The Maryland Medical Protocols for Emergency Medical Services Providers

Effective July 1, 2016

Maryland Institute for Emergency Medical Services Systems
I. GENERAL INFORMATION

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 providers, 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 providers 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 on-line 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 on-line 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 on-line 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 life-saving patient care is rendered, the EMT/CRT-(I)/paramedic must document the treatment rendered and the reason on-line medical consultation could not be obtained on the Patient Care Report (PCR), the equivalent of the MAIS runsheet, and on an additional narrative. In addition, the “exceptional call” area on the PCR must be marked, and the provider must immediately notify the EMS Jurisdiction. The EMS Jurisdiction must notify the State EMS Medical Director within 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 LETTER AND NUMERICAL OUTLINE FORMAT IS STRICTLY FOR RAPID AND UNIFORM REFERENCE AND DOES NOT IMPLY OR DIRECT A MANDATORY SEQUENCE FOR PATIENT CARE.
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.
D. MARYLAND TRAUMA AND SPECIALTY REFERRAL CENTERS

Trauma Centers

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

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

**Level II Trauma Centers**
- Johns Hopkins Bayview Medical Center, Baltimore
- Prince George’s Hospital Center, Cheverly
- Sinai Hospital of Baltimore
- Suburban Hospital (JHM), Bethesda

**Level III Trauma Centers**
- Meritus Medical Center, Hagerstown
- Peninsula Regional Medical Center, Salisbury
- Western Maryland Regional Medical Center, Cumberland

**Out-of-State Centers**
- Christiana Care Health System, Wilmington, DE
- MedStar Washington Hospital Center, Washington, DC

Specialty Referral Centers

**Eye Trauma**
- Wilmer Eye Institute at The Johns Hopkins Hospital, 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**
- Pediatric Trauma Center at The Johns Hopkins Children’s Center, Baltimore
- Pediatric Trauma Center at Children’s National Health System, Washington, DC

**Burns**
- Baltimore Regional Burn Center at Johns Hopkins Bayview Medical Center, Baltimore
- Burn Center at MedStar Washington Hospital Center, Washington, DC
- Pediatric Burn Center at The Johns Hopkins Children’s Center, Baltimore
- Pediatric Burn Center at Children’s National Health System, Washington, DC
Specialty Referral Centers

Perinatal Referral Centers

- Anne Arundel Medical Center, Annapolis
- Franklin Square Medical Center (MedStar), Baltimore
- Frederick Memorial Hospital, Frederick
- Greater Baltimore Medical Center, Towson
- Holy Cross Hospital, Silver Spring
- Howard County General Hospital (JHM), Columbia
- Johns Hopkins Bayview Medical Center, Baltimore
- Mercy Medical Center, Baltimore
- Prince George’s Hospital Center, Cheverly
- Saint Agnes Hospital, Baltimore
- Saint Joseph Medical Center (UM), Baltimore
- Shady Grove Adventist Hospital, Gaithersburg
- Sinai Hospital of Baltimore
- The Johns Hopkins Hospital, Baltimore
- University of Maryland Medical Center, Baltimore

Primary Stroke

- Anne Arundel Medical Center, Annapolis
- Atlantic General Hospital, Berlin
- Baltimore Washington Medical Center (UM), Glen Burnie
- Calvert Memorial Hospital, Prince Frederick
- Carroll Hospital Center, Westminster
- Charles Regional Medical Center (UM), La Plata
- Franklin Square Medical Center (MedStar), Baltimore
- Frederick Memorial Hospital, Frederick
- Good Samaritan Hospital (MedStar), Baltimore
- Greater Baltimore Medical Center, Baltimore
- Harbor Hospital (MedStar), Baltimore
- Harford Memorial Hospital (UMUCH), Havre De Grace
- Holy Cross Hospital, Silver Spring
- Howard County General Hospital (UMUCH), Columbia
- Johns Hopkins Bayview Medical Center, Baltimore
- Mercy Medical Center, Baltimore
- Meritus Medical Center, Hagerstown
- Midtown Campus (UM), Baltimore
- Montgomery Medical Center (MedStar), Olney
- Northwest Hospital, Baltimore
- Peninsula Regional Medical Center, Salisbury
- Saint Agnes Hospital, Baltimore
- Saint Joseph Medical Center (UM), Baltimore
- Saint Mary’s Hospital (MedStar), Leonardtown
- Shady Grove Adventist Hospital, Rockville
- Shore Medical Center at Easton (UMSRH)
MARYLAND TRAUMA AND SPECIALTY REFERRAL CENTERS (Continued)

Primary Stroke (Continued)
- Sinai Hospital of Baltimore
- Southern Maryland Hospital (MedStar), Clinton
- Suburban Hospital (JHM), Bethesda
- Union Hospital of Cecil County, Elkton
- Union Memorial Hospital (MedStar), Baltimore
- Upper Chesapeake Medical Center (UMUCH), Bel Air
- Washington Adventist Hospital, Takoma Park
- Western Maryland Regional Medical Center, Cumberland

Comprehensive Stroke
- The Johns Hopkins Hospital, Baltimore
- University of Maryland Medical Center, Baltimore

Cardiac Interventional
- Anne Arundel Medical Center, Annapolis
- Baltimore Washington Medical Center (UM), Glen Burnie
- Bayhealth Kent General, Dover, DE
- Carroll Hospital Center, Westminster
- Christiana Care Health System, Newark, DE
- Franklin Square Medical Center (MedStar), Baltimore
- Frederick Memorial Hospital, Frederick
- Holy Cross Hospital, Silver Spring
- Howard County General Hospital (JHM), Columbia
- Johns Hopkins Bayview Medical Center, Baltimore
- MedStar Washington Hospital Center, Washington, DC
- Meritus Medical Center, Hagerstown
- Nanticoke Memorial Hospital, Seaford, DE
- Peninsula Regional Medical Center, Salisbury
- Prince George’s Hospital Center, Cheverly
- Saint Agnes Hospital, Baltimore
- Saint Joseph Medical Center (UM), Baltimore
- Shady Grove Adventist Hospital, Rockville
- Sinai Hospital of Baltimore
- Southern Maryland Hospital (MedStar), Clinton
- Suburban Hospital (JHM), Bethesda
- The Johns Hopkins Hospital, Baltimore
- Union Memorial Hospital (MedStar), Baltimore
- University of Maryland Medical Center, Baltimore
- Upper Chesapeake Medical Center (UMUCH), Bel Air
- Washington Adventist Hospital, Takoma Park
- Western Maryland Regional Medical Center, Cumberland
E. PROTOCOL KEY

1. Basic Life Support Level Care

2. Advanced Life Support Level Care

3. Requires Medical Consultation

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

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

Response

Scene Arrival + Size Up

Personal Protective Equipment

Patient Approach

Initial Assessment

History + Physical Exam

Withhold Resuscitation

YES

Pronouncement of Death

NO

Palliative Care Protocol

OPTION A/B

YES

DNR/MOLST

NO

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

Termination of Resuscitation Efforts

Procedures

Pharmacology

Inability to Carry Out Physician’s Orders

Extraordinary Care

LEGEND

General Patient Care Section

Refer to Specific Protocols
G. 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 provider 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 Compliance Office, 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’ Compliance Office and State EMS Medical Director the written results of the Medical Review Committee QA investigation and recommendations.
H. INABILITY TO CARRY OUT PHYSICIAN ORDER

Occasionally a situation may arise in which a physician’s order cannot be carried out; e.g., the provider 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 provider 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 Administrator 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 MIEMSS’ Compliance Office 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.
I. PHYSICIAN ORDERS FOR EXTRAORDINARY CARE NOT COVERED BY MARYLAND PROTOCOL

Rarely, a physician providing on-line medical consultation may direct a prehospital provider to render care that is truly life-saving and is not explicitly listed within the protocols.

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

   a) During the consultation, both the consulting physician and the provider must acknowledge and 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 provider must feel capable of correctly performing the care directed by the consulting physician, based on the instructions given by the consulting physician.

   c) When such an order is carried out, the consulting physician and the provider must immediately notify the State EMS Medical Director (via SYSCOM, 800-648-3001) of the extraordinary care situation. In addition, the provider must fax documentation of the rationale for extraordinary care within **24 hours** to the State EMS Medical Director at 410-706-0853. Attendance at a subsequent review meeting shall be required.

   d) The prehospital provider 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). The prehospital provider must also notify his/her BLS/ALS Program Medical Director within **24 hours**.

   e) The public service local EMS jurisdiction and the Program Medical Director must then submit written notification of the incident to the Regional Medical Director with a copy to the State EMS Medical Director within **5 days** of the incident.

   f) The commercial ambulance company and the Program Medical Director must submit written notification of the incident to the Director of the State Office of Commercial Ambulance Licensing and Regulation and the State EMS Medical Director within **5 days** of the incident.
II. GENERAL PATIENT CARE (GPC)

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

   CORRECT LIFE-THREATENING PROBLEMS AS IDENTIFIED.
   STABILIZE CERVICAL SPINE WHEN APPROPRIATE.

   FOR PEDIATRIC PATIENTS, CONSIDER USING THE PEDIATRIC ASSESSMENT TRIANGLE.

1. Assess mental status
   a) Alert
   b) Responds to Verbal stimuli
   c) Responds to Painful stimuli
   d) Unresponsive

2. Airway
   a) Open and establish airway using appropriate adjunct.
   b) Place patient in appropriate position.
   c) Suction airway as needed, including tracheostomy tubes.
IF A PATENT AIRWAY CANNOT BE ESTABLISHED, THE PATIENT MUST BE TRANSPORTED TO THE NEAREST APPROPRIATE HOSPITAL-BASED EMERGENCY DEPARTMENT OR DESIGNATED FREESTANDING MEDICAL FACILITY. ONCE THE PATIENT PRESENTS TO THE HOSPITAL OR DESIGNATED FREESTANDING MEDICAL FACILITY FOR TREATMENT OF AN EMERGENCY CONDITION, TREATMENT AND TRANSFER DECISIONS ARE THE RESPONSIBILITY OF THE HOSPITAL UNDER APPLICABLE LAW. THE PROVIDER SHOULD STAND BY TO BE AVAILABLE FOR AND ASSIST WITH TRANSFER OF THE PATIENT IF THE HOSPITAL DETERMINES SUCH A TRANSFER IS APPROPRIATE.

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 CAUTIOUSLY AND HAVE FOREIGN BODY AIRWAY REMOVAL EQUIPMENT READY FOR IMMEDIATE USE IN CASE THE PATIENT’S AIRWAY BECOMES OBSTRUCTED.

3. Breathing
   a) Determine if breathing is adequate. Assess oxygen saturation (SpO₂) with portable pulse oximeter (required on all transport units since 2012).
      (1) If patient’s ventilations are not adequate, provide assistance with 100% oxygen using Bag-Valve-Mask (BVM).
         (i) For all ages except neonates, 1 breath every 5 seconds (8–12 breaths/min) (manually-activated positive pressure oxygen delivery device is not recommended for this group)
         (ii) For a neonate, 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) SpO₂ greater than or equal to 94% is considered normoxia in adults and children. 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 nonre-breather (NRB).
            Note: Respiratory distress is present if the patient has retractions, nasal flaring, wheezing, stridor, or difficulty speaking.
         (ii) 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
         (iii) CO exposure: Apply 100% oxygen via NRB mask. Maintain SpO₂ at 100%.
(3) If available, utilize EtCO₂ waveform monitoring in intubated patients (required on all ALS transport units for advanced airway management since 2015).

(4) Consider carbon monoxide measurement, if 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 (NEW ‘16)
   - Adult (including adolescent greater than 13 years): 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

(2) If hyperventilating, use EtCO₂ monitoring if available.

NEVER WITHHOLD OXYGEN FROM A PATIENT IN RESPIRATORY DISTRESS!

<table>
<thead>
<tr>
<th>DEVICE</th>
<th>FLOW RATE</th>
<th>CONCENTRATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal Cannula</td>
<td>2–6 lpm</td>
<td>24–44%</td>
</tr>
<tr>
<td>Venturi Mask</td>
<td>Variable</td>
<td>24–60%</td>
</tr>
<tr>
<td>Partial Rebreather Mask</td>
<td>6–10 lpm</td>
<td>35–60%</td>
</tr>
<tr>
<td>Simple Face Mask</td>
<td>6–10 lpm</td>
<td>35–60%</td>
</tr>
<tr>
<td>Pocket Mask</td>
<td>12–15 lpm</td>
<td>50–60%</td>
</tr>
<tr>
<td>Non-Rebreather Mask</td>
<td>12–15 lpm</td>
<td>80–100%</td>
</tr>
<tr>
<td>Bag-Valve-Mask</td>
<td>12–15 lpm</td>
<td>90–100%</td>
</tr>
</tbody>
</table>
4. Circulation

ONCE CONFIRMED PULSELESS, HIGH-QUALITY CONTINUOUS CPR WITH FREQUENT PROVIDER ROTATION IS AN ESSENTIAL COMPONENT IN THE SUCCESSFUL RESUSCITATION OF THE ARRESTED PATIENT. THIS MAY BE ACCOMPLISHED THROUGH MANUAL OR MECHANICAL MEANS AS APPROPRIATE.

PERFORM CPR WHILE PREPARING FOR RHYTHM ANALYSIS AND DEFIBRILLATION.

a) Assess pulse.

(1) Patients from one hour after birth (newly born) up to those who have not reached their 13th birthday (NEW ’16):

(a) If pulse is absent, use AED/manual defibrillator or begin CPR.

(b) If patient is symptomatic with poor perfusion (unresponsive or responds only to painful stimuli) and pulse is less than 60 bpm:

(i) Ventilate for 30 seconds.

(ii) If after 30 seconds the pulse is less than 60 bpm, begin CPR.

(c) If pulse greater than 60 bpm, continue assessment.

(2) Patients 13 years of age or older:

(a) If pulse is absent, use AED/manual defibrillator or begin CPR.

(b) If pulse is present, continue assessment.

b) Assess for and manage profuse bleeding.

c) Assess skin color, temperature, and capillary refill.

5. Disability

a) Perform Mini-Neurologic Assessment (Pulse/Motor/Sensory).

b) Spinal protection

(1) The provider shall determine the appropriate method to use in spinal protection of the patient. Infant or child car seats may NOT be used as a spinal immobilization device for the pediatric patient.

(2) 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 protection.

(a) Midline spinal pain, tenderness, or deformity

(b) Signs and symptoms of new paraplegia or quadriplegia

(c) Focal neurological deficit

(d) Altered mental status or disorientation

(e) Distracting injury

In addition to the above indicators for adults, the below apply to children who have not yet reached their 15th birthday.

(f) Neck pain or torticollis

(g) 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)

(h) Substantial torso injury

(i) Conditions predisposing to spine injury

(3) If NO to all of the above, transport as appropriate.

IF PATIENT IS UNABLE TO COMMUNICATE OR APPROPRIATELY RESPOND TO THE ABOVE QUESTIONS, APPLY SPINAL PROTECTION PROTOCOL.
6. Exposure
To assess patient’s injuries, remove clothing as necessary, considering condition and environment.

7. 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.

