIONS.....................N/A
(ix) DOSAGE.................................Per MD/DO

(i) Mupirocin topical ointment (Bactroban®)
(i) AVAILABILITY.........................2% topical ointment
(ii) ACTION...............................Other antibiotic
(iii) INDICATIONS........................Per MD/DO—infectious exposures
(iv) CONTRAINDICATIONS..............Known hypersensitivity
(v) PRECAUTIONS.........................Avoid eyes, limit prolonged use
(vi) OPERATIONAL STATUS..............Operational
(vii) SIDE EFFECTS......................Local irritation, GI discomfort
(viii) INTERACTIONS.....................N/A
(ix) DOSAGE.................................Per MD/DO
Optional Supplemental Protocol –
TACTICAL EMS (continued)

(j) Emtricitabine and tenofovir (Truvada) (high-risk post-exposure management)
   (i) AVAILABILITY........................................Tablet containing tenofovir DF 300 mg; emtricitabine 200 mg
   (ii) ACTION ...........................................Antiretroviral
   (iii) INDICATIONS ....................................Per MD/DO—infected exposures
   (iv) CONTRAINDICATIONS ......................Known hypersensitivity
   (v) PRECAUTIONS .................................Liver/kidney dx
   (vi) OPERATIONAL STATUS ......................Operational
   (vii) SIDE EFFECTS ..................................GI upset, nausea/vomiting, diarrhea
   (viii) INTERACTIONS ................................N/A
   (ix) DOSAGE ...........................................Per MD/DO

(5) Steroids
   (a) Prednisone (PO)
      (i) AVAILABILITY ..................................PO; 1/5/10/20/50 mg tablets
      (ii) ACTION .........................................Corticosteroid, anti-inflammatory
      (iii) INDICATIONS .................................Allergic reaction, auto-immune condition; per MD/DO
      (iv) CONTRAINDICATIONS ......................Known hypersensitivity
      (v) PRECAUTIONS ..................................PUD/GERD/GI bleed history
      (vi) OPERATIONAL STATUS ......................Operational
      (vii) SIDE EFFECTS ..................................GI upset/nausea
      (viii) INTERACTIONS ..............................N/A
      (ix) DOSAGE .........................................40 mg to 60 mg once daily; per MD/DO

   (b) Dexamethasone (Decadron®) (IV/IM and/or PO)
      (i) AVAILABILITY ..................................PO or IV/IM; tablets
      (ii) ACTION .........................................Corticosteroid, anti-inflammatory
      (iii) INDICATIONS .................................Allergic reaction, auto-immune condition; per MD/DO
      (iv) CONTRAINDICATIONS ......................Known hypersensitivity
      (v) PRECAUTIONS ..................................PUD/GERD/GI bleed history
      (vi) OPERATIONAL STATUS ......................Operational
      (vii) SIDE EFFECTS ..................................GI upset/nausea
      (viii) INTERACTIONS ..............................N/A
      (ix) DOSAGE .........................................10 mg once daily; per MD/DO

(6) Analgesics/Anesthetics
   (a) Acetaminophen (PO)
      (i) AVAILABILITY .................................Tablet: 325 and 500 mg
      (ii) ACTION .........................................Pain medication
      (iii) INDICATIONS .................................Mild to moderate pain
      (iv) CONTRAINDICATIONS ......................Known hypersensitivity, liver disease, PUD/GERD/GI bleed history
      (v) PRECAUTIONS .................................N/A
      (vi) OPERATIONAL STATUS ......................Operational
      (vii) SIDE EFFECTS .................................GI upset
      (viii) INTERACTIONS ..............................N/A
      (ix) DOSAGE .........................................650–1,000 mg / 6 hours
Optional Supplemental Protocol – TACTICAL EMS (continued)

(b) Ibuprofen (Motrin®/Advil®)
(i) AVAILABILITY..................................................200 mg tablet (OTC) and 100 mg/5mL suspension; 600 mg and 800 mg tablets
(ii) ACTION..................................................Non-steroidal anti-inflammatory pain medication
(iii) INDICATIONS.............................Mild to moderate pain
(iv) CONTRAINDICATIONS...............Known hypersensitivity, renal insufficiency (not failure), PUD/GERD/GI bleed history
(v) PRECAUTIONS..........................Do not use with other NSAIDs; caution with concomitant steroid use
(vi) OPERATIONAL STATUS...............Operational
(vii) SIDE EFFECTS..........................GI upset/nausea, GI bleeding risk
(viii) INTERACTIONS..........................N/A
(ix) DOSAGE.............................................400–600 mg / 4–6 hours or 600–800 mg / 6–8 hours

(c) Naproxen (Aleve®/Naprosyn®) (PO)
(i) AVAILABILITY..................................................Tablet: 220/375/500 mg PO tablets
(ii) ACTION..................................................Non-steroidal anti-inflammatory pain medication
(iii) INDICATIONS.............................Mild to moderate pain
(iv) CONTRAINDICATIONS...............Known hypersensitivity, renal insufficiency (not failure), PUD/GERD/GI bleed history
(v) PRECAUTIONS..........................Do not use with other NSAIDs; caution with concomitant steroid use
(vi) OPERATIONAL STATUS...............Operational
(vii) SIDE EFFECTS..........................GI upset/nausea, GI bleeding risk
(viii) INTERACTIONS..........................N/A
(ix) DOSAGE.............................................220–500 mg every 12 hours

(d) Tramadol (Ultram®) (PO)
(i) AVAILABILITY..................................................50 and 100 mg PO tablets
(ii) ACTION..................................................Pain medication
(iii) INDICATIONS.............................Moderate to moderately severe pain
(iv) CONTRAINDICATIONS...............Known hypersensitivity, seizure Disorder, SSRI/TCA/MAOI use, renal or hepatic insufficiency (adjust dose)
(v) PRECAUTIONS..........................Caution with concomitant opioid use
(vi) OPERATIONAL STATUS...............Operational (if no side effects reported)
(vii) SIDE EFFECTS..........................Potential dizziness/nausea
(viii) INTERACTIONS..........................Antidepressants, antipsychotics, Warfarin, Digoxin, Tegretol, Quinidine
(ix) DOSAGE.............................................50–100 mg every 4–6 hours; 400 mg per day maximum
Optional Supplemental Protocol –
TACTICAL EMS (continued)

15.33

(e) Ketamine

*Formulary per General Patient Care Protocols*

(f) Naloxone (Narcan®) (IN and/or IV)

*Formulary per General Patient Care Protocols*

(g) Lidocaine (transdermal for muscular relief, or IM/SQ for stapling as temporizing measure only, alternate dosing regimen)

(i) AVAILABILITY.................................1% (10 mg/mL) ampules/vials

(ii) ACTION........................................Injectable anesthetic

(iii) INDICATIONS.................................Local pain/injury

(iv) CONTRAINDICATIONS......................Known hypersensitivity

(v) PRECAUTIONS.................................Should not exceed 4 mg/kg or 300 mg

(vi) OPERATIONAL STATUS......................Operational

(vii) SIDE EFFECTS...............................With high doses: seizures, lightheadedness, ringing in ears

(viii) INTERACTIONS..............................N/A

(ix) DOSAGE........................................Topical application to site of dental pain

(h) Fentanyl Transmucosal (PO)

(i) AVAILABILITY.................................Lozenge / lollipop 800 mcg

(ii) ACTION.........................................Opioid analgesic

(iii) INDICATIONS.................................Severe pain/injury

(iv) CONTRAINDICATIONS......................Known hypersensitivity

(v) PRECAUTIONS.................................Controlled substance. Patient should not bite or chew the lozenge, but rather allow it to dissolve slowly in the mouth.

(vi) OPERATIONAL STATUS......................NON-OPERATIONAL

(vii) SIDE EFFECTS...............................Patient must be monitored for CNS/ respiratory depression

(viii) INTERACTIONS..............................N/A

(ix) DOSAGE........................................Oral application for patient directed analgesia; patient should remove the lollipop once pain is controlled

(i) Clove oil (for topical dental analgesia)

(i) AVAILABILITY.................................Topical liquid (OTC)

(ii) ACTION.........................................Topical (dental) anesthetic

(iii) INDICATIONS.................................Dental pain/injury

(iv) CONTRAINDICATIONS......................Known hypersensitivity

(v) PRECAUTIONS.................................Penetrating/open intra-oral wounds

(vi) OPERATIONAL STATUS......................Operational

(vii) SIDE EFFECTS...............................N/A

(viii) INTERACTIONS..............................N/A

(ix) DOSAGE........................................Topical application to site of dental pain
(j) **Ketorolac (Toradol®) (injectable)**

(i) **AVAILABILITY**.................................30 mg/mL IV/IM

(ii) **ACTION**........................................Non-steroidal anti-inflammatory pain medication

(iii) **INDICATIONS**..............................Mild to moderate pain

(iv) **CONTRAINDICATIONS**.........................Known hypersensitivity, renal insufficiency (not failure), PUD/GERD/GI bleed history

(v) **PRECAUTIONS**..............................Do not use with other NSAIDs; caution with concomitant steroid use

(vi) **OPERATIONAL STATUS**...................Operational

(vii) **SIDE EFFECTS**.............................GI upset/nausea; GI bleeding risk

(viii) **INTERACTIONS**..............................N/A

(ix) **DOSAGE**.................................15–30 mg IM/IV every 6–8 hours

---

(7) **Sleep/Wake**

(a) **Caffeine (No-Doz®)**

(i) **AVAILABILITY**.................................200 mg tablet

(ii) **ACTION**........................................Enhances alertness

(iii) **INDICATIONS**..............................Suspected caffeine withdrawal headache; to facilitate functioning with limited rest periods

(iv) **CONTRAINDICATIONS**.........................Known hypersensitivity

(v) **PRECAUTIONS**..............................N/A

(vi) **OPERATIONAL STATUS**...................Operational

(vii) **SIDE EFFECTS**.............................Insomnia

(viii) **INTERACTIONS**..............................N/A

(ix) **DOSAGE**.................................200 mg / 3–4 hours as needed

(b) **Zaleplon (Sonata®) (sleeper)**

(i) **AVAILABILITY**.................................10 mg capsule

(ii) **ACTION**........................................Anxiolytic/hypnotic; shortest t-1/2 of agents available

(iii) **INDICATIONS**..............................Facilitate rest during non-operational periods in prolonged deployment/transportation; minimum 4-hour block required for usage (6 hours preferred)

(iv) **CONTRAINDICATIONS**.........................Known hypersensitivity, unsecure location, lack of assured 4-hour non-operational period

(v) **PRECAUTIONS**..............................May not drive/operate machinery/use weapons for minimum 4 hours post-administration

(vi) **OPERATIONAL STATUS**.................NON-OPERATIONAL (x 4 hours after administration)

(vii) **SIDE EFFECTS**..............................Sedation
Optional Supplemental Protocol – TACTICAL EMS (continued)

(viii) INTERACTIONS........................................... Alcohol/other sedatives potentiate effect
(ix) DOSAGE................................................. 10–20 mg with assured 4-hour non-operational block, as approved by MD/DO and Team Commander

(c) Modafinil (Provigil®)
(i) AVAILABILITY........................................... 200 mg tablet
(ii) ACTION.................................................. Enhances alertness/concentration
(iii) INDICATIONS........................................ To facilitate functioning with limited rest periods
(iv) CONTRAINDICATIONS......................... Known hypersensitivity
(v) PRECAUTIONS........................................... N/A
(vi) OPERATIONAL STATUS............................. Operational
(vii) SIDE EFFECTS....................................... Insomnia, mild blood pressure elevation
(viii) INTERACTIONS................................. N/A
(ix) DOSAGE ........................................... 200 mg once daily