8. 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>
</tr>
<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>
</tr>
<tr>
<td>1 yr.</td>
<td>10 kg</td>
<td>120</td>
<td>26</td>
<td>90</td>
</tr>
<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>
</tr>
<tr>
<td>4 yrs.</td>
<td>17 kg</td>
<td>100</td>
<td>24</td>
<td>90</td>
</tr>
<tr>
<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>
</tr>
<tr>
<td>10 yrs.</td>
<td>35 kg</td>
<td>85</td>
<td>20</td>
<td>100</td>
</tr>
<tr>
<td>12 yrs.</td>
<td>40 kg</td>
<td>85</td>
<td>20</td>
<td>100</td>
</tr>
<tr>
<td>14 yrs.</td>
<td>50 kg</td>
<td>80</td>
<td>18</td>
<td>110</td>
</tr>
<tr>
<td>ADULT</td>
<td>Greater than 50 kg</td>
<td>80</td>
<td>18</td>
<td>120</td>
</tr>
</tbody>
</table>
# HISTORY AND PHYSICAL EXAMINATION

## TRAUMA PATIENT

<table>
<thead>
<tr>
<th>Significant MOI</th>
<th>Non-Significant MOI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Trauma Assessment</td>
<td>Determine Chief Complaint</td>
</tr>
<tr>
<td>Head</td>
<td>D</td>
</tr>
<tr>
<td>Crepitation</td>
<td>D</td>
</tr>
<tr>
<td>Chest</td>
<td>C</td>
</tr>
<tr>
<td>Crepitation</td>
<td>C</td>
</tr>
<tr>
<td>Respiration</td>
<td>A</td>
</tr>
<tr>
<td>Paradoxical Motion</td>
<td>A</td>
</tr>
<tr>
<td>Breath Sounds</td>
<td>P</td>
</tr>
<tr>
<td>Abdomen</td>
<td>B</td>
</tr>
<tr>
<td>Rigidity</td>
<td>B</td>
</tr>
<tr>
<td>Distention</td>
<td>T</td>
</tr>
<tr>
<td>Pelvis/GU</td>
<td>L</td>
</tr>
<tr>
<td>Pain on Motion</td>
<td>S</td>
</tr>
<tr>
<td>Blood, Urine, Feces</td>
<td>Extremities</td>
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<tr>
<td>Extremities</td>
<td></td>
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<tr>
<td>Posterior</td>
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## MEDICAL PATIENT

<table>
<thead>
<tr>
<th>Unresponsive Patient</th>
<th>Responsive Patient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Physical Examination</td>
<td>Obtain History of Episode</td>
</tr>
<tr>
<td></td>
<td>Onset</td>
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<tr>
<td></td>
<td>Provocation</td>
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<td>Quality</td>
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<td>Radiation</td>
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<td></td>
<td>Severity</td>
</tr>
<tr>
<td></td>
<td>Time</td>
</tr>
</tbody>
</table>

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

| | Baseline Vital Signs |
| | Obtain History of Episode |
| | Onset |
| | Provocation |
| | Quality |
| | Radiation |
| | Severity |
| | Time |

| | Obtain SAMPLE History |
| | Baseline Vital Signs |
| | Obtain History of Episode |
| | Onset |
| | Provocation |
| | Quality |
| | Radiation |
| | Severity |
| | Time |

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

| | Baseline Vital Signs |
| | Obtain History of Episode |
| | Onset |
| | Provocation |
| | Quality |
| | Radiation |
| | Severity |
| | Time |

| | Obtain SAMPLE History |
| | Baseline Vital Signs |
| | Obtain History of Episode |
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| | Pertinent History |
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| | Medications |
| | Pertinent History |
| | Last Oral Intake |
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| | Baseline Vital Signs |
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| | Radiation |
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| | Baseline Vital Signs |
| | Obtain SAMPLE History |
| | Signs & Symptoms |
| | Allergies |
| | Medications |
| | Pertinent History |
| | Last Oral Intake |
| | Events Prior |

## CONSIDER ALS, PERFORM INTERVENTIONS, AND TRANSPORT.
## Detailed and Ongoing Assessments

### Detailed Examination

| Head     | Scalp & Cranium Crepitation
|          | Eyes
|          | Discoloration Equality Foreign Bodies Blood in Anterior Chamber
| Ears & Nose | Fluid Drainage or Bleeding Discoloration
| Mouth    | Teeth & Foreign Bodies Swelling or Lacerations Breath Odor Discoloration
| Neck     | Jugular Vein Distention Trachea Position Crepitation
| Chest    | Paradoxical Motion Breath Sounds Crepitation
| Abdomen  | Rigidity Distention
| Pelvis/Gu | Pain on Motion
| Extremities | Pulse, Motor, Sensory Capillary Refill
| Posterior|  |

### Ongoing Assessment

#### Medical Patient

- **Repeat Initial Assessment**: Reassess AVPU, Reassess Airway, Monitor Breathing, Reassess Circulation, Monitor Skin, Confirm Clinical Priority

- **Repeat & Record Vital Signs**

- **Repeat Focused Assessment**

- **Check All Interventions**
  - Assure Oxygen Adequacy
  - Check Bleeding
  - Check Interventions
  - Check for Trending
  - Stable Pt.- Every 15 Min.
  - Unstable Pt.- Recommend Every 5 Min.

#### Trauma Patient

- **Repeat Initial Assessment**: Reassess AVPU, Reassess Airway, Monitor Breathing, Reassess Circulation, Monitor Skin, Confirm Clinical Priority

- **Repeat & Record Vital Signs**

- **Repeat Rapid Trauma Assessment**

- **Check All Interventions**
  - Assure Oxygen Adequacy
  - Check Bleeding
  - Check Neck Stabilization
  - Check Interventions
  - Check for Trending
  - Stable Pt.- Every 15 Min.
  - Unstable Pt.- Recommend Every 5 Min.

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**Consider ALS, Perform Interventions, and Transport.**
Combined START/JumpSTART Triage Algorithm

Able to walk?
  YES → MINOR → SECONDARY TRIAGE*
  NO → Breathing?
    NO → POSITION UPPER AIRWAY → BREATHING → IMMEDIATE
    YES → PEDI
      + PULSE → IMMEDIATE
      NO PULSE → ADULT
        APNEIC → DECEASED
    YES → 5 RESCUE BREATHS → APNEIC → DECEASED
          → BREATHING → IMMEDIATE

Respiratory Rate
  >30 ADULT → IMMEDIATE
  <30 ADULT
    15-45 PEDI → IMMEDIATE

Perfusion
  NO PALPABLE PULSE (PEDI) → IMMEDIATE
  YES → CR > 2 sec (ADULT) → IMMEDIATE
    T" INAPPROPRIATE POSTURING OR "I" (PEDIATRIC)
    DOESN'T OBEY COMMANDS (ADULT) → IMMEDIATE
    OBEYS COMMANDS (ADULT) → DELAYED

* Using the JS algorithm, evaluate first all children who did not walk under their own power.

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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. Obtain an EKG when appropriate.

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. An assessment must be conducted to determine if the patient meets Excited Delirium Syndrome. (NEW ‘16)

3. Providers may assist the patient or primary caregiver in administering the patient’s prescribed rescue medication.
   a) BLS providers may assist with the administration of the patient’s fast-acting bronchodilator MDI and sublingual nitroglycerin.
   b) ALS providers 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) Providers should obtain on-line medical direction to administer other prescribed rescue medications not specifically mentioned in The Maryland Medical Protocols for EMS Providers (e.g., 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.

   DO NOT ADMINISTER ORAL MEDICATIONS (EXCEPT GLUCOSE PASTE) TO PATIENTS WITH AN ALTERED MENTAL STATUS.

4. 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. (NEW ’16)
   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. (NEW ’16)
   d) Infants and children must be properly restrained prior to and during transport.
   e) When appropriate, family members should remain with pediatric patients.
G. COMMUNICATIONS (NEW ’16)

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 providers, hospitals, public safety communications centers, and EMRC/SYSCOM are being recorded.

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

ANY PATIENT WHO WHOM THE PROVIDER IDENTIFIES AS MEETING ANY “SPECIALTY” ALERT (E.G., TRAUMA, STEMI ALERT, STROKE ALERT, SEPSIS ALERT) REQUIRES AN ON-LINE MEDICAL CONSULTATION THROUGH EMRC ON A RECORDED LINE (RADIO OR PHONE).

3. All Priority 2 patients who have persistent symptoms or need further therapeutic intervention(s) require on-line medical consultation through EMRC on a recorded line (radio or phone).

4. Notification (“information only call” that can be through EOC or EMS communication system following local standard operating procedures) should be made to the receiving hospital for Priority 2 or Priority 3 patients whose symptoms have resolved and whose vital signs are within normal limits.

ON-LINE MEDICAL CONSULTATION MAY BE OBTAINED AT ANY TIME FOR ANY PATIENT, IF DESIRED BY THE PREHOSPITAL EMS PROVIDER. 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.

5. If medical consultation is genuinely unavailable, or if the time necessary to initiate consultation significantly compromises patient care, the provider 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).

6. Core essentials for communications:
   a) Assigned patient priority (1 to 4)
   b) Age
   c) Chief complaint
   d) Provider 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 physician findings
   g) ETA

In addition, for specialty center patients:

Trauma
h) Patient Trauma Decision Tree Category (Alpha, Bravo, Charlie, Delta)
i) Number of victims if more than one
j) Describe mechanism

Stroke
k) Last known well time
l) Specific neurological findings (sensory, motor, cognitive)
m) 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
n) 12-Lead interpretation
o) 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.
7. 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 providers 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 Communi-
      cator who shall establish appropriate communications.
   c) Reference the Multiple Casualty Incident/Unusual Incident 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 provider’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 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.
   c) Stable Priority 3 or 4 patients who do not need a time-critical intervention may
      also be transported to the local emergency department or freestanding 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 sus-
      pected 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. (NEW ’16)
   e) For Priority 2 and Priority 3 patients not meeting a specialty center destination
      care protocol, the EMS provider should ask if the patient has had a hospital ad-
      mission (inpatient service) within the last 30 days. If the answer is yes, the EMS
      provider 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). (NEW ’16)
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 not be a 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 Protection 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 provider-patient relationship is established when the ALS provider initiates patient assessment and
1. ALS medication(s)* is/are administered or
2. ALS procedure(s)* is/are performed or
3. Upon ALS provider assessment of the patient there is potential risk of deterioration.

* Based on the medication or procedure as listed in the protocol pages 182–185

ALS providers may only terminate their EMS provider-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 provider with a lower scope of practice.

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

Providers will relay assessment findings and treatment provided to the individual(s) assuming responsibility for the patient(s).

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, providers 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 provider 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.
III. TREATMENT PROTOCOLS

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

HOWEVER, THE GENERAL PATIENT CARE SECTION AND THE ALGORITHMS DO HAVE A SPECIFIC SEQUENCE TO BE FOLLOWED.

A. ABUSE/NEGLECT

1. Initiate General Patient Care.

ALL HEALTH CARE PROVIDERS ARE OBLIGATED BY LAW TO REPORT CASES OF SUSPECTED CHILD OR VULNERABLE ADULT ABUSE OR NEGLECT TO EITHER THE LOCAL POLICE OR SOCIAL SERVICE AGENCIES. DO NOT INITIATE REPORT IN FRONT OF THE PATIENT, PARENT, OR CAREGIVER.

DO NOT CONFRONT OR BECOME HOSTILE TO THE PARENT OR CAREGIVER.

2. Presentation

The patient may present with patterned burns or injuries suggesting intentional infliction, such as injuries in varying stages of healing, injuries scattered over multiple areas of the body, fractures, or injuries inconsistent with stated cause of injury. The patient, parent, or caregiver may respond inappropriately to the situation. Malnutrition or extreme lack of cleanliness of the patient or environment may indicate neglect. Signs of increased intracranial pressure (bulging fontanels and altered mental status in an infant) may also be seen.

3. Treatment

a) Stabilize injuries according to protocol.

b) Discourage patient from washing if sexual abuse is suspected.

c) Document the following information on the PCR:

(1) All verbatim statements made by the patient, the parent, or caregiver shall be placed in quotation marks, including statements made about the manner of the injuries.

(2) Any abnormal behavior of the patient, parent, and/or caregiver

(3) The condition of the environment and other residents present
A. ABUSE/NEGLECT (Continued)

(4) The time the police/welfare agency was notified and the name of the person notified

(5) The name of the receiving health care provider (RN, PA, MD) and any statements made

d) Treat injuries according to presentation.

4. Continue General Patient Care.
B. ALTERED MENTAL STATUS: SEIZURES

1. Initiate General Patient Care.

2. Presentation
   Seizures are a neuromuscular response to an underlying cause such as: epilepsy, hypoxia, hypoglycemia, hypoperfusion, head injury, CVA, alcohol or drug abuse. Consider recent history of possible illness, infection, fever, or stiff neck.

   DO NOT ATTEMPT TO FORCE ANY DEVICE INTO THE PATIENT’S MOUTH IF THE PATIENT IS STILL SEIZING.

3. Treatment
   a) If the patient is still seizing:
      (1) DO NOT RESTRAIN.
      (2) Protect patient from further injury.
      (3) Consider cause of seizure activity.
   b) When seizure activity has stopped:
      (1) Identify and treat injuries.
      (2) If patient is a known diabetic, glucose paste (10–15 grams) should be administered between the gum and cheek. Consider single additional dose of glucose paste if not improved after 10 minutes.
   c) Use glucometer and treat accordingly.
   d) Consider midazolam. (NEW ’16)
      If patient has no IV or IO in place, administer midazolam 5 mg IN or IM.
      If IV/IO is already in place: 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.
      If IV unavailable, 5 mg IM may be administered.
      Additional doses up to a maximum total dose 10 mg require medical consultation for all providers.
      If patient seizures are refractory to treatment, consider IO administration of midazolam.
      If midazolam is not available, consider diazepam.
      2.5 mg increments SLOW IVP/IM (IM requires all providers to obtain medical consultation).
B. ALTERED MENTAL STATUS: SEIZURES (Continued)

Maximum total dose 10 mg
If patient is in status, consider IO administration of diazepam.
If suspected severe nerve agent exposure, providers may administer midazolam 5 mg IM or diazepam (CANA) without medical consultation.

e) Establish IV access with LR.

f) If patient is pregnant, actively seizing, consider magnesium sulfate 4 grams IV/IO over 10 minutes.

g) If seizures persist, consult for second dose of magnesium sulfate.

IF PATIENT IS PREGNANT, USE MIDAZOLAM FOLLOWED BY MAGNESIUM SULFATE. MEDICAL CONSULTATION REQUIRED FOR PREGNANT PATIENTS WHO MAY REQUIRE LARGER DOSES OF MIDAZOLAM TO CONTROL SEIZURES.

IF, FOLLOWING ADMINISTRATION OF MAGNESIUM SULFATE, PATIENT EXHIBITS SIGNS OF TOXICITY, CONSIDER ADMINISTRATION OF CALCIUM CHLORIDE. CONSIDER CALCIUM CHLORIDE 500 MG IVP FOR RESPIRATORY DEPRESSION, DECREASED REFLEXES, FLACCID PARALYSIS, AND APNEA FOLLOWING MAGNESIUM SULFATE ADMINISTRATION. MEDICAL CONSULTATION REQUIRED.

h) If the patient is still seizing:

(1) DO NOT RESTRRAIN.

(2) Protect from further injury.

(3) Consider underlying cause of seizure.

i) When seizure activity has stopped:

(1) Identify and treat any injuries.

(2) If patient is a known diabetic, glucose paste (10–15 grams) should be administered between the gum and cheek. Consider single additional dose of glucose paste if not improved after 10 minutes.

j) Use glucometer and treat accordingly.

k) The paramedic may assist patients with the administration of their prescribed benzodiazepine.
B. ALTERED MENTAL STATUS: SEIZURES (Continued)

l) Consider midazolam for seizures lasting greater than 10 minutes. (NEW ’16)
   If patient has no IV or IO in place:
   Administer midazolam 0.2 mg/kg IN or IM. Maximum total dose 5 mg.

   If IV or IO is already in place:
   Administer midazolam 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2
   minutes. Maximum total dose 5 mg.

m) If patient is pregnant, actively seizing, consider magnesium sulfate 4 grams IV/IO
   over 10 minutes.

n) Establish IV/IO access with LR.

o) Administer fluid bolus, if appropriate
   20 mL/kg of LR IV/IO.

FOR A CHILD ACTIVELY SEIZING, ADMINISTER MIDAZOLAM IN/IM AND RESERVE IO FOR LIFE-
THREATENING ILLNESS.

p) Additional doses up to a maximum total dose 5 mg require medical consulta-
   tion for all providers.
   If patient’s seizures are refractory to treatment, consider IO administration
   of midazolam.
   If midazolam is not available, consider diazepam for seizures lasting
   greater than 10 minutes (paramedic may perform without consult for pa-
   tients with active seizures).
   Up to 0.2 mg/kg rectal
   Maximum total dose 10 mg
   OR
   0.1 mg/kg in 2.5 mg increments SLOW IVP/IO/IM (IM requires all
   providers to obtain medical consultation.)
   Maximum total dose 5 mg
   If suspected severe nerve agent exposure, providers may administer mid-
   azolam as above or diazepam (CANA) without medical consultation.

4. Continue General Patient Care.
C. ALTERED MENTAL STATUS: UNRESPONSIVE PERSON

1. Initiate General Patient Care.

2. Presentation
   Patients may exhibit confusion, focal motor sensory deficit, unusual behavior, unre- sponsiveness to verbal or painful stimulus.

   **ALERT** ALCOHOL CAN CAUSE ALTERED MENTAL STATUS BUT IS NOT COMMONLY A CAUSE OF TOTAL UNRESPONSIVENESS TO PAIN.

3. Treatment
   a) Obtain pulse oximetry, if available.
   
   b) Administer glucose paste (10–15 grams) between the gum and cheek. Consider single additional dose of glucose paste if not improved after 10 minutes.
   
   c) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.
   2 mg intranasal atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)
   
   d) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone
   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)
   Maximum single dose 0.4–2 mg (NEW ’16)
   
   e) Establish IV access with LR.
   Administer fluid bolus, if appropriate.
   20 mL/kg of LR IV
   
   f) Titrate to a systolic pressure of 100 mmHg.
   
   g) Consider obtaining blood sample using closed system.
   
   h) Use glucometer and treat accordingly.
   
   i) Consider an additional dose of naloxone. (NEW ’16)
   
   j) Consider additional fluid administration
   Maximum 2,000 mL without medical consultation.
C. ALTERED MENTAL STATUS: UNRESPONSIVE PERSON (Continued)

k) Obtain pulse oximetry if available.

l) Administer glucose paste (10–15 grams) between the gum and cheek. Consider single additional dose of glucose paste if not improved after 10 minutes.

m) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone

28 days to 4 years (NEW '16): Administer naloxone 0.8–1 mg intranasal atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

5 years to adult (NEW '16): Administer naloxone 2 mg intranasal atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

n) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.

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) (NEW '16)

Maximum single dose 0.4–2 mg

o) Consider repeating naloxone. (NEW '16)

p) Establish IV/IO access with LR.

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 IV/IO.

OR

For volume-sensitive children administer initial fluid bolus of 10 mL/kg LR IV/IO. If patient’s condition does not improve, administer the second bolus of fluid at 10 mL/kg LR IV/IO.

Volume-sensitive children include: neonates (birth to 28 days) (NEW '16), children with congenital heart disease, chronic lung disease, or chronic renal failure.

(1) Consider obtaining blood sample using closed system.

q) Use glucometer and treat accordingly.

r) Third and subsequent fluid boluses at 20 mL/kg IV/IO except in volume-sensitive children, then bolus at 10 mL/kg.

4. Continue General Patient Care.
D. APPARENT LIFE-THREATENING EVENT (ALTE)

1. Initiate General Patient Care.

2. Presentation
   An episode in an infant or child less than 2 years old that is frightening to the ob-
   server and is characterized by some combination of the following:
   a) Apnea (central or obstructive)
   b) Skin color change: cyanosis, erythema (redness), pallor, plethora (fluid overload)
   c) Marked change in muscle tone
   d) Choking or gagging not associated with feeding or a witnessed foreign body
      aspiration

   MOST PATIENTS WILL APPEAR STABLE AND EXHIBIT A NORMAL PHYSICAL EXAM UPON
   ASSESSMENT BY RESPONDING FIELD PERSONNEL. HOWEVER, THIS EPISODE MAY BE THE
   SIGN OF UNDERLYING SERIOUS ILLNESS OR INJURY. FURTHER EVALUATION BY MEDICAL
   STAFF IS REQUIRED AND IT IS ESSENTIAL TO TRANSPORT ALL PATIENTS WHO EXPERIENCED
   ALTE.

3. Treatment
   a) Perform an initial assessment utilizing the Pediatric Assessment Triangle.
   b) Obtain a description of the event including nature, duration, and severity.
   c) Obtain a medical history with emphasis on the following conditions:
      (1) Known chronic diseases
      (2) Evidence of seizure activity
      (3) Current or recent infections
      (4) Gastroesophageal reflux
      (5) Recent trauma
      (6) Medications (current or recent)
   d) Apply oxygen.
   e) Be prepared to assist with ventilation if this type of episode occurs again
      during transport.
   f) Assess environment for possible causes.
   g) Place patient on cardiac monitor.
   h) Consider establishing IV/IO access with LR.

   IF THE PARENT OR GUARDIAN REFUSES MEDICAL CARE OR TRANSPORT, PROVIDER SHALL
   CONTACT A PEDIATRIC BASE STATION PHYSICIAN.

4. Continue General Patient Care.
E. BEHAVIORAL EMERGENCIES

1. Initiate General Patient Care.

2. Presentation
   Behavior or actions that indicate the patient’s mental function is disturbed and may pose a threat to oneself or to others (suicide, threat of violence, or psychosis).

   THE PROVIDER SHOULD RECOGNIZE CRITICAL INCIDENT STRESS AS A STATE OF EMOTIONAL DISTRESS THAT DOES NOT NECESSARILY POSE A THREAT TO ONESELF OR OTHERS (E.G., DEATH IN THE FAMILY, BYSTANDERS AT A CRASH SCENE, OR REACTION TO VIOLENCE).

   THE PREHOSPITAL CARE PROVIDER SHOULD NOT BE PLACED IN ANY PHYSICAL JEOPARDY OR ASSUME ANY LAW ENFORCEMENT FUNCTIONS, ESPECIALLY WHEN WEAPONS AND/OR ACTS OF VIOLENCE ARE INVOLVED!

   LAW ENFORCEMENT SHOULD BE REQUESTED ON ALL CALLS INVOLVING POTENTIALLY VIOLENT PATIENTS.

3. Treatment
   a) When considering the prehospital use of restraints, a law enforcement officer should apply the device and accompany the provider and the patient in the ambulance.

   b) For interfacility transport, a physician order must be obtained for physical restraint.

   c) Implement SAFER model.

      (1) Stabilize the situation by containing and lowering the stimuli.

      (2) Assess and acknowledge the crisis.

      (3) Facilitate the identification and activation of resources (chaplain, family, friends, or police).

      (4) Encourage patient to use resources and take actions in his/her best interest.

      (5) Recovery or referral—leave patient in care of responsible person or professional or transport to appropriate facility.
E. BEHAVIORAL EMERGENCIES (Continued)

  d) Establish IV access with LR, if appropriate.
  e) Consider Chemical Restraint.

4. Continue General Patient Care.
1. The following algorithmic and standard formatted sections pertain to cardiac emergencies. Several guidelines apply to all algorithms when assessing and treating cardiac patients. These guidelines are:

   a) 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).

   b) As BLS/ALS guidelines indicate, definitive airway control is preferable; if this can be achieved, along with other initial interventions, then the earlier the better. However, defibrillation is more important initially if the patient can be ventilated without intubation.

   c) Cardiac Arrest:
      Immediately start CPR and apply AED or manual defibrillator as soon as possible; shock if indicated.
      The goal is to defibrillate as soon after stopping CPR as possible (ideally for manual defibrillator, in less than 5 seconds).
      After single shock, immediately restart CPR (do not perform pulse or EKG rhythm check) for 2 minutes, then assess for pulse and rhythm and apply single shock if indicated. Repeat this sequence of single shocks and 2 minutes of CPR.

   d) If unable to initiate an IV or perform endotracheal intubation within 5 minutes, continue with appropriate care and transport the patient as soon as possible to the appropriate hospital. Further attempts to initiate IV therapy or endotracheal intubation should be accomplished while en route to the receiving hospital.

   e) Only in a pediatric or neonatal arrest situation, naloxone, atropine, 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, and ten times the IV dose for epinephrine (1:1,000). All ET medications shall be diluted in 5 mL of LR for pediatric patients. (NEW '16)
UNIVERSAL ALGORITHM FOR ADULT EMERGENCY CARDIAC CARE FOR BLS

Unresponsive Not Breathing

Pulse?

YES
Support ventilation
ALS & transport

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
3. 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

YES

If unconscious and no trauma, place in recovery position

Pulse

NO

Begin CPR

VF/VT Present on Monitor

YES

Intubate Confirm Tube Placement Confirm Ventilations

Determine Rhythm & Possible Cause

Electrical Activity?

NO

YES

GO TO VT/VF ALGORITHM

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

Suspected Cause

Pulmonary Edema/CHF See Protocol

Chest Pain See Protocol

Dysrhythmia

Too Slow

Too Fast

GO TO BRADYCARDIA ALGORITHM

GO TO TACHYCARDIA ALGORITHM

NO

YES

GO TO ASYSTOLE ALGORITHM

GO TO PEA ALGORITHM

NO

YES

NO

YES

YES

NO

YES

NO

YES

NO

YES

NO

YES

NO

YES

NO

YES

NO
4. **UNIVERSAL ALGORITHM FOR PEDIATRIC (LESS THAN 18 YEARS OF AGE) EMERGENCY CARDIAC CARE FOR BLS**  
(If less than 1 hour old, refer to Newly Born Protocol)  
(NEW '16)

Unresponsive  
Not Breathing

**Pulse?**

- **YES**
  - Support ventilation
  - ALS & transport

- **NO**
  - Begin CPR  
    - Attach AED with pediatric capability
  - Analyze shockable rhythm?
    - **YES**
      - Defibrillate 1 time  
        - Resume CPR immediately for 2 minutes
    - **NO**
      - Resume CPR immediately for 2 minutes
5. **UNIVERSAL ALGORITHM FOR PEDIATRIC (LESS THAN 18 YEARS OF AGE)
EMERGENCY CARDIAC CARE FOR ALS
(If less than 1 hour old, refer to Newly Born Protocol) (NEW ’16)**

- **Assess Responsiveness**
  - Not Responsive: Call for Defibrillator, Assess Breathing
  - Responsive: Observe, Treat as Indicated

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

- **Pulse**
  - NO: Begin CPR, Attach AED with pediatric capability, 100 compressions/minute, 100% oxygen
  - YES: Oxygen as needed, **VENTILATE** as needed, Cardiac monitor, Vital signs, IV with LR, History & Physical, Detailed Assessment

- **Suspected Cause**
  - Altered Mental Status: See Protocol
  - Respiratory Distress
    - Allergic Reaction or Anaphylaxis: See Protocol, as appropriate
    - Asthma/COPD: See Protocol
    - Pulmonary Edema/CHF: See Protocol

- **Dysrhythmia**
  - Too Slow: **GO TO PEDIATRIC BRADYCARDIA ALGORITHM**
  - Too Fast: **GO TO PEDIATRIC TACHYCARDIA ALGORITHM**
G. CARDIAC EMERGENCIES: BRADYCARDIA

1. Initiate General Patient Care.

2. Presentation
   Patient may present with a slow heart rate and chest pain, shortness of breath, decreased level of consciousness, hypotension, hypoperfusion, pulmonary congestion, congestive heart failure, and/or acute myocardial infarction.

3. Treatment
   a) Place patient in position of comfort.
   b) Assess and treat for shock, if indicated.
   c) Continuously monitor airway and reassess vital signs every 5 minutes.
   d) Establish IV access with LR.
   e) If patient is hemodynamically unstable: initiate transcutaneous pacing (TCP).
   f) If TCP is unsuccessful or not available, administer atropine:
      0.5–1 mg IVP
      Atropine should be given in repeat doses in 3–5 minute intervals up to a total of 0.04 mg/kg.
   g) Consider dopamine
      2–20 mcg/kg/min
   h) If patient is hemodynamically stable and in Type II, second-degree AV Block or third-degree AV Block:
      (1) Consider/prepare for TCP.
      (2) If patient develops discomfort with TCP
         Administer opioid per Pain Management Protocol.
         OR
         Consider 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.)
   i) Refer to appropriate algorithm.
(a) - Serious signs and symptoms must be related to the slow rate. Signs and symptoms may include chest pain, shortness of breath, decreased level of consciousness, hypotension, hypoperfusion, pulmonary congestion, CHF, and/or AMI.

(b) - Do not delay TCP while awaiting IV or atropine to take effect if the patient is symptomatic.

(c) - Denervated transplanted hearts will not respond to atropine. Go at once to TCP.

(d) - Atropine shall be given in repeat doses in 3–5 minute intervals up to a total of 0.04 mg/kg. Consider shorter intervals in severe clinical conditions.

(e) - Never treat third-degree AV block or ventricular escape beats with amiodarone. (NEW '16)

(f) - In the presence of Mobitz II and third-degree AV block, medical consultation is required for atropine administration.

(g) - Requires medical consultation for administration of dopamine. Adults: titrate to systolic BP 100 mmHg or medical consultation directed BP. IV infusion pump is preferred.
5. PEDIATRIC BRADYCARDIA ALGORITHM
(If less than 1 hour old, refer to Newly Born Protocol) (NEW ’16)

Identify and treat underlying causes

Hemodynamically unstable? (a)

NO

Observe Support ABCs

YES

Begin CPR if HR less than 60 with poor perfusion despite oxygenation and ventilation

Bradycardia persists?

NO

Epinephrine (b)
IV/IO 0.01 mg/kg (1:10,000)
ET 0.1 mg/kg (1:1,000),
Dilute in 5 mL;
Repeat every 3–5 minutes

Atropine
IV/IO 0.02 mg/kg,
Maximum single dose 0.5 mg,
ET 0.04–0.06 mg/kg,
Dilute in 5 mL
Repeat once

Consider Transcutaneous Pacing

If pulseless arrest develops go to Cardiac Arrest Algorithm

YES

Possible causes of bradycardia
(Parenthesis) = Possible Therapies and Treatments

Hypovolemia (Volume Infusion) (c)
Hypoxia (Ventilation)
Hydrogen ion (acidosis) (d)
Hypo-/hyperkalemia (d,e)
Hypoglycemia (Glucometer Protocol)
Hypothermia (Warming)
Toxins (d,e)
Tamponade, cardiac (NDT)
Thrombus
Trauma

Pacer Age-Related Rates (NEW ’16)
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

(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 \(70 + (2 \times \text{years})\) = systolic BP for patients greater than 1 year of age.

(b) - Neonates (birth to 28 days) (NEW ’16), epinephrine ET 0.03 mg/kg (1:10,000) dilute with 1 mL.

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

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

(e) - Calcium chloride, 20 mg/kg (0.2 mL/kg) SLOW IVP/IO (50 mg/min). Max dose 1 gram.
H. CARDIAC EMERGENCIES: TACHYCARDIA

1. Initiate General Patient Care.

2. Presentation
   Patient may present with chest pain, shortness of breath, decreased level of consciousness, low blood pressure, hypoperfusion, pulmonary congestion, congestive heart failure, and/or acute myocardial infarction.

3. Treatment
   a) Place patient in position of comfort.

   b) Assess and treat for shock, if indicated.

   c) Continuously monitor airway and reassess vital signs every 5 minutes.

   d) Establish IV access with LR.

   e) Verify presence of pulse.

   f) If no pulse present, treat as pulseless VF/VT.

   g) If patient is hemodynamically unstable with a ventricular rate greater than 150, prepare for immediate cardioversion.

   h) If patient is hemodynamically stable, identify rhythm and proceed to appropriate algorithm.

   i) Place patient in position of comfort.

   j) Assess and treat for shock, if indicated.

   k) Continuously monitor airway and reassess vital signs every 5 minutes.

   l) Establish IV access with LR.

   m) Verify presence of pulse.

   n) If no pulse present, treat as pulseless VF/VT.
H. CARDIAC EMERGENCIES: TACHYCARDIA (Continued)

 o) If patient is hemodynamically unstable with a ventricular rate greater than 220 for an infant or 180 for a child, prepare for immediate cardioversion.

 p) If patient is hemodynamically stable, identify rhythm and proceed to appropriate algorithm.

 4. Continue General Patient Care.
5. ADULT TACHYCARDIA ALGORITHM (NEW ’16)

- Unstable condition must be related to the tachycardia. Signs and symptoms may include chest pain, shortness of breath, decreased level of consciousness, hypotension, hypoperfusion, pulmonary congestion, CHF, and/or AMI.

- Consider sedation (midazolam with medical consultation). However, overall patient status, including BP, may affect ability to administer sedative.

- Consider calcium chloride 500 mg IVP for hypotension induced by diltiazem. Medical consultation required. If rate does not slow in 15 minutes, administer a second dose of diltiazem (15–25 mg over 2 minutes). Medical consultation required.

- Be prepared for up to 40 seconds of asystole.

- If irregular, DO NOT administer amiodarone or adenosine. Cardiovert if unstable.