(8) Wound Management
(a) Cyanoacrylate tissue adhesive (Dermabond®)
(i) AVAILABILITY........................................... Single use ampoules
(ii) ACTION.................................................. Tissue adhesive
(iii) INDICATIONS........................................... Minor trauma
(iv) CONTRAINDICATIONS......................... Known hypersensitivity
(v) PRECAUTIONS........................................... Avoid near eyes
(vi) OPERATIONAL STATUS............................. Operational
(vii) SIDE EFFECTS....................................... Transient local discomfort
(viii) INTERACTIONS................................. N/A
(ix) DOSAGE ........................................... As required for wound closure, 2–4 layered applications

(b) Topical hemostatic dressing
(i) AVAILABILITY........................................... Individual use packages
(ii) ACTION.................................................. Promotes blood clotting
(iii) INDICATIONS........................................... Hemorrhage
(iv) CONTRAINDICATIONS......................... Known hypersensitivity
(v) PRECAUTIONS........................................... Standard/universal precautions for wound care
(vi) OPERATIONAL STATUS............................ NON-OPERATIONAL
(vii) SIDE EFFECTS....................................... N/A
(viii) INTERACTIONS................................. N/A
(ix) DOSAGE ........................................... Single or multiple dressings applied to bleeding wound

(c) Steri-strips
(i) AVAILABILITY........................................... Individual use packages
(ii) ACTION.................................................. Facilitates closure of wounds
(iii) INDICATIONS........................................... Superficial wounds
(iv) CONTRAINDICATIONS......................... Known hypersensitivity to adhesive
Optional Supplemental Protocol – TACTICAL EMS (continued)

(v) PRECAUTIONS...........................................Standard/universal precautions for wound care
(vi) OPERATIONAL STATUS.............................Operational
(vii) SIDE EFFECTS........................................N/A
(viii) INTERACTIONS........................................N/A
(ix) DOSAGE..................................................Single or multiple dressings applied for wound closure; per MD/DO

(d) Staples
(i) AVAILABILITY.............................................Individual use staple dispensers
(ii) ACTION....................................................Facilitates closure of wounds
(iii) INDICATIONS..........................................Wounds
(iv) CONTRAINDICATIONS...............................Contaminated wounds, wounds with foreign body material
(v) PRECAUTIONS.............................................Standard/universal precautions for wound care
(vi) OPERATIONAL STATUS.............................Operational
(vii) SIDE EFFECTS........................................N/A
(viii) INTERACTIONS........................................N/A
(ix) DOSAGE..................................................Single or multiple dressings applied for wound closure; per MD/DO

(9) ACLS/Resuscitation
(a) Albuterol MDI
(i) AVAILABILITY...........................................0.83 mcg metered dose inhaler
(ii) ACTION....................................................Bronchodilator
(iii) INDICATIONS..........................................Respiratory distress/bronchospasm
(iv) CONTRAINDICATIONS...............................Known hypersensitivity
(v) PRECAUTIONS.............................................Standard/universal precautions for respiratory patient
(vi) OPERATIONAL STATUS.............................NON-OPERATIONAL (without MD/DO consult)
(vii) SIDE EFFECTS........................................N/A
(viii) INTERACTIONS........................................N/A
(ix) DOSAGE..................................................2 puffs, may be repeated two additional times. Additional doses per MD/DO

(10) Anti-hypoglycemics
(a) Oral glucose
Formulary per General Patient Care Protocols
1. PURPOSE
To define the indications for use of a mechanical ventilator by a paramedic for the acute ventilated patient
a) The level of care required for the interfacility transport of the “acute ventilated interfacility patient” is beyond the routine training curriculum for a paramedic; this type of patient must be transported by a higher level health care clinician who is credentialed, educated, and competent in dealing with the ventilator and the ventilated patient. OR
b) When a critical interfacility transfer is needed and a credentialed, educated, and competent higher level health care clinician is genuinely unavailable, a credentialed, educated, and competent paramedic (through a MIEMSS-approved training program) may attend the ventilator and the ventilated patient with the addition of a second ALS clinician or advanced airway trained health care clinician when determined appropriate by the sending/referring physician.

2. INDICATIONS
ACUTE VENTILATED PATIENTS for the interfacility transport are defined as:
a) Intubated OR
b) Tracheostomy patient when the reason for transport is:
   (1) For increased level of care from a hospital, OR
   (2) To continue the same level of care in an acute care setting, OR
   (3) The new tracheostomy patient, within the last 7 days

3. VENTILATOR STANDARDS
a) ACUTE VENTILATOR DEVICE STANDARDS
(1) The ventilator that the service is to use for the acute ventilated patient should be able to match the existing ventilator settings. The following minimum device features (including circuit) must be present for this category of patient:
   (a) Set rate of ventilations
   (b) Adjust delivered Tidal Volume
   (c) Adjustable Pressure Support Settings
   (d) Adjustable Inspiratory and Expiratory ratios (I:E ratio)
   (e) Positive End-Expiratory Pressure (PEEP)
   (f) Peak airway pressure gauge
   (g) Continuous Expiratory Volume measurement (Required)
   (h) Modes
      (i) Assist Control (AC)
      (ii) Synchronized Intermittent Mandatory Ventilation (SIMV)
      (iii) Controlled Mechanical Ventilation (CMV)
   (i) Alarms
      (i) Peak airway pressure
      (ii) Disconnect
(2) Strongly recommended options are:
   Blend percentage oxygen
(3) Must perform periodic maintenance (including calibration) meeting the manufacturer's specifications

b) **ACUTE VENTILATOR USAGE**
   
   (1) A ventilator maintained by the ambulance service or health care facility must be specifically designed for transport use and capable of providing the required settings.
   
   (2) Continuous pulse oximeter and continuous capnography monitoring equipment must be used on all acute ventilated interfacility patients.
   
   (3) Tracheal suctioning kits/catheters must be available.
   
   (4) A tracheostomy replacement tube the same size and one size smaller shall be transported with the patient ventilated through a tracheostomy. (The endotracheal tube equivalent may be substituted.)

4. **POTENTIAL ADVERSE EFFECTS**

   a) Pneumothorax
   
   b) Barotrauma
   
   c) Hypoxemia
   
   d) Hyperventilation
   
   e) Hypoventilation
   
   f) Extubation of endotracheal or tracheostomy tube

5. **PRECAUTIONS**

   If any problems arise with mechanical ventilation, the patient shall be disconnected from the ventilator and manually ventilated.

6. **OPTIONAL PROGRAM REQUIREMENTS**

   a) A special “Ventilated Patient” report form will be completed for each mechanically ventilated patient and will include vital signs, pulse oximeter readings, and lung sounds (recorded a minimum of every 5 minutes), and documentation of any of the following:
      
      (1) cardiac arrest during transport,
      
      (2) dislodgment of tracheostomy tube or endotracheal tube,
      
      (3) equipment failure (with FDA report),
      
      (4) discontinuance of ventilator and conversion to BVM,
      
      (5) deterioration of patient, or
      
      (6) the upgrading of patient care to critical care.

   b) The Optional Program will require a training program that meets or exceeds the “Acute Ventilated Interfacility Patient” curriculum and is approved by the operational program medical director with skills validation. A copy of the training program shall be reviewed and be approved or disapproved by MIEMSS.
Optional Supplemental Protocol –
TRANSPORT OF CHRONIC AND SCENE VENTILATED PATIENTS

1. PURPOSE
To define the indications for use of a mechanical ventilator:

a) Chronic ventilated patient
   The level of care required for the interfacility transport of "chronic ventilated patients" is within the scope of practice of a paramedic who has been credentialed, is competent, and received adequate training specific to the patient’s condition and the equipment necessary to provide care. Exception: A CRT-I or EMT may transport a chronically ventilated patient who is going for routine medical care and has in attendance a patient provided attendant who can manage the patient’s own ventilator.

b) Patient ventilated at the scene of an emergency
   The level of care required for the transport of a ventilated patient from the "scene of an emergency" is within the scope of practice of a paramedic who has been credentialed, is competent, and received adequate training specific to the patient’s condition and the equipment to provide care.

2. INDICATIONS

a) CHRONIC VENTILATED PATIENTS are defined as:
   (1) Tracheostomy is more than 7 days old. If tracheostomy has been in place for 7 days or less, see Transport of Acute Ventilated Interfacility Patients protocol.
   (2) Ventilator settings that have no changes within 24 hours or changes reflecting improvement in the patient; and
   (3) Point of origin or destination is:
      (a) Long-term care facility,
      (b) Home,
      (c) Outpatient setting,
      (d) Hospital; and
   (4) Reason for transport is:
      (a) Return from or transport to a scheduled appointment, or
      (b) For extended care, or
      (c) For emergency treatment (but not complication of airway or respiratory distress); and
   (5) Ventilator settings are:
      (a) Positive End-Expiratory Pressure (PEEP) less than or equal to 10
      (b) Peak pressures less than or equal to 30, and
      (c) No changes in the ventilator settings are required during the transport.

b) SCENE OF AN EMERGENCY – Out-of-Hospital
   (1) Point of origin is at the scene of an out-of-hospital emergency
   (2) A paramedic may utilize mechanical ventilation once the patient is intubated.
   (3) Reason for mechanical ventilation is respiratory arrest or when the patient is intubated and not bucking the ventilator.
   (4) Once the patient is on a ventilator, a second clinician (EMT or higher) is required to assist with patient care.
   (5) Destination – closest appropriate hospital
   (6) Contraindicated in children 8 years of age or less.
3. VENTILATOR STANDARDS
   a) CHRONIC VENTILATOR DEVICE STANDARDS
      (1) The ventilator that the service is to use for the acute or chronically ventilated patient
          should be able to match the existing ventilator settings. The following minimum
          device features (including circuit) must be present for this category of patient:
          (a) Set rate of ventilations
          (b) Adjust delivered Tidal Volume
          (c) Adjustable Pressure Support Settings
          (d) Adjustable Inspiratory and Expiratory ratios (I:E ratio)
          (e) Positive End-Expiratory Pressure (PEEP)
          (f) Peak airway pressure gauge
          (g) Modes
              (i) Assist Control (AC)
              (ii) Synchronized Intermittent Mandatory Ventilation (SIMV)
              (iii) Controlled Mechanical Ventilation (CMV)
          (h) Alarms
              (i) Peak airway pressure
              (ii) Disconnect
      (2) Strongly recommended options are:
          (a) Continuous Expiratory volume measurement
          (b) Blend percentage oxygen
      (3) Must perform periodic maintenance (including calibration) meeting the manufacturer's specifications
   b) CHRONIC VENTILATOR USAGE
      (1) Ventilator used is:
          (a) The patient’s own ventilator intended for home/transport use and have the pa-
              tient, home-care clinician, or staff member from the health care facility manage
              the ventilator, or
          (b) A ventilator maintained by the ambulance service or health care facility specifi-
              cally designed for transport use and capable of providing the required settings.
              If the patient’s ventilator is the same as the company ventilator, the paramedic
              may manage the ventilator without the home-care clinician accompanying pa-
              tient. Exception: A CRT-I or EMT may transport a chronically ventilated patient
              who is going for routine medical care and has in attendance a patient provided
              attendant who can manage the patient’s own ventilator.
      (2) Monitoring equipment must include pulse oximeter (provided by family or service).
      (3) Tracheal suctioning kits/catheters must be available.
      (4) A replacement tracheostomy tube the same size and one size smaller shall be trans-
          ported with the patient ventilated through a tracheostomy. (The endotracheal tube
          equivalent may be substituted.)
Optional Supplemental Protocol –
TRANSPORT OF CHRONIC AND SCENE VENTILATED PATIENTS (continued)

15.35

3) SCENE OF AN EMERGENCY VENTILATOR DEVICE STANDARDS
   Mechanical ventilator used must:
   (1) Be intended for transport use,
   (2) Deliver 100% oxygen, and
   (3) Have minimal parameters to set rate and volume (both adjustable to meet the needs of pediatric and adult patients)

4. POTENTIAL ADVERSE EFFECTS
   a) Pneumothorax
   b) Barotrauma
   c) Hypoxemia
   d) Hyperventilation
   e) Hypoventilation
   f) Extubation of endotracheal or tracheostomy tube

5. PRECAUTIONS
   a) Any acutely ill or injured breathing patient at the “scene of an emergency” requiring assisted ventilation shall be manually ventilated.
   b) If any problems arise with mechanical ventilation, the patient shall be disconnected from the ventilator and manually ventilated.
   c) The Optional Program will require a training program that meets or exceeds the “Chronic and Scene Ventilated Patient” curriculum and be approved by the operational program medical director. A copy of that training program shall be reviewed and be approved or disapproved by MIEMSS.
1. PURPOSE
To define the type of patient an EMS service may transport to a MIEMSS-designated freestanding medical facility.