- If Torsades de Pointes, administer magnesium sulfate (1–2 grams IV/IO over 2 minutes).
Identify and treat underlying causes

Evaluate QRS duration

Narrow (less than or equal to 0.09 seconds)

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

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) (NEW '16), 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. Obtain 12-lead EKG prior to administration of amiodarone. (NEW '16)

(g) If Torsades de Pointes, administer magnesium sulfate (25 mg/kg IV/IO to a maximum of 2 grams over 2 minutes).
I. CARDIAC EMERGENCIES: CARDIAC ARREST

1. Initiate General Patient Care.

2. Presentation
   Patient must be unconscious, apneic, and pulseless.

3. Treatment
   a) Perform CPR.

   HIGH-QUALITY CONTINUOUS CPR WITH FREQUENT PROVIDER ROTATION IS AN ESSENTIAL COMPONENT IN THE SUCCESSFUL RESUSCITATION OF THE ARRESTED PATIENT. THIS MAY BE ACCOMPLISHED THROUGH MANUAL OR MECHANICAL MEANS AS APPROPRIATE.

   PERFORM CPR WHILE PREPARING FOR RHYTHM ANALYSIS AND DEFIBRILLATION.

   b) Utilize AED as appropriate.
   c) Transport
      (1) If no shock indicated, consider Termination of Resuscitation Protocol or transport immediately.
      (2) If shock indicated, defibrillate and resume CPR. Consider Termination of Resuscitation Protocol or transport ASAP.
      (3) If ROSC, refer to ROSC Protocol.
      (4) If no ROSC, consider Termination of Resuscitation Protocol or transport to the closest appropriate facility.

   d) Identify rhythm and treat according to appropriate algorithm.
   e) If no ROSC, consider Termination of Resuscitation Protocol or transport to the closest appropriate facility.
   f) If ROSC, refer to ROSC Protocol.

   g) Perform CPR.
   h) Utilize AED as appropriate (see AED Procedure).
   i) If no shock indicated, continue CPR and transport ASAP.
   j) If shock indicated, defibrillate, continue CPR, and transport ASAP.

   For patients who have not reached their 18th birthday:
   k) Identify rhythm and treat according to appropriate algorithm.
   l) If no ROSC, transport to the closest appropriate facility.
   m) If ROSC, perform 12-Lead EKG and transport the patient to Children’s National Health System 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 ED.
4. ADULT ASYSTOLE ALGORITHM

- Continue CPR
- Intubate \( \text{O}_2 \) (90–100%)
- Establish IV access with LR
- Confirm asystole in more than one lead

Consider Possible Causes

Epinephrine 1 mg IVP - Repeat every 3–5 minutes

Consider possible causes of asystole.
(Parenthesis) = Possible Therapies and Treatments

- Hypovolemia (Volume Infusion) (c)
- Cardiac Tamponade (Volume Infusion) (c)
- Tension Pneumothorax (Needle Decompression Thorocostomy–NDT)
- Massive Pulmonary Embolism
- Massive AMI
- Drug Overdose (a,b)
- Hypoxia (Ventilation)
- Hypothermia (Warming)
- Acidosis (a)
- Hyperkalemia (a,b)

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

(b) - Calcium chloride, 0.5–1 gram IVP, with medical consultation. See calcium chloride.

(c) - Volume infusion is 20 mL/kg.
5. PEDIATRIC CARDIAC ARREST ALGORITHM
(If less than 1 hour old, refer to the Newly Born Protocol) (NEW ’16)

Begin CPR
Attach monitor

VF/VT
Defibrillate 2 J/kg
Resume CPR immediately for 2 minutes

IV/IO Access
Defibrillate 4 J/kg
Resume CPR immediately for 2 minutes

Epinephrine (b)
IV/IO 0.01 mg/kg (1:10,000)
ET 0.1 mg/kg (1:1,000),
dilute with 5 mL
Repeat every 3–5 minutes

Defibrillate 4 J/kg
Resume CPR immediately for 2 minutes

Amiodarone 5 mg/kg IV/IO/
(Max single dose 300 mg) May repeat
twice to a maximum total dose of
15 mg/kg per day (a) (NEW ‘16)

Asystole/PEA
Consider possible causes

IV/IO Access
Epinephrine (b)
IV/IO 0.01 mg/kg (1:10,000)
ET 0.1 mg/kg (1:1,000),
dilute with 5 mL
Repeat every 3–5 minutes

Consider possible causes of asystole.
(Parenthesis) = Possible Therapies and Treatments
Hypovolemia (Volume Infusion) (e)
Hypoxia (Ventilation)
Hydrogen ion (acidosis) (c)
Hypo-/hyperkalemia (c,d)
Hypoglycemia (Glucometer Protocol)
Hypothermia (Warming)
Toxins (c,d)
Tamponade, cardiac
Tension pneumothorax (NDT)
Thrombus
Trauma

(a) - Continue cycle of epinephrine, defibrillation (at 4 J/kg), then amiodarone (NEW ’16). Defibrillate at increasing dosage: 6 J/kg, 8 J/kg, 10 J/kg.

(b) - Neonates (0–28 days), epinephrine ET 0.03 mg/kg (1:10,000) 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 two minutes before amiodarone).
6. ADULT PULSELESS ELECTRICAL ACTIVITY (PEA) ALGORITHM

Includes:
- EMD
- Pseudo EMD
- Brady-asystolic Rhythms
- Idioventricular Rhythms
- Ventricular Escape Rhythms
- Post-defibrillation Idioventricular Rhythms

Continue CPR
Intubate
IV with LR
Consider Possible Causes
Epinephrine 1 mg IVP. Repeat every 3–5 minutes.

Consider possible causes of PEA.
(Parenthesis) = Possible Therapies and Treatments

<table>
<thead>
<tr>
<th>Cause</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypovolemia</td>
<td>(Volume Infusion) (c)</td>
</tr>
<tr>
<td>Cardiac Tamponade</td>
<td>(Volume Infusion) (c)</td>
</tr>
<tr>
<td>Tension Pneumothorax</td>
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<td>Massive AMI</td>
<td></td>
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<tr>
<td>Drug Overdose</td>
<td>(a,b)</td>
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<tr>
<td>Hypoxia</td>
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<tr>
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</tr>
<tr>
<td>Acidosis</td>
<td>(a)</td>
</tr>
<tr>
<td>Hyperkalemia</td>
<td>(a,b)</td>
</tr>
</tbody>
</table>

(a) - Sodium bicarbonate 1 mEq/kg, with medical consultation. See sodium bicarbonate.
(b) - Calcium chloride, 0.5–1 gram IVP, with medical consultation. See calcium chloride.
(c) - Volume infusion is 20 mL/kg.
7. VENTRICULAR FIBRILLATION
PULSELESS VENTRICULAR TACHYCARDIA (NEW '16)

Perform CPR until defibrillator is attached
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

Intubate

IV with LR

Epinephrine
1 mg IVP
Repeat every 3–5 minutes

Defibrillate 1 time
Resume CPR immediately
for 2 minutes

(NEW '16) Amiodarone
300 mg IV/IO push
May repeat once 150 mg
IV/IO push (a) (b)

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, with medical consultation. See sodium bicarbonate.

(b) - If Torsades de Pointes is present, give magnesium sulfate 1–2 grams IV/IO over 2 minutes before amiodarone. (NEW '16)
J. RETURN OF SPONTANEOUS CIRCULATION (ROSC) (NEW ’16)

1. Initiate General Patient Care.

2. Presentation
   Patients revived from non-traumatic cardiac arrest.

3. Treatment
   a) Verify presence of carotid pulse. If absent, go to Cardiac Arrest Protocol.

   FREQUENTLY REASSESS FOR PRESENCE OF PULSE. IF ANY DOUBT AS TO PRESENCE OF PULSE, REINITIATE CHEST COMPRESSIONS AND RETURN TO APPROPRIATE ALGORITHM FOR CARDIAC ARREST.

   b) If apneic, or inadequate respirations, continue to support ventilations. Otherwise, use supplemental oxygen as needed.

   c) Reassess vital signs. Treat any abnormalities in accordance with relevant algorithms.

   d) If patient 18 years of age or older and comatose (GCS less than 8), initiate Neuroprotective Induced Hypothermia Protocol.

   e) Rendezvous with ALS or transport to nearest ED.

   f) Identify rhythm and treat according to appropriate algorithm.

   g) Obtain 12-lead EKG.

   h) Establish IV/IO access, if not yet obtained.

   i) Treat hypotension
      (1) If lungs are clear, consider fluid bolus. 20 mL/kg LR IV. Titrate to SBP of 100 mmHg.

      (2) With medical consultation, consider dopamine infusion.
         (a) Adjust infusion rate in accordance with blood pressure and clinical response.
         (b) Adult: Administer 2–20 mcg/kg/min IV/IO drip titrated to BP of 100 systolic or medical consultation selected BP; initial infusion rate 2–5 mcg/kg/min.
         (c) Pediatric: Administer 2–20 mcg/kg/min IV/IO drip titrated to age specific BP or medical consultation selected BP; initial infusion rate is 2 mcg/kg/min.

   j) Reasses need for intubation, if not yet performed.

   k) Evaluate for contributing causes. For example,
      (1) If chest pain currently or prior to arrest, see Chest Pain Protocol.
      (2) If signs and symptoms of CHF, follow Pulmonary Edema/CHF Protocol.
J. RETURN OF SPONTANEOUS CIRCULATION (ROSC) (continued)

l) If VF or VT during arrest and amiodarone not yet given, consider Amiodarone 150 mg IV/IO over ten minutes.\(^{(a)}\)

m) Initiate transport to appropriate facility. Most patients should go to a Cardiac Interventional Center. Consider helicopter transport.

(1) Except as under (2) below, most patients should be transported to a Cardiac Interventional Center.

(2) Transport to nearest ED

(a) If obvious non-cardiac cause for arrest (e.g., drowning, asphyxiation, opiate overdose), (If cause for arrest is in any way uncertain, patient must be transported to Cardiac Interventional Center, except as under b and c below)

OR

(b) If transport time to Cardiac Interventional Center is more than 45 minutes greater than transport time to nearest ED,

OR

(c) With medical consultation, if patient’s clinical instability will not allow for safe transport to Cardiac Interventional Center due to transport time.

(3) Medical Consultation may assist with determination of destination.

(4) Except as under (5) below, most pediatric patients should be transported to Children’s National Health System or Johns Hopkins Children’s Center. Consider helicopter transport.

(5) Transport to nearest ED

(a) If transport time to Children’s National Health System or Johns Hopkins Children’s Center is more than 30 minutes greater than transport time to nearest ED,

OR

(b) With medical consultation, if patient’s clinical instability will not allow for safe transport to one of the above centers due to transport time.

ALL POST-CARDIAC ARREST PATIENTS ARE PRIORITY 1, AND REQUIRE MEDICAL CONSULTATION. PEDIATRIC PATIENTS REQUIRE CONSULTATION WITH A PEDIATRIC BASE STATION AND MAY ASSIST IN DESTINATION DETERMINATION.

4. Continue General Patient Care.

(a) Presence of a perfusing sinus rhythm is necessary for the administration of amiodarone for the ROSC patient post VF/VT conversion.
K. TERMINATION OF RESUSCITATION (Medical and Traumatic)

IF ANY DOUBT EXISTS, INITIATE RESUSCITATION AND TRANSPORT

1. PURPOSE
   This protocol is designed to guide the provider in determining a futile resuscitation and managing the patient after this determination.

2. PROCEDURE
   a) Exclusions to this protocol.
      (1) If arrest is believed to be secondary to hypothermia or submersion, treat according to appropriate protocol and transport to the nearest appropriate facility.
      (2) If patient is pregnant, treat according to appropriate protocol and transport to the nearest appropriate facility.
      (3) If patient has not reached his/her 18th birthday, treat according to appropriate protocol and transport to the nearest appropriate facility.
   b) Medical Arrest
      (1) EMS providers may terminate resuscitation without medical consult when all three criteria are met.
         (a) The arrest was not witnessed by an EMS provider (and patient is unresponsive, pulseless, and apneic). AND
         (b) There is no shockable rhythm identified by an AED or there is asystole or PEA on a manual cardiac monitor. AND
         (c) There is no return of spontaneous circulation (ROSC) prior to decision to terminate resuscitation despite appropriate field EMS treatment that includes 15 minutes of minimally-interrupted EMS CPR. OR
      (2) EMS providers may terminate resuscitation with medical consult when there is no ROSC prior to decision to terminate resuscitation despite appropriate field EMS treatment that includes 15 minutes of minimally-interrupted CPR in the presence of an arrest witnessed by an EMS provider or the presence of a shockable rhythm.
   c) Trauma Arrest
      (1) EMS providers may terminate resuscitation without medical consult when both criteria are met. (If medical etiology is suspected, use “Medical Arrest” above.)
         (a) There are no signs of life. AND
         (b) The patient is in asystole. OR
      (2) EMS providers may terminate resuscitation with medical consult when both criteria are met in either blunt or penetrating trauma.
         (a) Blunt
            (i) There are no signs of life. AND
            (ii) The patient is in a rhythm other than asystole and there is no ROSC despite 15 minutes of appropriate treatment that includes 15 minutes of minimally-interrupted CPR.
K. TERMINATION OF RESUSCITATION (Medical and Traumatic) (Continued)

(b) Penetrating
   (i) There are no signs of life. **AND**
   (ii) The patient is in a rhythm other than asystole and there is no ROSC.

If less than 15 minutes from a trauma center, transport the patient. If transport time exceeds 15 minutes, consult.

**ALERT**

**THERE ARE SOME CAUSES OF TRAUMATIC CARDIOPULMONARY ARREST (E.G., PENETRATING TRAUMA) THAT MAY BE REVERSED IF APPROPRIATELY AND EMERGENTLY MANAGED. THEREFORE, EMS PROVIDERS SHOULD FOLLOW APPROPRIATE PROTOCOLS FOR TRAUMATIC ARREST INCLUDING APPROPRIATE AIRWAY MANAGEMENT AND CONSIDERATION FOR BILATERAL NEEDLE DECOMPRESSION THORACOSTOMY. HOWEVER, EVEN WITH THE APPLICATION OF THESE MANEUVERS, ASYSTOLE AND PULSELESSNESS FOR GREATER THAN 10 MINUTES ARE INDEPENDENT PREDICTORS OF MORTALITY.**

   d) Pronouncement of Death in the Field Protocol.
Exclusions to this Protocol
- Arrest secondary to hypothermia or submersion
- Patient is pregnant
- Patient has not reached his/her 18th Birthday

If any doubt exist, initiate resuscitation and transport.
Initiate Pronouncement of Death in the Field Protocol when termination occurs.
L. PRONOUNCEMENT OF DEATH IN THE FIELD

1. PURPOSE
This protocol is designed to guide the EMS provider in pronouncing death in the field.

Health General Article §5-202 provides that:

a) An individual is dead if, based on ordinary standards of medical practice, the individual has sustained either:
   (1) Irreversible cessation of circulatory and respiratory functions; or
   (2) Irreversible cessation of all functions of the entire brain, including the brain stem.

2. INDICATIONS
EMS providers may pronounce the death of a patient when one or more of the following criteria has been met.

a) Decapitation
b) Rigor mortis
c) Decomposition
d) Dependent lividity
e) Pulseless, apneic patient in a multi-casualty incident where system resources are required for the stabilization of living patients
f) Pulseless, apneic patient with an injury not compatible with life (with the exception of an obviously pregnant female where resuscitation attempts should be initiated and the patient transported to the nearest appropriate facility)
g) The EMS provider has terminated resuscitation per the Termination of Resuscitation Protocol.

3. PROCEDURE
a) Confirm that the patient is unresponsive, pulseless, and apneic.
b) The patient who meets criteria in 2.e may be “black” tagged during triage (by a BLS or ALS provider), but asystole must be confirmed by ALS provider before a formal pronouncement of death.
c) The patient who meets criteria in 2.f must be confirmed to be in asystole by ALS provider before a formal pronouncement of death. If the condition of the remains precludes obtaining a cardiac rhythm to confirm asystole (e.g., incineration, severe disruption of the torso, etc.), this must be documented on the patient care report.
d) Document the exact time and location of the pronouncement of death.
e) Notify law enforcement and follow local jurisdictional policies or, if death is pronounced during transport, deliver patient to emergency department and follow hospital policies.
M. EMS DNR/MOLST

THE FOLLOWING SECTION IS ABSTRACTED FROM THE ORIGINAL MARYLAND EMERGENCY MEDICAL SERVICES DO NOT RESUSCITATE PROGRAM 2ND REVISION (07/01/98). THE PAGE (pg.) AND THE CHAPTER (ch.) NUMBERS HAVE BEEN APPENDED TO THE FOLLOWING CHAPTER TITLES FOR EASY REFERENCE. BECAUSE THIS ABSTRACT IS CONDENSED FROM THE ORIGINAL DOCUMENT, SOME CHAPTER NUMBERS OR LETTERS WERE INTENTIONALLY LEFT OUT. PLEASE REFER TO THE ORIGINAL MARYLAND EMS/DNR DOCUMENT FOR FURTHER INFORMATION.

AS OF JANUARY 1, 2002, A COPY OF THE MARYLAND EMS DNR ORDER FORM CAN BE ACCEPTED IN LIEU OF THE ORIGINAL.

AS OF OCTOBER 1, 2011, THE MARYLAND MOLST FORM CAN BE ACCEPTED IN LIEU OF THE MARYLAND EMS/DNR FORM.

1. PREFACE EMS/DNR Order or MOLST forms, bracelets, and necklaces will recognize three patient options for care prior to arrest: (pg. 15 ch. A)
   a) **Option A (ALS) (MOLST A1)**—Maximal (Restorative) Care (with intubation) Before Arrest, then DNR
   b) **Option A (DNI) (MOLST A2)**—Comprehensive Efforts to Prevent Arrest But Do Not Intubate, then DNR
   c) **Option B (BLS) (MOLST B)**—Limited (Palliative) Care Only Before Arrest, then DNR

2. VALID EMS/DNR or MOLST BRACELET WITH INSERT or AUTHORIZED METAL EMBLEM HAS THE SAME EFFECT AS THE FORM. (pg. 17 ch. D)
   a) Typically only one EMS/DNR device is needed to initiate the EMS/DNR Protocol.
   b) EMS providers should only request a second instrument (e.g., a bracelet when a form has already been presented) if there is reason to question the validity of the first produced notification device.

3. RECIPROCITY (pg. 19 ch. E)
   a) A standardized EMS/DNR Order from another state may be honored.
   b) Out-of-state EMS/DNR Orders shall be followed to the full extent that is permissible by the Maryland Medical Protocols for Emergency Medical Services Providers. If there is misunderstanding with family members or others present at the scene or if there are other concerns about following the out of state EMS/DNR Order, contact online medical direction for assistance. (NEW ’16)
   c) See chart in “EMS/DNR Program” booklet for how other states will treat Maryland devices.

4. ORAL EMS/DNR ORDERS (pg. 19 ch. G)
   a) EMS providers may follow an oral EMS/DNR Order directly from a Maryland-licensed physician (MD or DO) or nurse practitioner who is physically present “on-site.” EMS shall not accept orders from private physician attendings or nurse practitioner by telephone.
   b) **EMS providers may follow an oral EMS/DNR Order from a Maryland-licensed physician “on-line” via the EMS Communications System (e.g., radio or telephone consult that is routed through a public service access point (PSAP) for audio recording).**
M. EMS DNR/MOLST (Continued)

5. ACCEPTABLE AND UNACCEPTABLE EMS/DNR ORDERS (pg. 19 ch. H)
   a) The following are acceptable for implementing the EMS/DNR Protocol:
      (1) Original Maryland EMS/DNR Order Form
      (2) Copy of the Maryland EMS/DNR Order Form
      (3) Other State EMS/DNR Order Form
      (4) Maryland EMS/DNR Bracelet Insert
      (5) Medic Alert DNR Bracelet or Necklace
      (6) Oral DNR Order from EMS System Medical Consultation
      (7) Oral DNR Order from other on-site physician or nurse practitioner
      (8) Maryland MOLST Form
      (9) Maryland MOLST Bracelet
   b) The following are not acceptable for implementing the EMS/DNR Protocol:
      (1) Advance directives without an EMS/DNR Order
      (2) Facility-specific DNR orders
      (3) Notes in medical records
      (4) Prescription pad orders
      (5) DNR stickers
      (6) An oral request from someone other than a physician or nurse practitioner
      (7) An oral order from an attending physician or nurse practitioner who is not on site
      (8) Any other device or instrument not listed above as acceptable

6. VALIDITY OF EARLIER VERSIONS OF EMS/DNR ORDERS (pg. 22 ch. K)
   a) Older versions of EMS/DNR Orders — i.e., initial version (1995 and first revision, 4/1/96) — continue to be valid and need not be updated unless the patient or authorized decision maker wishes to take advantage of new features available in the newer forms.
   b) EMS providers should treat older versions of EMS/DNR order (pre 7/1/98) as “Option B (BLS) - Limited (Palliative) Care Only Before Arrest, Then DNR.”

7. REVOCATION OF AN EMS/DNR ORDER (pg. 24 ch. M)
   a) An EMS/DNR Order may be revoked at any time by:
      (1) Physical cancellation or destruction of all EMS/DNR Order devices; or
      (2) An oral statement by the patient made directly to emergency medical services personnel requesting only palliative care or resuscitation. If the patient revokes an EMS/DNR order orally, the EMS/DNR Order notification devices do not need to be destroyed. EMS providers should thoroughly document the circumstances of the revocation. An oral revocation by a patient is only good for the single response or transport for which it was issued.
   b) An authorized decision-maker, other than the patient, cannot revoke an EMS/DNR Order orally. Because of the difficulty in identifying authorized decision makers in emergent situations, it is incumbent upon an authorized decision maker who has authority to revoke an EMS/DNR Order to either destroy or withhold all EMS/DNR Order devices, if they wish resuscitation for the patient.
M. EMS DNR/MOLST (Continued)

c) Section 5-610 of the Health Care Decision Act (Health General Article, Annotated Code of Maryland) makes willful concealment, cancellation, defacement, obliteration, or damage of an advance directive (including EMS/DNR Orders), without the patient’s or authorized decision maker’s consent, a misdemeanor subject to a fine not exceeding $10,000, imprisonment not exceeding one year, or both.

8. ANTICIPATED LOCATIONS FOR EMS/DNR ORDER FORMS: (pg. 25 ch. N)
EMS personnel shall be directed to look for an EMS/DNR Order in the following places:

a) About a patient’s wrist, hung from a necklace, or safety-pinned to a patient’s clothing.
b) At medical facilities, in the patient’s chart.
c) In residences and domicile facilities, by the bedside, behind the patient’s bedroom door, or on the refrigerator door.
d) In schools and educational institutions, in the nurse’s office, health room, or with the student’s attendant caregiver/aide.
e) Family or caregivers will be expected to retrieve the original EMS/DNR Order prior to the ambulance’s arrival.

9. IDENTIFICATION OF PATIENT (pg. 25 ch. O)

a) If the patient is able, the patient can self-identify during the initial assessment.
b) If the patient is unable to communicate, then family, caregivers, or bystanders can identify the patient for EMS providers.
c) If an EMS/DNR vinyl bracelet with insert or metal emblem (bracelet or necklace) is attached to a patient (on wrist, pendant from neck, pinned to clothing, etc.) the patient’s identity can be reasonably assumed by EMS providers.
d) If an EMS/DNR vinyl bracelet insert or metal emblem (bracelet or necklace) is found detached from the patient, EMS personnel must treat it as an EMS/DNR Order form and identify the subject of the EMS/DNR Order as the patient. A valid bracelet insert alone, without the vinyl bracelet, is a valid EMS/DNR Order so long as EMS providers confirm the patient’s identity (pg. 17 ch. D).
e) If EMS personnel are unable to ascertain with reasonable certainty, when required to do so, that the subject of the EMS/DNR Order is the patient, they may resuscitate the patient.

10. HEALTH PROVIDER/EMS PERSONNEL IMMUNITY (pg. 26 ch. R)

a) General immunity provisions, such as Good Samaritan immunity for volunteers and sovereign immunity for government employees, may apply under specific circumstances.
b) In addition to other immunity that may be provided for in law, the Health Care Decisions Act provides the following specific immunity in cases involving the provision, withdrawal, or withholding of care that may be life-sustaining in nature:

(1) EMS providers are not subject to criminal prosecution or civil liability or deemed to have engaged in unprofessional conduct as determined by the appropriate licensing, registering, or certifying authority as a result of withholding or withdrawing any health care under authorization obtained in accordance with the Health Care Decisions Act. See HG (5-609(a)(1)).

(2) EMS providers providing, withholding, or withdrawing treatment under authorization obtained under the Health Care Decisions Act do not incur liability arising out of any claim to the extent the claim is based on lack of consent or authorization for the action. See HG (5-609(a)(2)).

(3) EMS providers providing treatment because they reasonably believe that an EMS/DNR order, other than a bracelet, is not valid, do not incur liability arising out of any claim to the extent the claim is based on lack of consent or authorization for the action. See HG (5-608(d)).

11. EMS/DNR MEDICAL PROTOCOLS (pg. 29 ch. T)

a) DISPATCH

(1) Option B EMS/DNR patients (7/98 version) or patients with older version EMS/DNR orders (pg. 22 ch K) only require a BLS response. Once the on-scene BLS provider has determined the need for additional pain control, an ALS Rendezvous may be requested. Medevac requests are not appropriate for these patients.

(2) Option A or A (DNI) EMS/DNR patients (7/98 version) who are not in arrest may require a range of responses from BLS through the highest echelon of response available. This will depend on the information available to dispatch and the service requested. The response complement in these cases will be dictated by local standard operating procedures (SOP).

(3) If a dispatch center is unclear whether the DNR order is an EMS/DNR order or is unclear about the pre-arrest patient care option selected (A, A (DNI), or B), the dispatch center shall dispatch the appropriate resources based on the information available.

(4) In the absence of knowledge to the contrary, information from medical professionals at a health care facility about the EMS/DNR status of a patient may be presumed to be reliable.
M. EMS DNR/MOLST (Continued)

b) PERFORM LIMITED PATIENT ASSESSMENT
   Vital signs:
   (1) Check for absence of a palpable pulse.
   (2) Check for absence of spontaneous respirations in an unresponsive patient.
   (3) Check for a valid EMS/DNR Order or MOLST form; vinyl bracelet insert worn either on the wrist, as a necklace, or pinned to clothing; or for a metal em-blem (bracelet or necklace).

c) RESUSCITATE/DO NOT RESUSCITATE CRITERIA
   (1) If an EMS/DNR Order is not present, revoked, or otherwise void, the EMS provider shall treat and, if necessary, transport the patient.
   (2) If an EMS/DNR Order is not present, but the EMS provider believes that resuscitation or further resuscitation is futile, they may initiate the Termination of Resuscitation Protocol.
   (3) If a valid EMS/DNR order is found and the patient is in cardiac or respiratory arrest, no resuscitative measures shall be initiated.
   (4) If the patient is conscious and able to communicate that he/she revokes the EMS/DNR orally directly to EMS providers, EMS providers shall treat and, if necessary, transport the patient.
   (5) If the EMS/DNR patient (Option A, A (DNI), or B) arrests, withhold or withdraw further resuscitation and provide support to the family and caregivers. Consider notifying appropriate personnel.

d) OPTION A (MOLST A1) – MAXIMAL (RESTORATIVE) CARE PROTOCOL
   (1) When Option A - “Maximal (Restorative) Care (with intubation) Before Arrest, then DNR” is selected on an EMS/DNR Order or MOLST form, the patient shall receive the full scope of restorative interventions permissible under the Maryland EMS Medical Protocols (including Continuous Positive Airway Pressure (CPAP), cardiac monitoring, synchronized cardioversion for pulse-present ventricular or supraventricular tachycardia, cardiac pacing for pulse-present symptomatic bradycardia, insertion of IVs, and drug therapy), in an attempt to forestall cardiac or respiratory arrest.
   (2) This option was requested primarily by long-term care facilities for their patients who are on DNR orders for potentially prolonged periods of time. Many of these patients are less concerned about palliation of pain and more concerned about the quality of life after a stroke or heart attack. The primary medical conditions seen in the field necessitating this option have been the desire to administer dextrose for diabetic emergencies and epinephrine for anaphylactic reactions in patients who, upon arrest, are not to be resusci-tated.
M. EMS DNR/MOLST (Continued)

(3) If, despite these efforts, the patient becomes pulseless or stops breathing spontaneously, EMS providers shall then withhold or withdraw cardiopulmonary resuscitation including, but not limited to, no CPR, no cardiac pacing, no defibrillation, withdrawal of active ventilatory assistance upon cardiac arrest, and withholding or withdrawal of drug therapy (e.g., chemical resuscitation).

e) **OPTION A (DNI) (MOLST A2) – COMPREHENSIVE EFFORTS TO PREVENT ARREST BUT DO NOT INTUBATE, THEN DNR**

(1) Option A (DNI) is exactly the same as Option A, which may include limited ventilatory support by CPAP or BiPAP, but Do Not Intubate.

(2) Therefore, inappropriate care for “Option A (DNI) – Comprehensive Efforts to Prevent Arrest but Do Not Intubate, then DNR” would be nasal or oral intubation.

**IF MAXIMAL CARE IS SELECTED AND THE PATIENT’S CONDITION REQUIRES ALS, AN ALS UNIT SHOULD BE REQUESTED IF FEASIBLE GIVEN THE LOCATION OF THE INCIDENT RELATIVE TO THE NEAREST APPROPRIATE FACILITY, THE AVAILABILITY OF AN ALS UNIT, AND ITS ABILITY TO ARRIVE OR RENDEZVOUS IN A MEDICALLY APPROPRIATE PERIOD OF TIME.**

f) **OPTION B (MOLST B) – PALLIATIVE CARE PROTOCOL**

(1) Supportive Care for Control of Signs and Symptoms

(a) Respiratory distress

(i) Open the airway using non-invasive means (e.g., chin lift, jaw thrust, finger sweep, nasopharyngeal airway, oropharyngeal airway, and Heimlich maneuver, but no laryngoscopy, no Magill forceps, no cricothyroidotomy, and no tracheostomy).

(ii) Administer O₂ as follows:

a. If the patient is not on a ventilator and would benefit from oxygen therapy, provide passive oxygen via nasal cannula or non-rebreather mask (but no positive pressure oxygen via ambu bag, demand valve, or ventilator).

b. If the patient is found on an outpatient ventilator and is not in cardiac arrest, maintain ventilatory support during transport to the hospital.

c. If the patient is found on an outpatient ventilator and is in cardiac arrest, contact on-line medical direction to consult about disconnecting the ventilator.

(iii) Maintain an open airway by non-invasive means (e.g., chin lift, jaw thrust, finger sweep, nasopharyngeal airway, oropharyngeal airway, and Heimlich maneuver, but no laryngoscope, no Magill forceps, no cricothyroidotomy, and no tracheostomy).

(iv) Suction as necessary.

(v) Position for comfort.
M. EMS DNR/MOLST (Continued)

(b) External bleeding
   (i) Standard treatment (direct pressure with dressing, tourniquet)
   (ii) No IVs
(c) Immobilize fractures using skills and devices that minimize pain.
(d) Uncontrolled pain or other symptoms (e.g., severe nausea)
   (i) Allow patient, family, or health care providers (other than the prehospital provider) to administer patient’s prescribed medications. Such health care providers administering medication will not have to accompany the patient to the hospital.
   (ii) Patient controlled analgesia (PCA) systems for pain medication delivery and other patient-controlled medication (PCM) systems shall be left in place in DNR patients and monitored to the extent possible according to the provider’s level of certification or licensure.
   (iii) For the patient with significant pain and/or pain with a prolonged transport, opioid may be administered.
(e) Existing IV lines may be in place and if so, shall be monitored to the extent possible according to the provider’s level of certification and licensure.

(2) Inappropriate Care for a Palliative Care Patient
   (a) Cardiac monitoring, including 12-lead EKG, pacing, cardioversion, and defibrillation
   (b) Initiation of IV therapy (except for morphine and fentanyl administration for pain control as in 1 (d) (iii)) (NEW ’16)
   (c) EMS-initiated medications (except oxygen, and morphine or fentanyl administration for pain control as in 1 (d) (iii)) (NEW ’16)
   (d) CPR
   (e) Intubation (alternative airway device, endotracheal, nasotracheal, or gastric tube)
   (f) Active ventilatory assistance, unless on an outpatient ventilator (pg. 32 ch. 5)

g) TRANSPORT
   (1) Upon request of the patient, family, or caregivers and in lieu of transport to a hospital-based emergency department, EMS providers may transport Option B EMS/DNR patients who require transportation for pain control or symptom management or respite care to a specified inpatient hospice facility.
   (2) A current list of those facilities is available from the MIEMSS Program Development Office 410-706-4367 (4DNR). The receiving status of a particular facility can be ascertained from EMRC (24 hours a day) by EMS radio, EMSTEL, or red phone, or by calling 800-492-3805.
M. EMS DNR/MOLST (Continued)

(3) The State EMS Board may authorize additional facilities under 6.2.2 or 6.2.4 (pp. 35-36), if recognized in the future by DHMH in accordance with 42 CFR 418.98 and 42 CFR 418.100. EMS jurisdictions and commercial ambulance services will be notified by MIEMSS of any facilities that become eligible and elect to receive patients by ambulance, become ineligible, or elect to discontinue their participation.