2. INDICATIONS
A jurisdiction may allow transport of a patient, who meets one or more of the following indications, to a freestanding emergency medical facility.

a) A stable Priority 2, 3, or 4 patient as outlined in The Maryland Medical Protocols for Emergency Medical Services who does not need a time-critical intervention

b) Priority 1 patient with an unsecured airway or in extremis, who requires stabilization beyond the capability of the EMS crew (e.g., cardiac or respiratory arrest)

c) If the freestanding emergency medical facility is a MIEMSS-designated Acute Stroke Ready Facility, patients of all priority that meet stroke criteria may be transported to the Acute Stroke Ready Facility, as long as the transport time to a Primary Stroke or Comprehensive Stroke Center is greater than 15 additional minutes.

3. CONTRAINDICATIONS
Except as provided in Indications, above, the following patients shall not be transported to a freestanding emergency medical facility.

a) Any patient meeting the criteria for transport to a Trauma Center or Specialty Referral Center as defined in The Maryland Medical Protocols for Emergency Medical Services

b) A pregnant patient complaining of abdominal pain or a patient who is in active labor

c) Any patient in need of time-critical intervention that can be provided only at a hospital-based emergency department

4. PROCEDURE
The EMS clinician shall consult with a designated Base Station at the freestanding emergency medical facility, or the nearest Base Station if the freestanding emergency medical facility is not a designated Base Station, prior to arrival on all Priority 1 and 2 transports as provided in Indications and when otherwise unclear of the appropriate destination. The designated Base Station shall direct the clinician to the appropriate destination.

5. SPECIAL CONSIDERATIONS
None
1. **PURPOSE**
The purpose of this protocol is to define the type of patient an EMS service may transport to a MIEMSS-designated freestanding emergency medical facility.

2. **INDICATIONS**
A jurisdiction may allow transport of a patient, who meets one or more of the following indications, to a freestanding emergency medical facility.
   a) A stable Priority 2, 3, or 4 patient as outlined in *The Maryland Medical Protocols for Emergency Medical Services* who does not need a time-critical intervention
   b) Priority 1 patient with an unsecured airway or in extremis, who requires stabilization beyond the capability of the EMS crew (e.g., cardiac or respiratory arrest)

3. **CONTRAINDICATIONS**
Except as provided in INDICATIONS, above, the following patients shall not be transported to a freestanding emergency medical facility.
   a) Any patient meeting the criteria for transport to a Trauma Center or Specialty Referral Center as defined in *The Maryland Medical Protocols for Emergency Medical Services*
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5. **SPECIAL CONSIDERATIONS**
None
## Viral Syndrome Pandemic Triage Protocol

**EFFECTIVE March 17, 2020, until rescinded or superseded**  
For Use by BLS and ALS Clinicians

<table>
<thead>
<tr>
<th>YES</th>
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**ANY CHECKS in a shaded box indicate that patient transport should be encouraged.**

If **ALL CHECKS** are in non-shaded boxes, patient may provide self-care at home.  
Refer to no-transport instructions for patients.

Any patient may be transported at the EMS Clinician’s discretion.

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This emergency protocol was issued by the Maryland Institute for Emergency Medical Services Systems, after approval by the Executive Director and Chairman of the State Emergency Medical Services Board, in response to the COVID-19 pandemic, and in accordance with Education Article Section 13-516(d)(1) and COMAR 30.03.05.02(l) and a catastrophic health emergency proclamation.
Maximize the Use of Limited Resources Alternative Dispatch Protocols

<table>
<thead>
<tr>
<th>Dispatch Priority Level (match vendor or call center based dispatch protocol/tiered algorithm)</th>
<th>Response (Standard Operating Mode)</th>
<th>Level 1(A) Activation of Card 36 and ONLY for use in 6, 10, 18, and 26 DSS1 BELOW IS BACK UP STRATEGY FOR EMD WITHOUT CARD 36</th>
<th>Level 2(B) Implement Declining Response / Configuration CAD Table (Moderate) + Card 36 (6,10,18 &amp; 26) DSS2</th>
<th>Level 3(C) Implement Declining Response / Configuration CAD Table (Severe) + Card 36 (6,10,18 &amp; 26) DSS 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classification 1 (*Echo) Confirmed Cardiac Arrest (Not Breathing Unresponsive per 911 call) (MPD cards - 2, 6, 9, 11,15, 31)</td>
<td>Closest AED Unit and Closest 1st Responder and Closest ALS Ambulance</td>
<td>Closest AED Unit and Closest 1st Responder and Closest BLS Ambulance if available</td>
<td>-Closest AED Unit and -Closest 1st Responder if available</td>
<td>- Closest AED Unit if available - If no unit available, no response</td>
</tr>
<tr>
<td>Classification 2 (*Delta) Life Threatening Emergency/Potentially Life Threatening/Confirmed Unstable Patient(s)</td>
<td>Closest 1st Responder and Closest ALS Ambulance</td>
<td>- Closest 1st Responder and Closest ALS Ambulance if available; - BLS ambulance if ALS unit not available</td>
<td>Closest 1st Responder and Closest Ambulance available (ALS or BLS)</td>
<td>- Closest 1st Responder and - Closest Ambulance if available (ALS or BLS)</td>
</tr>
<tr>
<td>Classification 3 (*Charlie) Non-Critical/Currently Stable Patient(s) Requiring ALS Assessment</td>
<td>Closest ALS Ambulance</td>
<td>Closest Ambulance available (ALS or BLS)</td>
<td>Closest Ambulance available (ALS or BLS)</td>
<td>- Closest 1st Responder if available or - Closest stand-in responder unit</td>
</tr>
<tr>
<td>Classification 4 (*Bravo) BLS Assessment for unknown/possibly dangerous scenes</td>
<td>Closest 1st Responder and Closest BLS Ambulance</td>
<td>Closest 1st Responder and Closest BLS Ambulance if available</td>
<td>Closest 1st Responder</td>
<td>- Trauma Closest 1st Responder - Medical Referral to Nurse or Health Department Advice Phone service if available; or self-transport to Alternate Care Site</td>
</tr>
<tr>
<td>Classification 5 (*Alpha) BLS Treatment</td>
<td>BLS Ambulance</td>
<td>Alternate Care Referral</td>
<td>Alternate Care Referral</td>
<td>Alternate Care Referral</td>
</tr>
<tr>
<td>Classification 6 (*Omega) Non-Ambulance Care</td>
<td>Alternate care such as Poison Control Center; Police/Fire service call, etc.</td>
<td>Alternate care such as Poison Control Center; Police/Fire service call, etc.</td>
<td>Alternate care such as Poison Control Center; Police/Fire service call, etc.</td>
<td>Alternate care such as Poison Control Center; Police/Fire service call, etc.</td>
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A. INTRODUCTION
These protocols are complementary to The Maryland Medical Protocols for Emergency Medical Services. They are to be utilized only under the following conditions:
1. The protocols are being utilized in a defined wilderness environment.
2. The EMS jurisdiction has been authorized to utilize wilderness EMS protocols.
3. The EMS clinician has been credentialed as a wilderness EMS clinician (see B.1.b).
4. The EMS clinician is functioning under appropriate wilderness EMS medical direction.

B. DEFINITIONS
1. Wilderness Environment
   a) A wilderness environment is defined as “any geographic area where the typical urban resources are not adequate for the management of an injured or sick patient.” Some examples include woodland areas, mountainous terrain, uneven terrain where traditional urban EMS equipment and stretchers are not able to safely function, rivers, and ski hills.
   b) In order to be considered a Wilderness EMS (WEMS) clinician, the clinician needs to have completed additional training beyond that required to function in the urban environment. This training can be completed by any of the following methods:
      (1) Completion of the State of Maryland Wilderness EMS Course
      (2) Alternatively, the clinician may demonstrate proficiency in the skills of wilderness EMS after providing proof of completion of a nationally recognized wilderness EMS program. Five programs that are nationally recognized are:
          (a) National Outdoor Leadership School’s Wilderness Medical Institute
          (b) National Ski Patrol’s Outdoor Emergency Care program
          (c) Stonehearth Open Learning Opportunities
          (d) Wilderness Medical Associates
          (e) American Health Safety Institute
      (3) Basic Life Support (BLS) clinicians include both the EMTs and WEMRs who meet these credentialing processes
2. Wilderness EMS Physician
   a) In order to be considered a wilderness EMS physician, the physician needs to have fulfilled the requirements in order to function as a medical director under COMAR 30.03.03 and be recognized by the State EMS Medical Director as being qualified to provide medical direction in the wilderness environment. Expertise in wilderness EMS may be demonstrated by:
      (1) Completion of a recognized program in wilderness medicine
      (2) At least 2 years of experience functioning in the wilderness environment under the defined capacity of a wilderness medical practitioner
3. Wilderness EMS Jurisdiction
   a) In order to be recognized as a wilderness EMS jurisdiction the following parameters must be met:
      (1) A written request with a demonstrated need
      (2) EMS clinicians credentialed as Wilderness Clinicians
      (3) The clinicians are functioning under a state recognized wilderness EMS medical director
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

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b) As there is limited utility for a ground ambulance in the wilderness environment, the wilderness EMS jurisdiction need not be required to have a primary transport vehicle in order to be recognized as a wilderness EMS jurisdiction. However, since the patient will likely eventually need transport to definitive care by ground and/or air ambulance, the wilderness EMS jurisdiction needs to have a plan for transportation once the patient(s) is out of the wilderness environment. Thus, there must be readily available and functioning communication methods between the wilderness EMS jurisdiction and the local EMS jurisdiction. Further, in order to facilitate timely and appropriate post-wilderness care, if the WEMS program is not a section of a previously established public safety EMS transporting jurisdiction, the wilderness EMS jurisdiction must notify the jurisdiction that will be responsible for ground or air transport as soon as the need for transport has been confirmed. Ideally this communication should occur through direct communication with the transporting jurisdiction’s emergency communication center rather than simply dialing 9-1-1.

C. SCOPE OF PRACTICE

1. Provision of medical care in the wilderness environment is unique in that delays of care due to the remoteness of the environment may be detrimental to the patient. In order to address the unique needs and specialized skills required to manage a patient in the wilderness, these protocols and the training required to utilize these protocols will serve to define the scope of practice of the WEMS clinician. Therefore, THE TERM CLINICIAN IS GENERIC AND DOES NOT IMPLY A SPECIFIC LEVEL OF MEDICAL TRAINING. THE WILDERNESS CLINICIAN MAY BE TRAINED TO ANY LEVEL AND COULD BE A PHYSICIAN, PARAMEDIC, CARDIAC RESCUE TECHNICIAN, EMT, OR WILDERNESS EMERGENCY MEDICAL RESPONDER.

2. In order for the EMS clinician to use these wilderness EMS protocols there must be a need demonstrated in which it is documented that without these protocols:
   a) It would not be possible to safely extricate the patient from the environment or
   b) There is a high risk of the patient or other public safety personnel incurring permanent disability or death without the use of the WEMS Protocols

D. TRANSFER OF CARE

1. Care is transferred from the WEMS clinician to the transporting EMS clinician at the point at which the patient is either:
   a) No longer in the wilderness environment, or
   b) The wilderness EMS clinician has formally transferred care to the transporting clinician.

2. There may be times in which the WEMS clinician’s expertise is needed after transfer of care to the transporting jurisdiction. If this is the case:
   a) The highest trained WEMS clinician shall ride to the hospital with the patient.
   b) Conflicts shall be resolved by contacting the medical director for the WEMS jurisdiction and then the local EMS Base Station medical control.
E. DOCUMENTATION/QUALITY IMPROVEMENT

1. At the completion of the rescue, the WEMS clinicians must fill out a patient chart in compliance with the MIEMSS charting system.

2. A brief written report shall be provided to the transporting agency with the following information:
   a) Patient name, age, gender
   b) Pertinent history of the case
   c) Vital signs and other pertinent physical findings
   d) Care rendered

3. WEMS clinicians must demonstrate proficiency to the WEMS Medical Director on an annual basis via skills testing and/or documentation of the utilization of skills in the field. This may be demonstrated through regular field training exercises.

4. Review of each call:
   a) Upon completion of the WEMS event, notification of the utilization of the WEMS Protocols will be made to the appropriate EMS supervisor.
   b) The WEMS Medical Director will review 100% of WEMS calls as soon as is reasonably possible. Ideally this should be done within 48 hours of the event.
   c) The WEMS program will maintain a detailed WEMS database and will provide an annual report to the State EMS Medical Director.

TREATMENT PROTOCOLS

The wilderness EMS clinician shall have responsibilities for part or all of these protocols, summarized as follows, based on BLS or ALS level of certification/licensure:

<table>
<thead>
<tr>
<th>Intervention</th>
<th>BLS</th>
<th>ALS</th>
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<tbody>
<tr>
<td>Provision of access to medications:</td>
<td></td>
<td></td>
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<tr>
<td>Ibuprofen, Acetaminophen, Oral electrolytes, Calcium Carbonate tablets</td>
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<tr>
<td>(e.g. Tums), ranitidine, diphenhydramine, epinephrine, aspirin, albuterol</td>
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<tr>
<td>ondansetron ODT</td>
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<td></td>
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<tr>
<td>Administration of medications in Protocol, not listed above</td>
<td></td>
<td>•</td>
</tr>
<tr>
<td>Hemorrhage control with hemostatic agent and tourniquet</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>King Airway</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>Surgical Cricothyroidotomy</td>
<td></td>
<td>•</td>
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<tr>
<td>(Paramedic only)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wound closure with steri-strips or other tissue tape</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>Wound closure with tissue adhesive</td>
<td></td>
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<tr>
<td>Pelvic Binder</td>
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</tbody>
</table>
Optional Supplemental Protocol –  
WILDERNESS EMS PROTOCOLS (continued)  

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A. Airway  
1. Initiate general patient care as per The Maryland Medical Protocols for Emergency Medical Services. 
2. Assess the patient’s airway and determine if the patient’s airway is patent, intact, or compromised. 
3. If the airway is compromised, establish a patent airway using one of the following techniques: 
   a) Insert an oral-pharyngeal airway or naso-pharyngeal airway. 
   b) Tack the patient’s tongue to the patient’s lip using a safety pin. 
   c) Insert a KING airway per protocol. 
ALS SKILL (PARAMEDIC ONLY)  
   d) If unable to insert a KING airway and unable to keep the airway open with a non-invasive technique, then proceed to a surgical cricothyroidotomy. 

B. Cardiac Arrest  
1. Initiate general patient care as per The Maryland Medical Protocols for Emergency Medical Services. 
2. Perform CPR. 
3. If equipped with AED, utilize as appropriate. 
4. Continue CPR and utilization of AED per protocol until there is Return of Spontaneous Circulation (ROSC). 
5. If an AED is present, the resuscitation may be terminated per the TOR protocol. TOR conditions requiring physician consult are waived, such that clinicians may terminate without consult. 
6. If an AED is not present, the resuscitation may be terminated if there is no ROSC after 30 minutes of resuscitative efforts. 
7. Resuscitation may also be terminated if rescuers are exhausted or in danger. 

C. Asthma  
1. Initiate general patient care as per The Maryland Medical Protocols for Emergency Medical Services. 
2. Administer albuterol MDI – 2 puffs every hour as needed; may administer up to 4 puffs per hour. 
3. Consider administration of epinephrine (manual or auto-injector) for severe asthma. 
4. Pediatrics less than 30 kg estimated weight administer 0.15 mg IM 
5. Pediatrics greater than 30 kg estimated weight and adults administer 0.5 mg IM 
ALS SKILL  
6. Consider administration of dexamethasone 
   (a) Pediatrics – 0.5 mg/kg to max of 10 mg every 24 hours 
   (b) Adults – 10 mg every 24 hours 
All Clinicians  
7. Continue treatment and monitoring of patient. 
8. Transport to definitive care.
D. Acute coronary syndrome
   1. Initiate general patient care as per *The Maryland Medical Protocols for Emergency Medical Services*.
   2. Acute coronary syndrome may be difficult to diagnose in the wilderness environment without the use of a 12-lead EKG. WEMS clinicians should have a high index of suspicion in a patient complaining of chest pain, shortness of breath, or extreme fatigue without an alternate explanation for these symptoms.
   3. Closely monitor vital signs during patient contact.
   4. Provide oxygen if available at 2 liters per nasal cannula or as needed to treat symptoms or keep oxygen saturation above 90% if a pulse oximetry is available.
   5. Administer aspirin 324 mg (81 mg low-dose aspirin X 4) or 325 mg aspirin chewed
   6. Expedite transport out of the wilderness.

E. Shock
   1. Patients presenting with shock will exhibit signs of poor perfusion to critical organs.
   2. The patient may or may not be hypotensive.
   3. The most common reason for shock in trauma is hemorrhage.
   4. Treat the underlying cause. Control external bleeding.
   5. Control for environmental conditions.

ALS SKILL
   6. If carrying IV/IO fluids, establish IV access and administer parenteral fluids with Lactated Ringer's (LR).
   7. Pediatrics 20 mL/kg bolus to maintain a radial pulse and to maintain normal mentation
   8. Adults 500–1,000 mL bolus to maintain a radial pulse and to maintain normal mentation
   9. Continue fluids to maintain peripheral perfusion.

ALL CLINICIANS
   10. Expedite transport.

F. External Bleeding
   1. Initiate general patient care as per *The Maryland Medical Protocols for Emergency Medical Services*.
   2. Control external bleeding with direct pressure.
   3. If unable to control extremity bleeding with direct pressure, apply tourniquet proximally to the site of bleeding. Note the time and date of the tourniquet application. If time of delivery of patient to definitive care is expected to exceed 12 hours, then it is appropriate to release the tourniquet every 2 hours. However if tourniquet is released, closely observe area for bleeding and immediately reapply if bleeding resumes.
   4. If unable to control bleeding in site other than extremity, or if unable to get control of bleeding with a tourniquet, then apply hemostatic impregnated gauze or hemostatic agent (HemCon® or similar product) per manufacturer instructions.

G. Wound Care
   1. Initiate general patient care as per *The Maryland Medical Protocols for Emergency Medical Services*.
   2. Once bleeding has been controlled, assess the size and depth of the wound.
Assess for extent of contamination. In addition, assess for any suspicion of underlying broken bones or dislocated joints in association with the wound.

3. Irrigate the wound. Ideally the wound should be irrigated with high pressure. High pressure irrigation devices can be created with a syringe or a plastic bag with a small hole. Irrigate with water that is clean enough to drink. Irrigate until all visible foreign bodies have been removed.

4. Assess need for primary closure of wound.
   a) In the wilderness setting, large wounds may warrant primary closure if time to definitive treatment is greater than 4 hours.
   b) Primary closure can be achieved with:
      (1) Steri-strips or other tape (duct tape works well)

ALS SKILL
   (2) Tissue adhesive (Dermabond\textsuperscript{\textregistered} or similar product)
   (3) Staples (Physician only skill)
   (4) Sutures (Physician only skill)

 c) Wounds that persist with foreign bodies despite adequate irrigation should not be primarily closed.
 d) Unless there will be a significant delay of transport of patient to definitive care (i.e., greater than 12 hours) do not primarily close facial wounds in the wilderness environment.

5. Assess need for administration of antibiotics
   a) Wounds that warrant antibiotic prophylaxis include:
      (1) Grossly contaminated wounds
      (2) Wounds with obvious involvement of broken bones or joint spaces
      (3) Wounds with involvement of tendons or ligaments
      (4) Mammalian bites
   b) Antibiotic that may be used include:
      (1) Amoxicillin-clavulanate (Augmentin\textsuperscript{\textregistered}) – 10 mg/kg or 500 mg of the amoxicillin component every 8 hours
      (2) Cephalexin (Keflex\textsuperscript{\textregistered}) – 10 mg/kg or 500 mg every 6 hours
      (3) Bactrim\textsuperscript{\textregistered} 5 mg/kg every 12 hours or 1 DS every 12 hours
      (4) Clindamycin 10 mg/kg every 8 hours or 300 mg every 8 hours

ALL CLINICIANS
   6. Cover wound with bacitracin antibiotic ointment.
   7. Cover wound with sterile gauze and gauze wrap.

H. Altered mental status
   1. The differential of altered mental status is quite broad, including:
      a) Traumatic brain injury
      b) Stroke
      c) Infection
      d) Acute coronary syndrome
      e) Intoxication
      f) Hypoglycemia
2. If there is any possibility of trauma, protect the patient’s cervical spine.
3. If unable to check glucose with a glucometer, assume that the patient is hypoglycemic and treat accordingly.
   a) Gently rub oral glucose on the inside of the patient’s cheek, 10–15 grams.

ALS SKILL
   b) If carrying glucagon, administer 1 mg IM (0.5 mg if less than 25 kg).
   c) If carrying IV medications, administer dextrose.
   d) 1 amp D50 IV for adults
   e) 1–2 mL/kg D50 for children greater than 2 years old
   f) 2–4 mL/kg D25 for children less than 2 years old

ALL CLINICIANS
4. Transport out of the wilderness.

I. Traumatic Brain Injury
1. Initiate general patient care as per the MIEMSS protocols.
2. Any patient with a blow to the head and the following findings should prompt the WEMS clinician to initiate rapid transportation to a trauma center:
   a) GCS less than 13 or a motor score less than 6
   b) Rapidly declining GCS
   c) Debilitating headache
   d) Profuse vomiting
   e) Raccoon’s eyes
   f) Battle’s signs
   g) Seizures
3. Control the cervical spine and airway as needed.
4. In a patient with a blow to the head, no loss of consciousness, but at least a brief period of confusion or loss of memory, closely observe and extricate from the wilderness environment. Watch for deterioration of mental status. The patient should be cleared by a physician prior to resuming activities at risk for head injury.