(4) Take a copy of EMS/DNR Order or MOLST form, vinyl bracelet with insert, or metal emblem (bracelet or necklace) to the hospital with the patient. If returning the patient from a previous transport, be sure to request a copy of the EMS/DNR Order form, vinyl bracelet with insert, or metal emblem (bracelet or necklace) from the staff (see pg. 20 ch H2 and the “EMS/DNR Order Retrieval Strategies” on pg. 58 of the EMS/DNR program booklet).

h) COMMUNICATIONS
   (1) Consultation requirements for Option A EMS/DNR patients shall be dictated by the Maryland EMS Medical Protocols in accordance with the patient’s medical needs. EMS providers shall notify the hospital of the patient’s EMS/DNR status (i.e., Option A) and the identity of patient’s physician or nurse practitioner.
   
   (2) No consultation is required for the Option B EMS/DNR patients. The receiving hospital or inpatient hospice facility should be notified to expect the patient and prepare accordingly. Also make the hospital or inpatient facility aware of the patient’s EMS/DNR status (i.e., Option B) and the identity of the patient’s physician or nurse practitioner.
   
   (3) If there is misunderstanding with family members or others present at the scene or if there are other concerns about following the EMS/DNR Order and the patient’s condition permits, contact the physician or nurse practitioner signing the order, or the patient’s hospice program, or on-line medical direction for assistance.

i) DOCUMENTATION
   (1) If possible, make or retain a copy of the EMS/DNR Order or MOLST form and attach it to the official copy of the call runsheet that is kept by the EMS service. Having a copy of the EMS/DNR Order or MOLST form can significantly reduce documentation requirements. Encourage sending facilities to provide you with a copy of the EMS/DNR order or MOLST form, in addition to an original of the order, with the patient’s transfer documents.
M. EMS DNR/MOLST (Continued)

(2) If the EMS/DNR Protocol is initiated:
   (a) Document, in the narrative section:
      (i) Who gave you the EMS/DNR Order or MOLST form (as an applicable
          person physically providing the written order, name of on-site physi-
          cian or nurse practitioner, or name of on-line medical direction physi-
          cian) or
      (ii) Where the EMS/DNR Order or MOLST form was found;
   (b) Document the EMS/DNR order number, the effective date of the order,
       the name of the patient, the patient’s date of birth, and the name of the
       physician, nurse practitioner, or physician assistant who signed the order;
   (c) Document the time the EMS/DNR Protocol was initiated;
   (d) Document any care rendered;
   (e) If the patient arrests while under your care, document the time the
       patient lost spontaneous respirations or palpable pulse, if able to deter-
       mine, and
   (f) If the patient arrests while under your care, document the chain of cus-
       tody until the body is out of custody of EMS.

(3) If resuscitation protocols are initiated, document:
   (a) Care rendered as per normal practice;
   (b) The reason the EMS/DNR Protocol was not initiated, if relevant (e.g., un-
       able to find EMS/DNR Order, EMS/DNR is not or does not appear to be
       valid, patient request);
   (c) If resuscitation was started because there was reasonable doubt as to
       the validity of an EMS/DNR Order;
      (i) The EMS/DNR Order number, the effective date of the order, the
          name of the patient, the patient’s date of birth, and the name of the
          physician, nurse practitioner, or physician assistant signing the order;
       and
      (ii) Who gave you the EMS/DNR Order or where the EMS/DNR Order or
          MOLST form was found.

(4) Transfer any EMS/DNR Order or MOLST form to the appropriate
    authorities (e.g., to hospital or in-patient hospice personnel of the facility
    where the patient was transferred or, if the patient is deceased, to the physi-
    cian/police/medical examiner). If possible at the receiving facility, and if not
    already done, make a copy of the EMS/DNR Order or MOLST form.
    DO NOT RETAIN an original EMS/DNR Order or MOLST form.
M. EMS DNR/MOLST (Continued)

(5) If a copy of the EMS/DNR Order or MOLST form is available to EMS providers, it shall be attached to the official copy of the call runsheet that is retained by the EMS service.

(6) A vinyl bracelet with insert or metal emblem (bracelet or necklace) shall be left where found on the patient. Bracelets or metal emblems shall not be removed without the permission of the patient or the patient’s authorized decision maker and, when possible, shall be returned with the patient to the sending facility (see pg.16 ch. C of the EMS/DNR Program booklet).

j) PATIENT DISPOSITION IF NOT TRANSPORTED
If the EMS/DNR Protocol is implemented and the patient is not transported because the patient arrested at the response site, EMS personnel shall:

(1) Follow local operational procedures for handling deceased patients (see “How to Best Tell the Worst News” on pp. 105–106 of the EMS/DNR program booklet);

(2) Do not remove an EMS/DNR vinyl bracelet or metal emblem (bracelet or necklace) from the deceased patient;

(3) Law enforcement personnel or a representative of the medical examiner’s office needs to be notified only in the case of sudden or unanticipated death that occurs:
   (a) By violence
   (b) By suicide
   (c) As a result of an accident
   (d) Suddenly, if the deceased was in apparent good health, or
   (e) In any suspicious or unusual manner.
N. EMS DNR 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 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.
O. CARDIAC EMERGENCIES: CHEST PAIN/ACUTE CORONARY SYNDROME

1. Initiate General Patient Care.

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.

ACUTE CORONARY SYNDROME (ACS) IS DEFINED AS PATIENTS PRESENTING WITH ANGINA OR
ANGINAL EQUIVALENTS SUCH AS SHORTNESS OF BREATH; CHEST, EPIGASTRIC, ARM, OR JAW
PAIN OR DISCOMFORT; DIAPHORESIS; AND/OR NAUSEA.

3. Treatment
   a) Place patient in position of comfort.
   b) Assist patient with administration of patient’s own prescribed nitroglycerin. May
      be repeated in 3–5 minutes if chest pain persists, blood pressure is greater than
      90 mmHg, and pulse is greater than 60 bpm. Maximum three doses total (pa-
      tient and EMT assisted).
   c) Assess and treat for shock if indicated.
   d) Continuously monitor airway and reassess vital signs every 5 minutes.
   e) Consider aspirin 324 mg or 325 mg chewed, if acute myocardial infarction is
      suspected.

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.
MEDICAL CONSULTATION IS REQUIRED TO OVERRIDE THIS CONTRAINDICATION.

IF THE PATIENT’S BLOOD PRESSURE DROPS MORE THAN 20 mmHg AFTER ADMINISTRATION
OF NITROGLYCERIN, OBTAIN MEDICAL CONSULTATION BEFORE FURTHER ADMINISTRATION.

f) Additional doses of nitroglycerin require medical consultation.

   g) Establish IV access with LR.
   h) Shall perform a 12-lead EKG for patients with ACS.
      (If trained, providers may perform a 15-lead EKG.)
   i) If patient has a prescription or previous history of nitroglycerin use, administer
      nitroglycerin: 0.4 mg SL. May be repeated if symptoms persist, and BP is greater
      than 90 mmHg and pulse is greater than 60 bpm, to a maximum dose of 1.2 mg.
j) If patient does not have a prescription or previous history of nitroglycerin use, an IV must be established prior to administration; then administer nitroglycerin as above.

k) If IV cannot be established, nitroglycerin may be administered with medical consultation.

l) Identify rhythm and treat according to appropriate algorithm.

m) Administer additional doses of nitroglycerin.


CONSULT A PEDIATRIC BASE STATION FOR CHILDREN (WHO HAVE NOT REACHED THEIR 18TH BIRTHDAY) WITH CHEST PAIN WITH ASSOCIATED DYSRHYTHMIAS, CARDIAC DISEASE, OR BLUNT CHEST TRAUMA.

4. Continue General Patient Care.
P. CARDIAC EMERGENCIES: HYPERKALEMIA
(RENAL DIALYSIS/FAILURE OR CRUSH SYNDROME)

1. Initiate General Patient Care.

2. Presentation
   Certain conditions may produce an elevated serum potassium level that can cause hemodynamic complications.

3. Treatment
   a) Patients must meet the following criteria:
      (1) Suspected hyperkalemia patient
         (a) Renal dialysis/failure with poor or non-functioning kidneys or
         (b) Crush syndrome or patients with functional kidneys by history
         **AND**
         (2) Hemodynamically unstable renal dialysis patients or patients suspected of having an elevated potassium with bradycardia and wide QRS complexes.
   b) Place patient in position of comfort.
   c) Assess and treat for shock, if indicated.
   d) Continuously monitor airway and reassess vital signs every 5 minutes.
   e) Establish IV access with LR.
   f) Initiate Bradycardia Protocol.
   g) Consider calcium chloride 0.5–1 gram SLOW IV over 3–5 minutes. Maximum dose 1 gram or 10 mL.
   h) Consider sodium bicarbonate 50 mEq IV over 5 minutes.
   i) Consider albuterol 20 mg (high dose) via nebulizer (if available).
   FLUSH IV WITH 5 ML OF LR BETWEEN CALCIUM AND SODIUM BICARBONATE ADMINISTRATION.
   j) Crush syndrome or patients with functional kidneys by history
      Consider 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 (reserve for patient suspected of crush syndrome or patients with functional kidneys by history).
P. CARDIAC EMERGENCIES: HYPERKALEMIA (Continued)

k) Place patient in position of comfort.

l) Assess and treat for shock, if indicated.

m) Continuously monitor airway and reassess vital signs every 5 minutes.

n) Establish IV access with LR.

o) Initiate Bradycardia Protocol.

p) Administer calcium chloride 20 mg/kg (0.2 mL/kg) slow IVP/IO (50 mg/min)
   Maximum dose 1 gram or 10 mL.

q) Consider albuterol via nebulizer
   (1) For patients 2 years of age or greater, administer albuterol 2.5 mg.
   (2) For patients less than 2 years of age, administer albuterol 1.25 mg.

   FLUSH IV WITH 5 ML OF LR BETWEEN CALCIUM AND BICARBONATE ADMINISTRATION.

r) Crush syndrome or patients with functional kidneys by history
   Consider sodium bicarbonate 1 mEq/kg IV over 5 minutes. Maximum
dose 50 mEq. (Reserve for patient suspected of crush syndrome or pa-
tients with functional kidneys by history.)

4. Continue General Patient Care.
Q. CARDIAC EMERGENCIES: IMPLANTABLE CARDIOVERTER DEFIBRILLATOR (ICD) MALFUNCTION

1. Initiate General Patient Care.

2. Presentation
   An implantable cardioverter defibrillator (ICD) is a device that delivers an internal de-
   fibrillation (shock) whenever the patient’s heart rhythm/rate exceeds defined limits. EMS
   providers may encounter ICD devices that are appropriately or inappropriately delivering shock therapy. Internal shocks cause patient discomfort but DO NOT pose a danger to EMS personnel even when in direct contact with patient receiving an internal shock.

3. Treatment
   a) Place patient in position of comfort.
   b) Assess and treat for shock if indicated.
   c) Continuously monitor airway and reassess vitals every 5 minutes.

   IF PATIENT IS IN CARDIAC ARREST, PERFORM CPR AND USE AED AS APPROPRIATE DESPITE THE PATIENT’S ICD, WHICH MAY OR MAY NOT BE DELIVERING SHOCKS.

   d) Establish IV access with LR.
   e) Monitor cardiac rhythm and treat according to appropriate algorithm(s).
   f) ICD deactivation: Patient must meet the following criteria:
      (1) Three or more distinct shocks and
      (2) Obvious device malfunction with an EMS provider-witnessed inappropriate shock (e.g., alert patient in atrial fibrillation with rapid ventricular rate or SVT)
   g) Place an EMS donut magnet directly over device. Magnet placed directly over will deactivate device and shocks will not be delivered. After defibrillator is deactivated, tape magnet firmly in place and treat according to the appropriate algorithm(s).

   IF THE PATIENT HAS A COMBINATION ICD AND PACEMAKER, DEACTIVATING THE ICD MAY OR MAY NOT DEACTIVATE THE PACEMAKER.

   h) Regardless of the decision to deactivate the ICD device, be prepared to manage the underlying rhythm (e.g., treat wide complex tachycardia with cardioversion or amiodarone per protocol as appropriate). (NEW ’16)
Q. CARDIAC EMERGENCIES: IMPLANTABLE CARDIOVERTER DEFIBRILLATOR (ICD) MALFUNCTION (Continued)

IF 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.

CONTINUE CHEST COMPRESSIONS FOR PEDIATRIC PATIENTS WHO REMAIN POORLY PERFUSED DESPITE PACEMAKER CAPTURE.

i) 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.
   **OR**
   (2) Midazolam 0.1 mg/kg SLOW IV/IN/IM/IO. Maximum single dose is 5 mg. (Paramedic may perform without consult.)
   IN administration max 1 mL per nare
   IM administration requires all providers to obtain consultation

j) Transport to the closest appropriate facility.

Consult a Pediatric Base Station for children (who have not reached their 18th birthday) with an ICD device delivering shock therapy or malfunctioning.

k) 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.
   **OR**
   (2) 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 providers to obtain medical consultation.)
   Maximum total dose 5 mg.

l) Transport to the closest appropriate facility.

4. Continue General Patient Care.
R. CARDIAC EMERGENCIES: ST ELEVATION MYOCARDIAL INFARCTION (STEMI)

1. Initiate General Patient Care.

2. Presentation

**ACUTE CORONARY SYNDROME (ACS) IS DEFINED AS PATIENTS PRESENTING WITH ANGINA OR ANGINAL EQUIVALENTS SUCH AS SHORTNESS OF BREATH; CHEST, EPIGASTRIC, ARM, OR JAW PAIN OR DISCOMFORT; DIAPHORESIS; AND/OR NAUSEA.**

Inclusion Criteria:
Patient presents with Acute Coronary Syndrome (ACS) symptoms and has one of the following in a diagnostic quality EKG:

a) Greater than 1 mm of ST elevation in two or more contiguous limb leads
b) Greater than 1.5 mm of ST elevation in two or more precordial leads (in women)
c) Greater than 2 mm of ST elevation in two or more precordial leads (in men)
d) Anterior, Inferior, or Lateral MI: ST elevation greater than 1 mm in two or more contiguous leads and
   QRS complex is narrower than 0.12 seconds; (if wider than 0.12, you are unable to diagnose as STEMI)

   OR
   e) Posterior MI: ST depression greater than 1 mm in V1 and V2 with an R/S ratio of greater than or equal to one and
      QRS complex is narrower than 0.12 seconds; (if wider than 0.12, you are unable to diagnose as STEMI)


Detection of right ventricular and posterior wall infarction is important, as approximately 40% of patients with inferior wall infarctions have right ventricular and/or posterior wall involvement, which predisposes them to more complications and increased mortality.
Consider the following presentations as indicative of increased cardiovascular risk and request guidance from the closest appropriate EMS Base Station or Cardiac Interventional Center.

a) **Left bundle branch block (LBBB):** LBBB is rare in the setting of acute myocardial infarction and often indicates underlying cardiovascular disease. LBBB is more likely to signal a myocardial infarction if the one of the following conditions are met:
   1) Patient presents in cardiogenic shock
   2) EKG shows excessive ST segment elevation greater than 5 mm
   3) EKG shows ST segment deviation (elevation or depression) in the same direction as the QRS complex. This concept is known as inappropriate concordance.

b) **Wellens’ Wave:** Biphasic T waves or deeply inverted T waves in precordial leads (V2-V3, +/-V4).

c) **ST segment elevation in Lead aVR:** Multilead ST segment depression with coexisting ST segment elevation in lead aVR.

d) **Hyperacute T waves:** Peaked, broad-based T waves.

### 3. Treatment

a) Follow Chest Pain Protocol for nitrate, aspirin, and pain management.

b) If patient meets above STEMI criteria, this patient is a Priority 1 patient and requires a medical consult.

c) If a patient meets one of the above condition sets for STEMI inclusion criteria, the patient 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.

(1) When indicated and based on the EMS provider’s report, the Base Station physician at the receiving Cardiac Interventional Center will activate its Cardiac Interventional Team.

(2) The receiving ED physician will determine if the patient can bypass the ED and go directly to the cardiac catheterization lab to meet the cardiac interventional team.