J. Back Injury/Spinal Cord Injury
1. Extrication of a fully immobilized patient from the wilderness environment can be quite difficult and pose increased risks to both the patient and rescuers. Therefore, despite a significant mechanism of injury, patients who have concern for spinal column injury and/or meet criteria for the Spinal Protection Protocol should be allowed to ambulate on their own volition as long as the patient is alert, reliable, and has no major neurological deficits.
2. Patients who have evidence of neurological deficit and/or those who are not able to safely ambulate on their own volition shall be secured in an extrication device in a manner that conforms, as much as possible, to the normal contours of the spine and minimizes, as much as possible, movement of the spinal column.
3. Any patient who has been secured in an extrication device should have placement of a diaper for control of urine, especially if the transport time to definitive care is expected to be greater than one hour.
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

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K. Diagnosis of fractures in the wilderness will be based on clinical findings rather than radiologic studies.

1. Things to assess when considering if a patient has a possible fracture requiring immobilization are:
   a) Ability of the patient to bear weight or use the affected limb
   b) Evidence of angulations, deformities, crepitus, bruising
   c) Did the patient hear a breaking sound or feel the bone breaking?

2. Assess distal neurological as well as vascular function.

3. If the patient does NOT have intact distal pulses, then manually reduce by bringing the affected area back to a near anatomic alignment.

4. The general principle of splinting is to immobilize the joint above and below the site of suspected fracture. Provide adequate padding. Splints may be commercially designed or improvised. Assess pulses before and after splinting. Perform frequent vascular checks during transportation.

5. Consider placing a diaper on the patient to catch urine—especially for fractures of the lower extremities that will prevent the patient from being able to urinate unaided.

6. Specific splinting guidelines are as follows:
   a) Shoulder and upper arm
      (1) Immobilize as needed for comfort.
      (2) Place in a sling and swath.
   b) Lower arm
      (1) Immobilize, including the wrist and elbow.
      (2) Place in sling and swath.
   c) Hand
      (1) Realign misangulated digits as needed.
      (2) Place a soft roll of gauze in the hand.
      (3) Wrap with a bandage.
   d) Hip
      (1) Immobilize both upper legs together, placing padding between the legs.
      (2) Place on a stretcher.
      (3) Carry out.
      (4) Do not place patient in traction.
   e) Pelvis
      (1) Assess for injury to vagina or penis.
      (2) Pelvic fracture is noted by instability of the pelvis.
      (3) Immobilize with commercially available pelvic binder or improvised pelvic binder.
      (4) Expedite transport to a trauma center.
   f) Femur
      (1) Immobilization of femur fractures with traction splints is no more effective than immobilization to the unaffected leg and transport on a stretcher. In the WEMS setting, the clinician should use judgment and either use a traction splint or immobilize the injured leg to the unaffected leg.
      (2) Immobilize the fractured leg to the uninjured leg with adequate padding or use a traction splint.
Optional Supplemental Protocol – WILDERNESS EMS PROTOCOLS (continued)

(3) Place padding behind the knees.
(4) Carry the patient out on a stretcher.

g) Knee
(1) Patellar fractures typically occur due to a direct blow to the patella.
(2) The patient is likely to have significant pain and not want to fully extend the knee.
(3) Immobilize with a circumferential splint ensuring that the popliteal artery behind the knee is not compromised.
(4) The patient may be able to ambulate out on own with a crutch and assistance.

L. Dislocations
1. Considerations for reducing a dislocated joint in the wilderness:
   a) Reductions are typically easier immediately after an injury, before the joint has become swollen and muscles are in spasm.
   b) Extrication of a patient from the wilderness with a dislocated joint can be quite difficult, presenting increased risks to the patient and the rescuers.
   c) Dislocated joints can result in compromise to vascular and/or neurological structures.
2. Always check neurological and vascular integrity before and after an attempted reduction.
3. Consider placing a diaper on the patient for control of urine—especially for dislocations of the lower extremities that may prevent the patient from being able to urinate unaided.
4. Specific reductions are as follows:
   a) Shoulder
      (1) The greater majority of shoulder dislocations are anterior. Mechanism is typically external rotation and abduction. The patient will complain of pain in the shoulder and will be resistant to bringing the arm into a position of rest across the body.
      (2) Check for motor and vascular integrity in the hand.
      (3) Also check for sensation in the outer aspect of the shoulder.
      (4) Reduction technique
         External Rotation
            (a) Lie the patient supine on a flat surface.
            (b) Secure the patient’s affected arm adducted to the patient’s side.
            (c) The elbow should be flexed to 90 degrees.
            (d) Hold the patient’s wrist and gently guide the arm into a slow external rotation while holding the upper arm fixed to the patient’s side.
            (e) Whenever the patient experiences pain, halt the procedure momentarily then continue.
            (f) Continue guiding the forearm until it is lying perpendicular to the patient’s side on the flat surface.
      (5) Place the patient in a sling and swath.
   b) Fingers
      (1) Clinically diagnosed by obvious deformity and loss of function
      (2) Reduction technique
         (a) Maintain digit in partial flexion.
         (b) Apply traction to the flexed digit while pushing the base of the phalanx back into place.
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

(3) Splint the fingers in an anatomic position with a roller gauze splint.

(c) Hip
(1) Hip dislocations tend to be posterior. The patient’s hip will be internally rotated and adducted. You may also notice the affected limb to appear shorter than the other limb.
(2) If equipped with ALS medications, pretreat with midazolam 5 mg IM. Alternatively pre-medicate with an oral analgesic.
(3) Reduction technique
   (a) The patient should be lying supine flat on the ground.
   (b) Flex the hip and knee to 90 degrees.
   (c) Straddle the patient and apply traction in an upward direction while another clinician is providing counter traction by holding the pelvis fixed to the ground.
(4) Once reduced, the hip should be immobilized to the uninjured leg and the patient carried out on a stretcher.

d) Knee
(1) Knee dislocations carry great risk of injury to the popliteal artery behind the knee.
(2) Assess for pulses in the foot.
(3) Reduction technique
   Gently exaggerate the injury and then apply gentle traction to bring the joint to anatomic position.
(4) Splint the knee slightly flexed and carry the patient out.
(5) Expedite transport to a trauma center.

e) Patella
(1) The patella will typically displace laterally with the knee held flexed by the patient for comfort.
(2) Reduction technique
   (a) Gently extend the knee so that the lower leg is straight to the upper leg. This movement may result in the reduction of the dislocated patella.
   (b) If the patella remains dislocated after extension of the knee, then apply gentle pressure on the lateral edge of the patella pushing the patella back into its anatomic location. Do not force the patella if it is not easily reducible.
(3) Splint the leg in extension.
(4) The patient may be able to ambulate with a crutch and assistance.

f) Ankle
(1) Ankle dislocations are typically associated with fractures.
(2) There will be obvious deformity.
(3) There may be compromise of vascular structures.
(4) Reduction technique
   Apply gentle traction to place the ankle back into its anatomic location.
(5) The ankle will likely remain unstable after reduction and may easily dislocate without splinting. Therefore, be prepared to splint the ankle immediately after reduction. Have one clinician maintain the reduction, while another clinician applies a splint.
(6) Carry the patient out of the wilderness.
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

M. Ankle sprain
1. An ankle sprain typically is described by the patient as twisting of the ankle after walking or tripping over a ledge. The patient will often be able to ambulate on the ankle with assistance. There should be no instability to the ankle.
2. Management
   a) Support the ankle with an ACE wrap or other supportive device.
   b) Provide a walking aid for the patient such as a crutch or walking stick.
   c) Assist the patient in ambulating out of the wilderness.

N. Foot Care – Blister management
1. Blisters typically develop from a hiker wearing a shoe that has not been broken in and/or is not fitted properly. Wearing two pairs of socks often helps to prevent blisters.
2. Management
   a) Cover the blister with mole-skin or mole foam.
   b) In most cases you should NOT open the blister, as this increases the risk of infection.
   c) You may open the blister with a scalpel or clean knife if the location of the blister is impeding the ability for the patient to self-extricate from the wilderness. Cut in the lines of the skin, drain the fluid, and then cover with antibiotic ointment and a sterile dressing.
   d) Assist the patient in ambulating out of the wilderness.

O. Eye
1. Non-painful acute loss of vision
   a) Patients with acute non-painful loss of vision may have occlusion of the artery to the eye or vasculitis of the artery.
   b) If available, administer oxygen at high flow.
   ALS SKILL
   c) Administer aspirin 325 mg po (adults only).
   ALL CLINICIANS
   d) Expedite transport to the ophthalmology referral center.
2. Globe rupture
   a) Rupture of the eye globe may be obvious or occult.
   b) Obvious globe rupture will be diagnosed by bleeding from the orbit and irregularly shaped orbit and/or pupil that is not reactive to light.
   c) Cover the affected eye with eye dressing, being careful not to put pressure on the globe, and expedite transport to the ophthalmology referral center.
3. Red Eye
   a) Differential diagnosis of red eye includes:
      (1) Foreign body
      (2) Infection—either bacterial or viral
      (3) Allergic reaction
      (4) Globe rupture
      (5) Acute angle closure glaucoma
   b) Cover eye and expedite transport to ophthalmology referral center.
4. Foreign body in eye
   a) If the clinician is sure that the patient’s discomfort is due to a foreign body, the clinician may attempt to remove the foreign body.

ALS SKILL
   b) Numb the eye with 2 drops tetracaine 0.5% ophthalmic solution (peds and adults).

ALL CLINICIANS
   c) Evert the eyelid.
   d) Remove any foreign particles with a moist cotton applicator or equivalent.
   e) DO NOT FORCEFULLY REMOVE PARTICLES STUCK TO THE EYE.
   f) Irrigate the eye with water clean enough to drink.

P. Nose - Epistaxis
   1. Control bleeding by pinching nose until bleeding stops.
   2. If unable to control bleeding, pack.

ALS SKILL
   3. If you anticipate the packing to be in for greater than 24 hours, initiate antibiotic prophylaxis with either Augmentin® or Bactrim®.

ALL CLINICIANS
   4. Transport out of wilderness.

Q. Teeth
   1. Fractured tooth
      a) A fractured tooth that is bleeding is a dental emergency.
      b) The exposed nerve roots will typically be quite painful.
      c) Place a small piece of aspirin on the top of the exposed nerve roots. This will initially be painful to the patient, but the pain should quickly decrease and then be followed by significant relief of pain. You can also cover the exposed nerve roots with sugarless gum or wax.
      d) Have patient cover tooth with gauze.
      e) Transport out of wilderness.

   2. Tooth avulsion
      a) Pick the tooth up by the top rather than the root.
      b) Irrigate tooth and socket gently with water clean enough to drink.
      c) DO NOT SCRUB THE TOOTH.
      d) Replace tooth in socket and have patient maintain tooth by keeping mouth closed as much as possible. You may fix the tooth in place with a piece of sugarless gum.
      e) Alternatively place tooth inside of cheek ensuring that the patient does not aspirate or swallow the tooth.
      f) If traveling in difficult terrain, it is acceptable to place tooth in container with clear liquid.

R. Burns
   1. Clean burns with water clean enough to drink and gentle scrubbing as needed to remove debris.
2. If you expect to get the patient to a burn center within 24 hours, do not cover with antibiotic ointment. If transport to a burn center is expected to exceed 24 hours, then cover with antibiotic ointment.
3. Cover burn with sterile dressing.

ALS SKILL
4. Treat pain
   a) Ibuprofen 600 mg po every 6 hours; 10 mg/kg
   b) Acetaminophen 3–5 yrs old 160 mg/5mL; 6–9 yrs old 320 mg/10mL; greater than 9 yrs old 640 mg/20mL or 650 mg po tab. May repeat dose every 6 hours as needed.
   c) Oxycodone 5–10 mg every 6 hours as needed
   d) For pediatrics administer 0.1 mg/kg of oxycodone every 6 hours as needed.
   e) Morphine 0.1 mg/kg IV/IM to max dose 20 mg with repeat dose of 0.05 mg/kg to max dose of 10 mg every 1 hour as needed
   f) Administer fentanyl 1 mcg/kg IN/IV/IM to a max dose of 200 mcg with a repeat dose of 1 mcg/kg to a max dose of 200 mcg every 1 hour as needed.