(3) If the patient cannot be delivered to a Cardiac Interventional Center within the allotted time, complete the Fibrinolytic Therapy Checklist for STEMI.

(a) If the patient meets all of the criteria for fibrinolytic therapy, transport to the nearest ED.

(b) If the patient does not meet all of the criteria for fibrinolytic therapy, consult with the nearest Cardiac Interventional Center and the nearest ED to determine the most appropriate receiving facility.
R. CARDIAC EMERGENCIES: ST ELEVATION MYOCARDIAL INFARCTION
(STEMI) (Continued)

d) If patient does not have EKG ST elevations greater than 1 mm in two contiguous
leads, the patient shall be transported to the closest appropriate facility.
e) If a patient presents with IWMI, obtain a tracing of V4R to rule out right ventricu-
lar involvement. If ST elevation noted in V4R, withhold nitrates. The triad of RVMI
often includes clear lung sounds, hypotension, and JVD. 40% of IWMI have right
ventricular involvement. If hypotensive with clear lung sounds, administer 250–
500 mL of LR. For additional bolus, perform medical consultation.

CONSULT A PEDIATRIC BASE STATION FOR CHILDREN WITH ST ELEVATIONS
WHO HAVE NOT REACHED THEIR 18TH BIRTHDAY.

<table>
<thead>
<tr>
<th>Fibrinolytic Therapy Checklist for STEMI</th>
</tr>
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<tbody>
<tr>
<td>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.</td>
</tr>
</tbody>
</table>

**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, provider 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
S. SUDDEN INFANT DEATH SYNDROME (SIDS)

1. Initiate General Patient Care.

2. Presentation
   The unexpected arrest of an apparently healthy infant in which resuscitation is unsuccessful and there is no attributable cause of death. The infant is often discovered by a caretaker in the early morning hours after having been uneventfully laid down to sleep the night before.

3. Treatment
   a) Perform an initial patient assessment, assign a treatment priority, and perform CPR, if indicated.

   RIGOR MORTIS MAY BE PRESENT (SEE PRONOUNCEMENT OF DEATH IN THE FIELD PROTOCOL).

   b) Move patient to the transport unit.

   c) Establish communications and obtain medical direction.

   d) If physician consultation is genuinely unavailable, monitor cardiac rhythm and treat according to the appropriate algorithm(s).

   e) Transport quickly to the closest appropriate facility.

4. Continue General Patient Care.
T. ENVIRONMENTAL EMERGENCIES: COLD EMERGENCIES (FROSTBITE)

1. Initiate General Patient Care.

2. Presentation
   Exposure to cold environment (not necessarily outdoors). Frostbite usually affects the feet first followed by the hands, face, and/or ears. The skin initially appears reddened, then turns mottled, bluish, white and/or gray with continued freezing of the flesh. Pain persists during initial stages followed by numbness.

3. Treatment
   a) Remove patient from cold environment.
   b) Handle potential frostbitten areas gently.
   c) Cover lightly with gauze.
   d) Protect from further heat loss.
   e) Establish IV access with LR.

PEDIATRIC SECTION ON NEXT PAGE
T. ENVIRONMENTAL EMERGENCIES: COLD EMERGENCIES (FROSTBITE) (Continued)

g) Remove patient from cold environment.

h) Handle potential frostbitten areas gently.

i) Cover lightly with gauze.

j) Protect from further heat loss

k) Establish IV/IO access with LR, if appropriate.


4. Continue General Patient Care.
U. ENVIRONMENTAL EMERGENCIES: COLD EMERGENCIES (HYPOTHERMIA)

1. Initiate General Patient Care.

2. Presentation
   a) Mild to moderate hypothermia (90°–95° F)
      Core body temperature (if available) less than 95° F but greater than 90° F. Patient may present with a history of exposure to cold, altered level of consciousness, shivering, stiffness of muscles, stumbling or staggering gait, cool or cold skin, mottled or pale skin, absent or difficult to detect respiratory effort and/or peripheral pulses, respiratory and/or cardiac arrest.
   
   b) Severe hypothermia (less than 90° F)
   
   c) Core body temperature (if available) less than 90° F. Patient may present with any of the symptoms listed above except shivering.

HANDLE ALL HYPOTHERMIC PATIENTS CAREFULLY. ROUGH HANDLING MAY PRECIPITATE CARDIAC ARREST.

IF HYPOTHERMIA IS SUSPECTED AND THE PATIENT DOES NOT HAVE INJURIES INCOMPATIBLE WITH LIFE, THE PATIENT SHOULD BE RESUSCITATED.

3. Treatment
   a) Remove the patient from the cold environment.
   b) Avoid further heat loss by removing wet clothing, replacing with dry blankets and insulating material. Use a thermal type blanket and special attention to covering the patient’s head.
   c) PASSIVELY rewarm patient within a warm environment.
   d) If available, administer warmed oxygen.

ADMINISTER SHOCK(S) WITH THE AED IF INDICATED.

   e) For further AED shocks, obtain medical consultation.
U. ENVIRONMENTAL EMERGENCIES: COLD EMERGENCIES (HYPOTHERMIA)  
(Continued)

f) Monitor EKG closely.

g) Establish IV access with LR, if appropriate.

h) Identify rhythm and treat according to appropriate algorithm.

CONSIDER, WITH MEDICAL CONSULTATION, CONTINUED CARDIOPULMONARY ARREST PROTOCOLS WITH LONGER MEDICATION INTERVALS.

4. Continue General Patient Care.
V. ENVIRONMENTAL EMERGENCIES: DEPRESSURIZATION

1. Initiate General Patient Care.

2. Presentation
   History of SCUBA, breathing in a pressurized environment, or altitude chamber usage with sudden depressurization. Patients may present with any of the following symptoms: fatigue and itching, pain, vertigo, focal weakness, visual disturbances, speech difficulty, marbled rash, numbness, tingling, confusion, seizure, and/or cardiac arrest.

CONSIDER TRANSPORT TO HYPERBARIC MEDICINE SPECIALTY CENTER.

AEROMEDICAL TRANSPORT MAY BE APPROPRIATE FOR PATIENTS WITH BAROTRAUMA.

FOR ADDITIONAL INFORMATION CONCERNING SCUBA INJURIES, CONTACT THE DIVING ALERT NETWORK VIA EMRC 1-800-648-3001.

3. Treatment
   a) Remove patient from water.
   b) Protect patient from and/or treat for hypothermia.
   c) Establish IV access with LR.

4. Continue General Patient Care.
W. ENVIRONMENTAL EMERGENCIES: HAZARDOUS MATERIALS EXPOSURE

1. Initiate General Patient Care.

2. Presentation
   Exposure to a known or unknown hazardous material. Patient may present with a wide array of signs and symptoms due to the variables of substance exposure. Any patient who is exposed to a hazardous material is considered contaminated until the patient is decontaminated thoroughly.

3. Treatment

   **DO NOT ENTER THE SCENE UNLESS PROPERLY TRAINED AND EQUIPPED TO DO SO.**

   PROPER LEVELS OF PERSONAL PROTECTIVE EQUIPMENT (PPE) ARE TO BE WORN BY ALL PERSONNEL, DEPENDING ON THE MATERIAL INVOLVED AND THE ZONE OCCUPIED.

   IT IS ESSENTIAL TO HAVE THE EMS PROVIDER IN CHARGE NOTIFY EMRC AND POTENTIAL RECEIVING HOSPITALS OF A HAZARDOUS MATERIALS EVENT IN WHICH THEY MAY BE CONSULTED. NOTIFY EMRC/RECEIVING HOSPITALS ABOUT THE FIRST PATIENT’S ETA, THE NUMBER OF VICTIMS, AND THE TYPE OF HAZARDOUS MATERIAL AS SOON AS INFORMATION BECOMES AVAILABLE.

   a) Transport of patients even after decontamination will be by ground units only.

   **THE USE OF AEROMEDICAL TRANSPORT IS CONTRAINDICATED FOR ANY POTENTIALLY CONTAMINATED PATIENT**

   b) Triage and decontaminate if indicated.

   c) Protect the patient from the environment and ensure the patient is not/does not become hypothermic.

   d) Establish IV access with LR in a clean area if medication administration is anticipated.

   e) Consider antidote to specific agent if available.

   f) Consider antibiotic specific to agent in mass casualty incident, if available.
W. ENVIRONMENTAL EMERGENCIES: HAZARDOUS MATERIALS EXPOSURE
(Continued)

g) Medical Follow-Up
All public safety personnel who come into close contact with hazardous materials should receive an appropriate medical examination, post-incident, based on information from the designated poison control center. This should be completed within 48 hours of the incident and compared with the findings of any recent pre-incident examination. Personnel who routinely respond to hazardous materials emergencies should have periodic pre-incident examinations. Personnel should be advised of possible latent symptoms at the time of their exams.

4. Continue General Patient Care.
X. ENVIRONMENTAL EMERGENCIES: HEAT-RELATED EMERGENCIES

1. Initiate General Patient Care

2. Presentation
   a) **Heat Cramps**: Moist, cool skin temperature, cramps, normal to slightly elevated temperature
   
   b) **Heat Exhaustion**: Moist, cool skin, cramps, weakness, dizziness, normal to elevated temperature, nausea
   
   c) **Heat Stroke**: Hot, dry skin (25% of patients will still be moist), seizures, altered mental status, dilated pupils, rapid heart rate, or arrhythmia

3. Treatment
   a) Remove patient from hot environment.
   
   b) Cool patient as appropriate.

   **DO NOT GIVE ANYTHING BY MOUTH TO A PATIENT WITH AN ALTERED MENTAL STATUS.**
   
   c) If patient is fully conscious and not nauseated, give electrolyte-rich fluid by mouth if available. *(NEW ’16)*
   
   d) If **heat stroke**, aggressively cool patient and place patient in semi-fowler’s position.

   e) Establish IV access with LR.
   
   f) Administer fluid bolus, if appropriate.
      20 mL/kg of LR IV
      Titrate to a systolic pressure of 100 mmHg.

4. Continue General Patient Care.
Y. ENVIRONMENTAL EMERGENCIES: NEAR-DROWNING

1. Initiate General Patient Care.

2. Presentation
   Confirmed or suspected near drowning, altered level of consciousness, dyspnea, cyanosis, vomiting, seizures, or cardiopulmonary arrest.

3. Treatment
   a) Remove patient from water.

   ALERT
   ABDOMINAL THRUSTS ARE CONTRAINDICATED, UNLESS THE PATIENT HAS A FOREIGN BODY AIRWAY OBSTRUCTION.

   ALL NEAR-DROWNING VICTIMS SHOULD BE TRANSPORTED EVEN IF THEY APPEAR UNINJURED OR APPEAR TO HAVE RECOVERED.

   ENTER WATER ONLY IF TRAINED AND AS A LAST RESORT. (REACH, THROW, ROW, GO WITH ASSISTANCE)

   b) Protect from and/or treat for hypothermia.

   c) Establish IV access with LR.

   d) Identify rhythm and treat according to appropriate algorithm.

   e) Protect from and/or treat for hypothermia.

   f) Establish IV/IO access with LR.

   g) Identify rhythm and treat according to appropriate algorithm.

   IF THE PARENT OR GUARDIAN REFUSES MEDICAL CARE OR TRANSPORT, PROVIDER SHALL CONTACT A PEDIATRIC BASE STATION PHYSICIAN. (NEW ’16)

4. Continue General Patient Care.
Z. ENVIRONMENTAL EMERGENCIES: OVERPRESSURIZATION

1. Initiate General Patient Care.

2. Presentation
   History of SCUBA, breathing in a pressurized environment and altitude cham-
   ber or exposure to blast concussion waves. Patients may present with any of
   the following symptoms: fatigue and itching, pain, vertigo, visual disturbances,
   dyspnea, bleeding from any body orifice, hearing difficulty, speech
   difficulty, numbness, tingling, confusion, seizure, and/or cardiac arrest.

   ASSOCIATED INJURIES MAY MAKE ASSESSMENT AND COMMUNICATION DIFFICULT.
   SYMPTOMS MAY BE SLOW TO PRESENT.
   AEROMEDICAL TRANSPORT MAY BE APPROPRIATE FOR PATIENTS WITH BAROTRAUMA.

   FOR ADDITIONAL INFORMATION CONCERNING SCUBA INJURIES, CONTACT THE DIVING ALERT
   NETWORK VIA EMRC 1-800-648-3001.

3. Treatment
   a) Treat associated trauma.

   b) Establish IV access with LR.

   c) Administer fluid bolus, if appropriate.
      20 mL/kg of LR IV
      Titrate to a systolic pressure of 100 mmHg.

4. Continue General Patient Care.
AA. HYPERBARIC THERAPY PROTOCOL

1. Initiate General Patient Care.

2. Presentation
   a) Patients involved in a closed space fire and/or explosion incident with exposure to products of combustion or toxic gas inhalation are more likely to have carbon monoxide toxicity.
   b) Patients with a recent history of scuba diving exhibiting signs of decompression complications.

3. INDICATIONS FOR REFERRAL TO A HYPERBARIC MEDICINE SPECIALTY CENTER
   a) Patients presenting with altered mental status or nausea with vomiting, seizures, loss of consciousness, or marked dyspnea in the face of suspected carbon monoxide or toxic inhalation with or without minor burns should be considered for transport to the hyperbaric specialty center. Patients in closed space incidents are more likely to manifest these symptoms.
   b) Patients experiencing pain, paralysis, respiratory distress, altered mental status with a history of scuba diving in the last 48 hours.

4. CONTRAINDICATIONS FOR REFERRAL TO A HYPERBARIC MEDICINE SPECIALTY CENTER
   a) Patients who meet the criteria for referral to a burn center.
   b) Patients with injuries that meet the criteria for a trauma center.

   PATIENTS WITH BURNS AND TRAUMA SHOULD BE REFERRED TO THE NEAREST APPROPRIATE TRAUMA CENTER, NOT A BURN CENTER.

   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.

   CHILDREN WHO MEET INCLUSION BASED ON THE TRAUMA DECISION TREE AND WHO HAVE NOT REACHED THEIR 15TH BIRTHDAY SHOULD BE TRANSPORTED TO A PEDIATRIC TRAUMA CENTER.

5. Treatment
   a) Remove patient from toxic environment or eliminate source of toxic gas.
   b) Administer as high a concentration of oxygen as possible.
c) Establish IV access with LR.

(1) If hypoperfusion exists, initiate IV LR fluid therapy 20 mL/kg bolus in unburned area, if possible.
Titrate to a systolic pressure of 100 mmHg.

(2) Obtain medical consultation to initiate an IV in an area of burn, if unable to obtain an IV in unburned area.

(3) Consider additional fluid administration.
(Max 2,000 mL without medical consultation)

___________________________

d) Establish IV/IO access with LR.

(1) If age-related vital signs and patient’s condition indicates hypoperfusion, administer initial fluid bolus of 20 mL/kg LR IV/IO in unburned area, if possible.
If patient’s condition does not improve, administer the second bolus of fluid at 20 mL/kg LR IV/IO.

(2) Obtain medical consultation to initiate an IV in an area of burn, if unable to obtain an IV in unburned area.

6. Transportation
a) Priority 1 Patients (immediate threat to life)
(1) Consider air transportation if the patient will ARRIVE at the appropriate receiving facility more quickly than could be accomplished by ground transportation.

(2) The provider should consider all of the following:
(a) Time for helicopter response
(b) Patient turnover (loading time)
(c) Flight time to appropriate facility
(d) Weather conditions

b) Priority 2 Patients (no immediate threat to life)
Consider air transport if drive time is greater than 30 minutes.

7. Continue General Patient Care.
BB. NAUSEA AND VOMITING

1. Initiate General Patient Care.

2. Presentation
Patients presenting with nausea and/or vomiting due to underlying injury, medical condition, active motion sickness, or medication side effect/complication.