ALL CLINICIANS
5. Transport to burn center if meeting burn center criteria (see Burn protocol in MIEMSS treatment protocols).

S. Anaphylaxis
1. Severe allergic reactions present with diffuse hives, airway swelling, and signs of hypoperfusion.
2. Goals of treatment are to counteract the effects on the airway, respiratory system, and cardiovascular system.
3. Specific treatment
   a) Epinephrine (manual or auto-injector)
      (1) Less than 30 kg estimated weight, administer 0.15 mg IM
      (2) Greater than 30 kg estimated weight and adults, administer 0.5 mg IM
   b) Albuterol MDI 2 puffs may repeat every 5 minutes as needed

ALS SKILL
   c) Benadryl®: Pediatric 1 mg/kg every 6 hours; Adults 25–50 mg every 8 hours
   d) Dexamethasone: Pediatric 0.5 mg/kg; Adults 10 mg po

ALL CLINICIANS
4. Expedite transport out of the wilderness.

T. Hypothermia
1. Hypothermia occurs when the body’s ability to conserve and generate heat is not able to compensate for loss of heat.
2. The conditions that are most favorable for development of hypothermia mirror the most efficient methods for losing heat—wet and windy conditions. Therefore, temperatures just above freezing are often more favorable for the development of hypothermia than temperatures below freezing.
3. The beginning stages of hypothermia are clinically evident when a patient is cold and shivering. During this stage the patient will be able to re-warm themselves with passive warming techniques.
a) Remove the patient from the wet and windy conditions.
b) Remove any wet clothes.
c) Place the patient in sleeping bags or cover the patient with blankets (foil safety blankets work well). Another option is to place the patient’s body into garbage bags, ensuring that the head is not covered with the bag.

4. The point at which the patient is no longer shivering marks the beginning of severe hypothermia. If the patient is not shivering, the patient will not be able to self-generate heat. Also during this stage the patient may develop confusion and other neurological findings. Treatment will need to be active replacement of heat. Follow the steps in #3 above. In addition, add heat to the patient. Possible methods for adding heat include:
   a) Have another person join the patient in a sleeping bag or under blankets.
   b) Pack the patient’s axilla and groin with warm packs or water bottles filled with warm liquids.

5. Profound hypothermia is marked by cardiac instability progressing to arrhythmias—ventricular fibrillation, severe bradycardias, and asystole. Handle the patient carefully so as to not induce ventricular fibrillation, but nevertheless remove the patient from the environment. If suspicious of cardiac arrest, check for a pulse for at least 30 seconds. If the patient is in cardiac arrest, attempt to warm the patient while performing CPR. Continue CPR until the patient is warm, he or she is transferred to the transporting EMS agency, or the rescuers are fatigued.

6. If the patient is alert and there is no concern for airway compromise, feed the patient per the nutrition guidelines. The treatment of hypothermia is aided by the patient having fuel to self-generate heat.

U. Frostbite
1. Frostbite is a localized tissue injury from freezing of tissue. Whereas hypothermia can occur in temperatures above freezing, tissue will not freeze unless temperatures are below freezing.
2. The beginning stages of frostbite are marked by periods of intermittent pain and swelling of the affected tissue. This period is actually called “frostnip” and does not require intervention other than removing the affected tissue from the cold environment.
3. Once the tissue is frostbitten the skin will be pale, cold, and numb. Underlying tissue may be soft and pliable or firm depending on the depth of the freezing.
4. Treatment should only be initiated if the clinician is confident that there is no chance of the affected tissue refreezing. If the tissue is likely to continue to be exposed to a cold environment prior to the patient reaching definitive care, then the affected tissue should, as much as possible, be protected from the environment and covered with warm clothes and/or sterile dressing.
5. If the clinician is reasonably sure the tissue will not be further exposed to the cold, then active treatment may be initiated.
   a) Actively warm the affected tissue in warm water that has been measured with a thermometer to a temperature of 100.4–104 degrees Fahrenheit.
   b) Give ibuprofen 600 mg po every 6 hours for management of the frostbite (Peds dosing 10 mg/kg up to max of 600 mg).
   c) Manage pain as needed—see Pain Management protocol.
ALL CLINICIANS

6. Transport the patient to definitive care.

V. Heat Exhaustion

1. Heat exhaustion is marked by intravascular volume depletion due to dehydration and excessive sweating in a hot environment.
2. Symptoms include dizziness, excessive sweating, headache, confusion, nausea, and weakness.
3. Treatment
   a) Remove the patient from the hot environment and keep in the shade.
   b) Cool the patient by getting the patient wet and fanning.
   c) Replace fluids.
4. Transport out of the wilderness.

W. Heat Stroke

1. Heat stroke is a true environmental emergency marked by injury to the neurological system as a result of excessive heat.
2. The patient may or may not be sweaty.
3. Symptoms include confusion, ataxia, and tachycardia.
4. Skin will be red and hot.
5. Treatment mirrors that for heat exhaustion.
   a) Remove patient from the hot environment and keep in the shade.
   b) Cool patient with water and fanning.
   c) Place ice packs in axilla and groin; if shivering, remove the ice packs.
   d) If the patient is alert, orally replace fluids.

X. Snake Bites

1. There are two wild snakes indigenous to the State of Maryland that are poisonous:
   a) Northern Copperhead – The Northern Copperhead is identified by the coppery color to its head and the alternating tan and dark brown on its body. It likes to hide within woodpiles or under logs.
   b) Timber Rattlesnake – The Timber Rattlesnake is a large, stout-bodied snake that can grow up to 5 feet or more. It is typically identified by bands of dark chevrons on its back. Generally, the snake likes to live in wooded areas, but gravid females may be found sunning on open rocks.
2. Snake bites may or may not present with paired fang puncture wounds. A snake bite may also present with a single puncture wound or just a scratch.
3. The greater majority of bites will present with immediate onset of pain at the site of the bite. The bite will become swollen and erythematous.
4. Mark the site of erythema and monitor its progression.
5. Treatment
   a) Gently clean the area and cover with a sterile dressing.
   b) Do NOT attempt to suck out the venom with a commercial or improvised device.
   c) Do not apply a distal and proximal constricting band for poisonous snakebite to an extremity. Splint the extremity. Remove any jewelry on affected extremity.
   d) As much as possible keep the affected area below the level of the heart.
   e) Unless absolutely necessary, the patient should be carried out rather than
Optional Supplemental Protocol – WILDERNESS EMS PROTOCOLS (continued)

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Optional Supplemental Protocol: Wilderness EMS Protocols

walked out on their own accord.

f) Calmly expedite transport out of the wilderness.

6. Do NOT try to catch the snake for identification purposes.

Y. Tick Bites

1. Tick bites in the State of Maryland are at high risk for transmission of Lyme disease and/or Rocky Mountain spotted fever.
2. In order for a tick to transmit Lyme, the tick has to be attached to the patient for at least 36 hours. Ticks found on a patient that are engorged with blood pose a much higher risk than ticks that are not engorged with blood.
3. Lyme disease presents with a circular red rash with the center clear of redness. Patients will have fevers and non-specific flu-like symptoms. The patient may also have neurological finding such as a facial droop.
4. To remove a tick, directly pull the tick up from the skin using a pair of tweezers or a tick key in a single firm steady pull.

ALS SKILL

5. If there is high suspicion for Lyme, start the patient on antibiotic treatment with doxycycline 100 mg twice a day; 2.2 mg/kg 8 years or greater. If less than 8 years old use Augmentin® 10 mg/kg every 12 hours.
6. If there is suspicion for Rocky Mountain spotted fever (the patient has fever and petechiae), then doxycycline is the antibiotic of choice for all age groups. If less than 45 kg estimated weight, administer 2.2 mg/kg every 12 hours to max dose of 100 mg. If greater than 45 kg then administer 100 mg every 12 hours.

Z. Large Animal Attacks (e.g., bear, wild cat, fox)

1. Ensure that the area is safe and that the animal is not still a threat to the patient or rescuers.
2. Patients typically die from large animal attacks secondary to injury to airway structures or hemorrhagic shock from large, gaping wounds.
3. Ensure the patient has an intact airway.
4. Control for any external bleeding.
5. Clean and dress wounds.
6. Transport out of the wilderness.
7. Do NOT attempt to capture the animal for identification purposes.

AA. Plants

1. Patients may develop localized skin reactions after contact with a plant.
   a) Remove the patient from the plant.
   b) Wash the area clean.

ALS SKILL

   c) For mild reactions, use a topical steroid. Cover the area with betamethasone valerate 0.1% ointment twice a day.
   d) For severe reactions administer dexamethasone 10 mg po; 0.5 mg/kg for pediatrics.
   e) Transport

2. Ingestion of plants and mushrooms can be life-threatening.
   a) Patients will present with nausea and vomiting.
b) Provide supportive care.

c) Transport

BB. Oral Rehydration
1. Oral rehydration with a glucose-sodium solution may be indicated in one of three conditions.
   a) Excessive sweat loss from intense exercise
   b) Mild to moderate heat illness, or severe heat illness as long as the airway is intact and the patient is able to tolerate oral fluids
   c) Dehydration from diarrhea
2. The patient will likely feel dehydrated. Mucus membranes will be dry. Skin may tent.
3. Replacement of fluids with only water and no electrolytes may lead to a dilution of intravascular sodium levels. This risks the development of cerebral edema. Therefore, fluids should be replaced with a solution of glucose and salts.
4. The ideal solution will contain 2–6% glucose and 30 mEq/Liter of sodium. Commercial sports drinks generally contain about 6% glucose and 25 mEq/Liter of sodium. While commercial sports drinks contain more than the ideal amount of glucose and less than the ideal amount of sodium, these solutions are better than just water.
5. If a glucose/sodium solution is not available, hydrate with water judiciously.
6. Replace fluids at a rate of 50–100 mL/kg over the first 4–6 hours.

CC. Nutrition
1. In rescues that are expected to be prolonged (i.e., greater than 4 hours) it may be necessary to provide nutritional support to the patient.
   a) Ensure that the patient has an intact airway and that the patient is not experiencing nausea or vomiting.
   b) Only feed the patient if you are reasonably sure that the patient will not be going to surgery in the next 12 hours.
   c) Provide nutrition with a combination of protein and carbohydrate.
      (1) Energy bars are a good choice.
      (2) A mixture of dried fruits and nuts is also a good choice.

DD. Nausea
1. Patients with traumatic injuries and/or medical illness may experience nausea. All clinicians should refer to the treatment protocols for ODT ondansetron.

ALS SKILL
2. If carrying ALS medications and IVs, follow Nausea and Vomiting Protocol in MIEMSS treatment protocols.
3. Alternatively, may administer
   a) Promethazine pediatric greater than 2 years old 0.5 mg/kg every 12 hours; adults 25 mg po every eight hours
   b) Zofran® pediatric 0.1 mg/kg; adults 4 mg IM

EE. Diarrhea
1. Diarrhea in the wilderness can result in significant dehydration to the patient.
2. Orally rehydrate the patient.
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

ALS SKILL
3. Administer loperamide
4. Pediatric – (loperamide is generally not indicated for pediatric populations. However, in the wilderness it may be needed to prevent profound dehydration or to facilitate extrication. Use judiciously.)
5. 2–6 years of age or 13–20 kg 1 mg po three times a day
6. 6–8 years of age or 20–30 kg 2 mg bid
7. Adults–4 mg po for the first dose then 2 mg po after each subsequent loose stool up to a total of 16 mg in a 24 hour period
8. Contraindications for loperamide are diarrhea with fevers and bloody diarrhea.

FF. Abdominal Pain
1. Non-traumatic abdominal pain may indicate a surgical emergency.
2. In women, a ruptured ectopic pregnancy is a true emergency that may present with abdominal pain.
   a) Check a female patient’s urine for beta hCG using a commercial urine pregnancy test.
   b) If the patient with abdominal pain is pregnant, expedite transport.
3. In non-pregnant females and all males with abdominal pain, monitor vital signs and patient symptoms. Concerning findings suggestive of a surgical abdomen include:
   a) Instability of vital signs
   b) Progressing pain
   c) Rebound pain–pain with movement
   d) Nausea and vomiting
4. If there is high concern for surgical abdomen, do not feed the patient and expedite transport.
5. All other patients with abdominal pain should be transported so as to not miss occult surgical disease.

GG. Gastroesophageal reflux
1. Gastroesophageal reflux (GERD) (or heartburn) is typically identified by the patient complaining of a burning, substernal chest pain. The patient also may complain of having a sour taste.
2. It is important to note that the patient with symptoms of GERD may actually have an acute coronary syndrome. Therefore, as you are treating the patient’s symptoms, also assess for possible acute coronary syndrome and manage appropriately. Relief of symptoms with the recommended treatment for GERD does NOT rule out the possibility of acute coronary syndrome.
3. Management of GERD
   Tums 1–2 chewed every hour as needed to a max dose of 4 tablets

HH. Pain Management
1. Treatment of pain in the wilderness may at times be necessary in order to facilitate extrication and transport out of the wilderness. Therefore, treatment of pain not only benefits the patient by simply decreasing pain, treatment of pain also improves the safety of the patient and rescuers by decreasing the time spent in the wilderness.
2. Mild to moderate pain can be treated with ibuprofen and/or acetaminophen.
   a) Ibuprofen 600 mg every 6 hours orally; 10 mg/kg to max dose 600 mg for pediatric dosing
   b) acetaminophen up to 650 mg every 6 hours orally; 160 mg/5mL for 3–5 years old; 320 mg/10 mL 6–9 years old

ALS SKILL

3. Management of severe pain will often require treatment with an opiate analgesic. While intravenous opiates may have a quicker onset and more easily titratable, oral opiate analgesics tend to have less acute respiratory depression.
   a) If carrying parenteral morphine, administer 0.1 mg/kg IV/IM up to 20 mg IM. May repeat dose of 0.05 mg/kg every hour as needed.
   b) Administer fentanyl 1 mcg/kg IN/IV/IM to a max dose of 200 mcg with a repeat dose of 1 mcg/kg to a max dose of 200 mcg every 1 hour as needed.
   c) Alternatively, administer oxycodone 5–10 mg every 6 hours as needed. Pediatric dosing for oxycodone – 0.1 mg/kg every 6 hours

FORMULARY

acetaminophen (Tylenol®)
- Availability: 325 mg tablet; 160 mg/5 mL
- Action: analgesic; anti-pyretic
- Indication: mild to moderate pain; fever
- Contraindication: known end stage liver disease
- Precautions: N/A
- Side effects: N/A
- Dose: 3–5 years old 160 mg/5 mL every 6 hours as needed
  6–9 years old 320 mg/10 mL every 6 hours as needed
  10 years and above 640 mg/20 mL or 650 mg tab every 6 hours as needed

albuterol
- Availability: 90 mcg/metered spray
- Action: bronchodilator
- Indication: shortness of breath; exacerbation of asthma/COPD; wheezing
- Contraindication: N/A
- Precautions: N/A
- Side effects: N/A
- Dose (Peds & Adult): start with 2 puffs every four hours as needed; may use up to 4 puffs every hour
amoxicillin-clavulanate (Augmentin®)

- Availability: 500 mg–125 mg tablet; 125 mg–31.5 mg/5 mL
- Action: antibiotic
- Indication: suspected respiratory infection
- Contraindication: hypersensitivity to penicillin
- Precautions: N/A
- Side effects: diarrhea
- Dose: Pediatrics – 10 mg/kg every 12 hours
  
  Adult - 1 tablet every 8 hours

Aspirin

- Availability: 325 mg; 81 mg
- Action: anti-platelet
- Indication: suspected acute coronary syndrome or stroke
- Contraindication: hypersensitivity to salicylates
- Precautions: N/A
- Side effects: N/A
- Dose: No pediatric dosing
  
  Adults - one 325 mg tab po qd or four 81 mg tabs po qd

bacitracin

- Availability: 1 ounce (28 gram) ointment tube
- Action: topical antibiotic
- Indication: soft tissue wounds
- Contraindication: N/A
- Precaution: N/A
- Side effects: N/A
- Dose: cover the affected area 2–3 times a day

betamethasone valerate

- Availability: 0.1% topical ointment
- Action: topical steroid anti-inflammatory
- Indication: contact dermatitis
- Contraindication: N/A
- Precautions: N/A
- Side effects: N/A
- Dose: apply to affected area twice a day

calcium carbonate (Tums®)

- Availability: 500 mg; 750 mg chewable
- Action: neutralizes stomach acid
- Indication: upset stomach; gastroesophageal reflux
- Contraindication: N/A
- Precautions: N/A
- Side effects: N/A
- Dose: Pediatric – 1 every four hours as needed
  
  Adult – 1–2 every hour as needed up to max dose of 8 tabs
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

15.39

cephalexin (Keflex®)
• Availability. 500 mg tablets; 125 mg/5mL
• Action. antibiotic
• Indication. suspected skin infection or prophylaxis for skin wound
• Contraindication. hypersensitivity to penicillin
• Precautions. N/A
• Side effects. diarrhea
• Dose. Pediatric – 10 mg/kg every 6 hours
  Adult – 500 mg every 6 hours

chitosan (Hemcon®)
• Availability. 2"X2"; 2"X4"; 4"X4" bandages
• Action. hemostatic
• Indication. severe bleeding
• Contraindication. N/A
• Precautions. N/A
• Side effects. N/A
• Dose (Peds and Adult). apply to severe bleeding as needed

ciprofloxacin (Cipro®)
• Availability. 500 mg tablets
• Action. antibacterial
• Indication. suspected urinary tract infection; skin infection if patient is hypersensitive to penicillin
• Contraindication. hypersensitivity to floroquinolone
• Precautions. N/A
• Side effects. N/A
• Dose. no pediatric dosing
  Adult – 500 mg every 12 hours

clindamycin (Cleocin®)
• Availability. 150 or 300 mg/tablet, reconstituted liquid 75 mg/5 mL
• Action. antibiotic
• Indication. suspected pharyngitis or respiratory infection; cellulitis
• Contraindication. hypersensitivity to clindamycin
• Precautions. N/A
• Side effects. diarrhea
• Dose. Pediatrics – 10mg/kg every 8 hours
  Adult – 300mg every 8 hours

cryanoacrylate tissue adhesive (Dermabond®)
• Availability. single use ampoules
• Action. tissue adhesive
• Indication. minor wound repair
• Contraindication. known hypersensitivity
• Precaution. avoid near eyes
• Side effects. transient local discomfort
• Dose. as required for wound closure; may need 2–4 layers
Optional Supplemental Protocol –
WILDERNESS EMS PROTOCOLS (continued)

**dexamethasone (Decadron®)**
- **Availability.** 1 mg/1 mL solution
- **Action.** Steroidal anti-inflammatory
- **Indication.** Asthma, allergic reactions
- **Contraindication.** N/A
- **Precautions.** N/A
- **Side effects.** N/A
- **Dose.** Adults 10 mg po every 24 hours as needed
  Pediatrics 0.5 mg/kg po every 24 hours as needed

**diphenhydramamine (Benadryl®)**
- **Availability.** 25 mg tablets; 12.5 mg/5 mL
- **Action.** Antihistamine
- **Indication.** Allergic reactions
- **Contraindication.** N/A
- **Precautions.** N/A
- **Side effects.** Sedating
- **Dose.** Pediatric – 1 mg/kg to max dose 50 mg every 8 hours
  Adult – 25–50 mg every 8 hours as needed

**doxycycline (Doxy®)**
- **Availability.** 100 mg tablets; 25 mg/5 mL
- **Action.** Antibacterial
- **Indication.** Suspected respiratory infection with contraindication to Augmentin®
- **Contraindication.** N/A
- **Precautions.** N/A
- **Side effects.** N/A
- **Dose.** 8–14 years old - 2.2 mg/kg every 12 hours
  Adults – 100 mg every 12 hours

**epinephrine auto-injector**
- **Availability.** 0.3 mg; 0.15 mg auto-injector
- **Action.** Antihistamine; anti-inflammatory; vasoconstrictor
- **Indication.** Moderate to severe allergic reaction
- **Contraindication.** N/A
- **Precautions.** N/A
- **Side effects.** Tachycardia; hypertension
- **Dose.** Pediatric less than 30 kg estimated weight – 0.15 mg IM
  greater than 30 kg estimated weight and adults – 0.3 mg IM

* All levels of clinicians shall be authorized to manually draw up epinephrine with a needle and syringe from an ampule or vial after education and credentialing by the wilderness jurisdictional medical director.
fentanyl
- Availability .................. prefilled syringe, multidose vial
- Action ...................... opioid analgesic
- Indication ................... severe pain
- Contraindication .......... N/A
- Precautions ................ N/A
- Side effects .................. depressed level of consciousness; hypoxia; hypotension
- Dose .......................... 1 mcg/kg IN/IV/IM to a max dose of 200 mcg with a repeat dose of 1 mcg/kg to a max dose of 200 mcg every 1 hour as needed

glucagon
- Availability .................. 1 mg injector
- Action ...................... facilitates release of glucose from glycogen stores in the liver
- Indication ................... suspected hypoglycemia in patient that is not able to take oral glucose
- Contraindication .......... N/A
- Precautions ................ N/A
- Side effects .................. N/A
- Dose .......................... Pediatric less than 25 kg – 0.5 mg IM
  greater than 25 mg and adults – 1 mg IM

glucose gel (Glutose 15°)
- Availability .................. 15 grams oral gel
- Action ...................... raises blood glucose levels
- Indication ................... suspected hypoglycemia
- Contraindication .......... N/A
- Precautions ................ N/A
- Side effects .................. N/A
- Dose (Peds and Adult) ...... give to patient by mouth
  in patient with depressed level of consciousness, rub the gel on the patient’s gums, but use caution

hemostatic agent
All levels of clinicians are authorized to use gauze impregnated with hemostatic agent.

ibuprofen (Advil®; Motrin®)
- Availability .................. 200 mg; 400 mg; 600 mg; 40 mg/mL
- Action ...................... anti-inflammatory; analgesic
- Indication ................... mild to moderate pain
- Contraindication .......... hypersensitivity; known renal disease; history of GI bleeding
- Precautions ................ N/A
- Side effects ................ N/A
- Dose .......................... Pediatric – 10 mg/kg to max dose 600 mg every 6 hours as needed
  Adult – 200 mg–600 mg every 6 hours as needed
loperamide (Imodium®)
- Availability: 2 mg tablets
- Action: anti-diarrheal
- Indication: diarrhea
- Contraindication: N/A
- Precautions: N/A
- Side effects: constipation
- Dose: Pediatric – 2 mg after first watery stool, then 1 mg after each subsequent watery stool; max dose 8 mg per day
  Adult – 4 mg after first watery stool; then administer 2 mg after each subsequent watery stool; max dose 16 mg per day

metoclopramide (Reglan®)
- Availability: 10 mg tablets; 5 mg/mL
- Action: anti-emetic
- Indication: nausea and vomiting
- Contraindication: N/A
- Precautions: N/A
- Side effects: N/A
- Dose: Pediatric – 0.1 mg/kg every 8 hours as needed
  Adult – 10 mg every 8 hours as needed

morphine
- Availability: 4 mg carpuject
- Action: opiate analgesic
- Indication: severe pain
- Contraindication: N/A
- Precautions: N/A
- Side effects: depressed level of consciousness; hypoxia; hypotension
- Dose: Pediatric – 0.1 mg/kg IM every hour as needed up to max dose 16 mg per day
  Adult – 4 mg IM every hour as needed up to max dose of 32 mg per day

ondansetron (Zofran®)
- Availability: 4 mg injectable solution
- Action: anti-emetic
- Indication: severe nausea and vomiting
- Contraindication: N/A
- Precautions: N/A
- Side effects: N/A
- Dose: Pediatric – 0.1 mg/kg IM every 1 hour as needed up to max dose 16 mg per day
  Adult – 4 mg IM every 1 hour as needed up to max dose of 32 mg per day
oxycodone
• Availability. 5 mg tablet
• Action. opiate analgesic
• Indication. moderate to severe pain
• Contraindication. N/A
• Precautions. N/A
• Side effects. depressed level of consciousness
• Dose. Pediatric – 0.05–0.15 mg/kg every 6 hours
  Adult – 1–2 tablets by mouth every 4 hours as needed

promethazine (Phenergan®)
• Availability. 25 mg tablets; 6.25/5 mL
• Action. anti-emetic
• Indication. mild to moderate nausea
• Contraindication. N/A
• Precautions. N/A
• Side effects. N/A
• Dose. Pediatric – 0.5 mg/kg every 8 hours as needed
  Adult – 25 mg every 8 hours by mouth as needed

tetracaine
• Availability. 0.5% ophthalmic solution
• Action. topical anesthetic
• Indication. severe eye pain; foreign body removal from the eye
• Contraindication. hypersensitivity
• Precautions. N/A
• Side effects. N/A
• Dose (Peds and Adult). 2 drops to the affected eye

trimethoprim/sulfamethoxazole (Bactrim®)
• Availability. 160 mg TMP/800 mg SMX (DS tab); 40 mg/200 mg/5 mL
• Action. antibiotic
• Indication. sinus infection, upper respiratory infection, urinary tract infection
• Contraindication. hypersensitivity to sulfa
• Precautions. N/A
• Side effects. N/A
• Dose. Pediatric – 5 mg/kg TMP every 12 hours
  Adult – 1 DS tab po bid
1. **PURPOSE**
   This evidence-based decision support tool is designed to assist clinicians in choosing the facility type most likely to deliver definitive care for pediatric patients requiring transport. This represents an ideal destination choice. Destination selection for any individual patient will include other factors, including transport time, unit availability, and patient/family requests.

2. **INDICATIONS**
   Current *Maryland Medical Protocols for Emergency Medical Services* (MMP) should take precedence. The PDTree should be applied to patients considered “pediatric” ages by the MMP. For medical pediatric patients, this is birth up to the 18th birthday. For trauma patients, the PDTree may be used for patients from birth up to the 15th birthday. For this research protocol, both trauma and medical pediatric patients will be called “child.”

3. **CONTRAINDICATIONS**
   a) Pregnant patients
   b) Newly born infants should be transported (with their mother) to the closest appropriate facility able to receive the post-partum mother.

4. **DEFINITIONS**
   a) Pediatric Base Stations currently designated by MIEMSS include Johns Hopkins Hospital Children’s Center and Children’s National Medical Center. These Pediatric Base Stations may be consulted at any time by any Maryland EMS clinician for online medical direction and assistance with destination decision-making.
   b) Specialty or Trauma Center is defined by current MIEMSS facility designations for Trauma, Eye, Burn, and Pediatric Specialty Centers.
   c) Medical Home is defined as the ED/hospital where the patient has their medical records and has established care by specific physicians to address the patient’s unique needs. Existing MMP suggests that EMS clinicians should transport (repatriate) the patient to that hospital as long as that hospital is not more than 15 additional minutes further than nearest hospital (or greater if allowed for by the EMS Operational Program).
   d) Comprehensive Pediatric Center is defined as a hospital ED with pediatric ICU on-site.
   e) Regional Pediatric Care Center is defined as a hospital ED with inpatient pediatric services and/or a designated pediatric ED staffed by pediatric specialty trained physicians 24/7 or a Freestanding Emergency Medical Facility (FEMF) with designated pediatric ED staffed by pediatric specialty trained physicians 24/7.
   f) Nearest Appropriate Facility is defined as the closest hospital ED or FEMF that is available as an EMS transport destination.
   g) Feasibility of transport to the suggested destination type is left to the discretion of the EMS Operational Program.

5. **PEDIATRIC DESTINATION DECISION TREE (See facing page)**

   **ALERT**
   CHILDREN WHO ARE IN CARDIAC ARREST, OR IF A PATENT AIRWAY CANNOT BE ESTABLISHED, MUST BE TRANSPORTED TO THE NEAREST APPROPRIATE HOSPITAL-BASED EMERGENCY DEPARTMENT OR DESIGNATED FREESTANDING EMERGENCY MEDICAL FACILITY.
Research Protocol –
PEdiATRIC DEStINATION DECISION TREE (PDTree) (continued)

PD Tree

Closest ED/FEMF

• Cardiac Arrest
• Unable to Establish a Patent Airway
• Patient in Need of Specialty Care but Prolonged Transport Time

YES

NO

Transport patient to nearest hospital or FEMF; consider consultation with pediatric base station

Consider Specialty or Trauma Center Needs

Specialty Center Criteria
• Cardiac arrest with ROSC
• Stroke patient under age 18
• Eye injury
• Hand injuries meeting criteria
• Burns meeting burn center criteria

Trauma Center Criteria
• Trauma categories A, B, C, D
• Suspected neck injury with paresthesia, weakness, or other neurologic deficits

YES

NO

Transport patient to trauma or specialty center based on protocol; alert trauma team; consider aviation if quicker and of clinical benefit

Consider Need for Transport to Child’s Medical Home

• Does the child have an emergency related to a known condition previously treated at a specific facility?

YES

NO

If feasible, transport patient to their medical home

Consider Need for Comprehensive Care

Medical
• Child ≤ 2 yr Altered Mental Status and no known seizure disorder
• Shock with abnormal Pediatric Assessment Triangle
• DKA/hyperglycemia with nausea/vomiting OR altered mental state
• Respiratory distress in child with technology dependence (CPAP, Bi-PAP, trach)

Trauma (not meeting Trauma Decision Tree)
• Significant soft-tissue injury/ complex wound
• Elbow injury with deformity
• Long bone deformity
• Femur fracture with intact pulse/motor/sensory

YES

NO

If feasible, transport patient to comprehensive pediatric center; consider aviation if faster and of clinical benefit

Consider Need for Regional Pediatric Care

Medical
• ALTE/brief, resolved, unexplained event
• Seizure patient requiring benzodiazepine
• Altered Mental Status, no trauma, no seizure, > 2 yr
• Respiratory distress with hypoxia or serious signs and symptoms
• Sepsis

Trauma (not meeting Trauma Decision Tree)
• Suspected child abuse

YES

NO

If feasible, transport patient to regional pediatric center

Transport per protocol to nearest appropriate facility
1. Indications

Research Protocol for use in Prince George's County only
• Inclusion: Child 0 months-17 years of age with active seizure during EMS care
• Exclusion:
  (1) Prior history of benzodiazepine allergy
  (2) Known or presumed pregnancy
  (3) Severe growth restriction based on paramedic assessment
• If patient meets any exclusion criteria, obtain medical consultation from a Pediatric Base Station.

2. Seizure has stopped

a) Identify and treat injuries.
b) Check blood glucose and treat per Hypoglycemia protocol. If blood glucose is less than 70 mg/dL, administer oral glucose paste (10-15 grams) between the gum and cheek. Administer single additional dose of oral glucose if not improved after 10 minutes.

3. Active seizure

a) Do not restrain the patient or place and device into the patient’s mouth.
b) Protect the patient from injury.
c) Identify and treat potential underlying cause of seizure: epilepsy, head injury, hypoxia, hypoglycemia, hypoperfusion, infection (fever/stiff neck), stroke, alcohol or drug abuse or withdrawal, CVA, head injury.

4. Active seizure

For a patient with ongoing seizure activity or recurrent seizure activity during EMS care:
  a) Administer midazolam with age-appropriate dosing per table below.
b) Do not delay midazolam to place an IV or obtain blood glucose.
### Research Protocol – PEDIATRIC SEIZURE STUDY: PEDIDOSE (continued)

<table>
<thead>
<tr>
<th>Age</th>
<th>First Dose mg</th>
<th>First Dose ml</th>
<th>Route</th>
<th>Second Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-5 months or for patient that does not meet study criteria</td>
<td>0.2 mg/kg OR Max dose 5 mg</td>
<td>NA</td>
<td>IN/IM* *Preferred</td>
<td>**Medical Consultation REQUIRED; Total max dose 5 mg</td>
</tr>
<tr>
<td></td>
<td>0.1 mg/kg Max dose 5 mg</td>
<td></td>
<td>IV/IO Slow IVP over 1-2 minutes</td>
<td></td>
</tr>
</tbody>
</table>

**Age Greater Than or Equal to 6 months**

<table>
<thead>
<tr>
<th>Age</th>
<th>First Dose mg</th>
<th>First Dose ml</th>
<th>Route</th>
<th>Second Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>6-16 months</td>
<td>1.25 mg</td>
<td>0.25 ml</td>
<td>IN/IM</td>
<td>1.25 mg</td>
</tr>
<tr>
<td>17 months-5 years</td>
<td>2.5 mg</td>
<td>0.5 ml</td>
<td>IN/IM</td>
<td>2.5 mg</td>
</tr>
<tr>
<td>6-11 years</td>
<td>5 mg</td>
<td>1 ml</td>
<td>IN/IM</td>
<td>5 mg</td>
</tr>
<tr>
<td>12-13 years</td>
<td>10 mg</td>
<td>2 ml</td>
<td>IN/IM</td>
<td><strong>Medical Consultation REQUIRED</strong></td>
</tr>
<tr>
<td>14-17 years MIEMSS Pediatric Seizure protocol</td>
<td>0.2 mg/kg OR Max dose 5 mg</td>
<td>NA</td>
<td>IN/IM* *Preferred</td>
<td>**Medical Consultation REQUIRED; Total max dose 5 mg</td>
</tr>
<tr>
<td></td>
<td>0.1 mg/kg Max dose 5 mg in 2 mg increments</td>
<td></td>
<td>IV/IO Slow IVP over 1-2 minutes</td>
<td></td>
</tr>
</tbody>
</table>

**18 Years of Age or Older – See MIEMSS Adult Seizure protocol**

If age unknown, use Handtevy app to estimate age:

a) **Airway and vital signs must be closely monitored after midazolam administration.**

b) **Initiate oxygen per protocol, cardiac monitoring, continuous ETCO₂, and pulse oximetry. Maintain all monitoring until transfer of care at the hospital.**

c) **Check blood glucose (after midazolam administered)**

d) **Children who require midazolam administration by EMS should be transported to a pediatric regional level facility or higher. If feasible, transport patient to Children’s National Medical Center or Children’s National at UMC location.**

- See table above regarding need for medical consultation.
- Obtain medical consultation from the Children’s National Medical Center Base Station.