Under certain injury or medical conditions, vomiting or intense nausea can complicate the existing injury or medical condition. Preventative administration of an anti-nausea/anti-emetic should be considered (e.g., penetrating eye injury, high risk for aspiration, side effects of opioid administration). (NEW '16)

3. Treatment
a) Place patient either in position of comfort or in left lateral position if not prevented by spinal protection or packaging.

b) Perform acupressure on P6 point either digitally or with commercial wrist band.

c) Establish IV access with LR, if appropriate.

d) Administer fluid bolus, if appropriate.
20 mL/kg of LR IV
Titrte to a systolic pressure of 100 mmHg.

e) Adult (NEW '16): 8 mg SLOW IV over 2–5 minutes OR 4–8 mg IM OR 8 mg ODT
May repeat once without medical consultation.
For third repeat dose to a patient with maximum total dose of 24 mg.

f) Establish IV access with LR, if appropriate.

g) If age-related vital signs and patient’s condition indicate hypoperfusion, administer initial fluid bolus of 20 mL/kg LR IV/IO.

h) Pediatric (NEW ‘16):
For patients 28 days – 12 years old: 0.1 mg/kg SLOW IV over 2–5 minutes
For patients 13–18 years of age: 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.

4. Continue General Patient Care.
CC. NON-TRAUMATIC SHOCK: HYPOPERFUSION

1. Initiate General Patient Care.

2. Presentation
   The body responds in various ways to a state of inadequate blood flow to meet the oxygen demands of the cells. A patient may exhibit an altered mental status; cool, clammy skin; diaphoresis; dilated pupils; a rapid, weak pulse; shallow, labored respirations; general weakness; and/or a decreasing pulse pressure.

3. Treatment
   a) Continue General Patient Care.

   b) Establish IV access with LR.
      (1) If lungs are clear, administer fluid bolus.
          20 mL/kg of LR IV
          Titrate to a systolic pressure of 100 mmHg.

      (2) If rales are present, administer fluid bolus.
          Maximum of 250 mL of LR IV
          Titrate to a systolic pressure of 100 mmHg.
          More fluid requires medical consultation.

   c) Consider dopamine (2–20 mcg/kg/min).
      Titrate to a systolic pressure of 100 mmHg.

   d) Consider additional fluid administration.
      Maximum Dose 2,000 mL without medical consultation.
e) The pediatric patient may present hemodynamically unstable or with hypoperfusion evidenced by hypotension and signs such as altered mental status, delayed capillary refill greater than 2 seconds, pallor, and/or peripheral cyanosis. Hypotension is defined as a systolic blood pressure less than 60 in neonates (patients birth to 28 days of age) (NEW '16), 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.

f) Continue General Patient Care.

g) Establish IV/IO access with LR.
   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 IV/IO.
   OR
   For volume-sensitive children administer initial fluid bolus of 10 mL/kg LR IV/IO. If patient’s condition does not improve, administer the second bolus of fluid at 10 mL/kg LR IV/IO.
   Volume-sensitive children include: neonates (birth to 28 days) (NEW '16), children with congenital heart disease, chronic lung disease, or chronic renal failure.

h) Third and subsequent fluid boluses at 20 mL/kg IV/IO.

i) Consider dopamine.
   2–20 mcg/kg/min IVP/IO
   Titrate to age-specific vital signs.

4. Continue General Patient Care.
1. Initiate General Patient Care.

2. Presentation
   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.

3. Treatment

   Pre-Arrival Information
   Excessive Bleeding? YES \[\rightarrow\] Absorb Bleeding Treat for Shock
   NO
   Seizures YES \[\rightarrow\] Transport Left Lateral Position Maintain Body Temp. Have Suction Ready (d)
   NO
   Baby’s Head Presents? NO
   YES
   Hand/Foot Presents? YES \[\rightarrow\] Left Lateral Position
   NO
   Feet or Butt Presents? YES \[\rightarrow\] Deliver Body Support Baby’s Wt. Form V to Open Airway
   NO
   Cord Presents? YES \[\rightarrow\] Position Mother Face Down & Butt Up Wrap Cord Keep Moist Insert Gloved Hand to Lift Baby (a,b)
   NO
   Amniotic Sac Broken? YES
   NO \[\rightarrow\] Puncture Sac
   Suction mouth then nose; if meconium present, multiple suction attempts should be made.
   Support Head

(Continued on next page)
(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.

4. Continue General Patient Care.
EE. NEWLY BORN PROTOCOL

1. Initiate General Patient Care.

2. Presentation
   This protocol applies to the infant within the first hour after delivery. (NEW '16)

UNIVERSAL ALGORITHM FOR THE NEWLY BORN FOR BLS (NEW ’16)

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 (NEW ’16)

ALS Care for Rhythm Management & Treatment Medications (ALS Only)
3. **UNIVERSAL ALGORITHM FOR NEWLY BORN FOR ALS (NEW ’16)**

```
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 SpO₂ (a)
  - Evaluate skin color
  - APGAR at 1 min, repeat at 5 mins

**Reassess**

- **IV/IO with LR (f)**
- Epinephrine IV/IO 0.01 mg/kg (1:10,000)
- Neonates (0–28 days)
- Epinephrine ET 0.03 mg/kg (1:10,000) dilute with 1 mL
- Repeat every 3–5 minutes
- Consider intubation (a)
- Consider causes (b)

- **Administer supplemental O₂**
- Monitor IV/IO with LR if poor perfusion (f)

- **Monitor and maintain body temperature**
- Transport

**Reassess**

- **Medical consult**

- **Monitor and maintain body temperature**
- Transport
```
EE. NEWLY BORN PROTOCOL (Continued)

(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 (c, d)
   Hypoglycemia (e) (Threshold for treatment = 30 mg/dL)
   Hypothermia (Warming)
   Hypovolemia (Volume infusion – see “f”, below)

(c) - Premature infants less than 32 weeks gestation will likely require ongoing BVM ventilations due to immature lungs.

(d) - Naloxone 0.1 mg/kg ET/IV/IO

(e) - D10W 2–4 mL/kg IV/IO (D10W is prepared by mixing one part of D50W with four parts LR.)

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

4. Apgar Chart

APGAR Chart

<table>
<thead>
<tr>
<th>SIGN</th>
<th>0</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>MUSCLE TONE (ACTIVITY)</td>
<td>LIMP</td>
<td>SOME FLEXION</td>
<td>ACTIVE, GOOD FLEXION</td>
</tr>
<tr>
<td>PULSE</td>
<td>ABSENT</td>
<td>LESS THAN 100/MIN</td>
<td>GREATER THAN 100/MIN</td>
</tr>
<tr>
<td>REFLEX IRRITABILITY* (GRIMACE)</td>
<td>NO RESPONSE</td>
<td>SOME GRIMACE OR AVOIDANCE</td>
<td>COUGH, CRY OR SNEEZE</td>
</tr>
<tr>
<td>COLOR (APPEARANCE)</td>
<td>BLUE, PALE</td>
<td>PINK BODY, BLUE HANDS/FEET</td>
<td>PINK</td>
</tr>
<tr>
<td>RESPIRATIONS</td>
<td>ABSENT</td>
<td>SLOW/IRREGULAR, INEFFECTIVE</td>
<td>CRYING, RHYTHMIC EFFECTIVE</td>
</tr>
</tbody>
</table>

*Nasal or Oral Suction Catheter Stimulus
**OBSTETRICAL/GYNECOLOGICAL EMERGENCIES: VAGINAL BLEEDING**

1. Initiate General Patient Care.

2. Presentation
   Unusually heavy vaginal bleeding as a result of possible pregnancy, miscarriage, postpartum bleeding, or sexual assault. Patient may exhibit the signs and symptoms of hypoperfusion.

3. Treatment
   a) Place absorbent pads underneath patient.
   b) Treat for hypoperfusion.
   c) If post-partum bleeding, consider uterine massage from pubis toward umbilicus only.
   d) Reconsider ALS.

**PRODUCTS OF CONCEPTION SHOULD BE BROUGHT TO THE HOSPITAL!**

DO NOT PULL CONCEPTUAL PRODUCTS FROM VAGINAL OPENING WITHOUT MEDICAL CONSULTATION!

   e) Establish IV access with LR, if appropriate.
   f) Administer fluid bolus, if appropriate.
      20 mL/kg of LR IV
      Titrate to a systolic pressure of 100 mmHg.
   g) Consider additional fluid administration.
      Maximum dose 2,000 mL without medical consultation.

4. Continue General Patient Care.
GG. OVERDOSE/POISONING: ABSORPTION

1. Initiate General Patient Care.

2. Presentation
   Patient may exhibit any of the following: nausea, vomiting, diarrhea, altered mental status, abdominal pain, rapid heart rate, dyspnea, seizures, arrhythmias, sweating, tearing, defecation, constricted/dilated pupils, rash, or burns to the skin.

3. Treatment
   a) Remove patient from the toxic environment by appropriately trained personnel using proper level PPE.
   b) Identify agent and mechanism of exposure.
   c) Decontaminate as appropriate.
   d) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.
      2 mg intranasal atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)
      Consider additional doses of naloxone. (NEW ’16)
   e) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone
      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)
      Maximum single dose 0.4–2 mg (NEW ’16)
   f) Consider repeating naloxone. (NEW ’16)
   g) Establish IV access with LR in a clean area, if appropriate.
   h) If organophosphate poisoning, consider atropine
      2–4 mg IV or IM every 5–10 minutes.
   i) Consider antidote to specific agent if available.
   j) Consider antibiotic specific to agent in mass casualty incident, if available.
GG. OVERDOSE/POISONING: ABSORPTION (Continued)

k) **If patient has respiratory depression with decreased LOC, constricted pupils, and provider strongly suspects an opioid/narcotic overdose,** Administer naloxone.

28 days to 4 years *(NEW ’16)*: Administer naloxone 0.8–1 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

5 years to adult *(NEW ’16)*: Administer naloxone 2 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

Consider additional doses of naloxone. *(NEW ’16)*

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

m) **Identify agent and mechanism of exposure.**

n) **Decontaminate as appropriate.**

o) **Establish IV access with LR in a clean area, if appropriate.**

p) **If patient has respiratory depression with decreased LOC, constricted pupils, and provider strongly suspects an opioid/narcotic overdose,** Administer naloxone.

0.1 mg/kg SLOW IVP/IO/IN (Divide administration of the IN dose equally between nares to a maximum of 1 mL per nare.)

Maximum single dose 2 mg

ET dose 0.2–0.25 mg/kg

q) **If organophosphate poisoning,** consider atropine

0.02 mg/kg IV/IO or IM every 5–10 minutes.

r) **Consider antidote to specific agent if available.**

s) **Consider antibiotic specific to agent in mass casualty incident,** if available.

4. **Continue General Patient Care.**
HH. OVERDOSE/POISONING: INGESTION

1. Initiate General Patient Care.

2. Presentation
   Patient may exhibit any of the following: nausea, vomiting, diarrhea, altered mental status, abdominal pain, rapid or slow heart rate, dyspnea, seizures, arrhythmias, chemical burns around or inside the mouth, or abnormal breath odors.

3. Treatment

   DO NOT GIVE ANYTHING BY MOUTH WITHOUT MEDICAL CONSULTATION!

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

   a) Identify substance and amount ingested.

   b) Consider activated charcoal without Sorbitol 1 gram/kg PO.

   c) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, administer naloxone.
      2 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

      Consider additional doses of naloxone. (NEW ’16)

   d) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, administer naloxone.
      0.4–2 mg SLOW 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.) (NEW ’16)
      Titrate to adequate respiratory effort.

   e) Establish IV access with LR in a clean area, if appropriate.

   f) If dystonic, extrapyramidal, or mild allergic reaction, consider diphenhydramine.
      25 mg IV or IM
HH. OVERDOSE/POISONING: INGESTION (Continued)

g) If beta-blocker overdose, consider glucagon.
   1 mg every 5 minutes IVP

h) If calcium channel blocker overdose, consider calcium chloride.
   0.5–1 gram SLOW IVP (50 mg/min)

CALCIUM CHLORIDE IS CONTRAINDICATED IN A CALCIUM CHANNEL BLOCKER OVERDOSE PATIENT TAKING DIGOXIN.

i) If organophosphate poisoning, consider atropine.
   2–4 mg IVP or IM every 5–10 minutes

j) If tricyclic overdose, consider sodium bicarbonate.
   1 mEq/kg IVP bolus initially with 0.5 mEq/kg at 10 minute intervals

k) Consider antidote to specific agent if available.

l) Consider antibiotic specific to agent in mass casualty incident, if available.

m) Identify substance and amount ingested.

n) Consider activated charcoal without Sorbitol 1 gram/kg PO.

o) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.

   28 days to 4 years (NEW ‘16): Administer naloxone 0.8–1 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

   5 years to adult (NEW ‘16): Administer naloxone 2 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

   Consider additional doses of naloxone. (NEW ‘16)
HH. OVERDOSE/POISONING: INGESTION (Continued)

p) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, administer naloxone.
   0.1 mg/kg SLOW IVP/IO (titrate)/IN (Divide administration of the IN dose equally between nares to a maximum of 1 mL per nare.) *(NEW ’16)*
   Maximum single dose 2 mg
   ET dose 0.2–0.25 mg/kg

q) Establish IV/IO access with LR in a clean area, if appropriate.

r) If dystonic, extrapyramidal, or mild allergic reaction, consider diphenhydramine 1 mg/kg IVP/IO or IM.
   Maximum single dose 25 mg

s) If beta-blocker overdose, consider glucagon.
   1 mg IVP (25–40 kg)
   0.5 mg IVP (less than 25 kg)
   Every 5 minutes as necessary

CALCIUM CHLORIDE IS CONTRAINDICATED IN A CALCIUM CHANNEL BLOCKER OVERDOSE PATIENT TAKING DIGOXIN.

u) If organophosphate poisoning, consider atropine.
   0.02 mg/kg IVP/IO or IM
   Maximum single dose 2 mg
   May be repeated every 5–10 minutes

v) If tricyclic overdose, consider sodium bicarbonate.
   1 mEq/kg diluted 1:1 SLOW IVP/IO

w) Consider antidote to specific agent if available.

x) Consider antibiotic specific to agent in mass casualty incident, if available.

4. Continue General Patient Care.
II. OVERDOSE/POISONING: INHALATION

1. Initiate General Patient Care.

2. Presentation
   Presentation may vary depending on the concentration and duration of exposure. Symptoms may include, but are not limited to, the following: nausea, vomiting, diarrhea, altered mental status, abnormal skin color, dyspnea, seizures, burns to the respiratory tract, stridor, sooty sputum, known exposure to toxic or irritating gas, sweating, tearing, constricted/dilated pupils, and/or dizziness.

   PULSE OXIMETRY MAY NOT BE ACCURATE FOR TOXIC INHALATION VICTIMS!

   PATIENTS PRESENTING WITH ALTERED MENTAL STATUS OR NAUSEA WITH VOMITING, SEIZURES, LOSS OF CONSCIOUSNESS, OR MARKED DYSPNEA IN THE FACE OF SUSPECTED CARBON MONOXIDE OR TOXIC INHALATION WITH OR WITHOUT MINOR BURNS SHOULD BE CONSIDERED FOR TRANSPORT TO THE HYPERBARIC SPECIALTY CENTER. PATIENTS IN CLOSED SPACE INCIDENTS ARE MORE LIKELY TO MANIFEST THESE SYMPTOMS.

3. Treatment
   a) Remove patient from the toxic environment by appropriately trained personnel using proper level PPE.

   b) Identify agent and mechanism of exposure.

   c) Decontaminate as appropriate.

   d) Consider obtaining blood sample using closed system, if indicated.

   e) Establish IV access with LR in a clean area, if appropriate.

   f) If organophosphate poisoning, consider atropine 2–4 mg IVP or IM every 5–10 minutes.

   g) Consider antidote to specific agent if available.

   h) Consider antibiotic specific to agent in mass casualty incident, if available.
II. OVERDOSE/POISONING: INHALATION (Continued)

   i) Remove patient from the toxic environment by appropriately trained personnel using proper level PPE.

   j) Identify agent and mechanism of exposure.

   k) Decontaminate as appropriate.

   l) Establish IV/IO access with LR in a clean area, if appropriate.

   m) If organophosphate poisoning, consider atropine.  
      0.02 mg/kg IV/IO or IM every 5–10 minutes.

   n) Consider antidote to specific agent if available.

   o) Consider antibiotic specific to agent in mass casualty incident, if available.

4. Continue General Patient Care.
JJ. OVERDOSE/POISONING: INJECTION

1. Initiate General Patient Care.

2. Presentation
Patient may exhibit any of the following: local pain, puncture wounds, reddening skin, local edema, numbness, tingling, nausea, vomiting, diarrhea, altered mental status, seizures, muscle twitching, hypoperfusion, metallic or rubbery taste.

3. Treatment
   a) Identify markings (insects, bites, needlestick, etc.).
   b) Do not apply distal and/or proximal constricting bands for a poisonous snakebite to an extremity. Do remove any jewelry on the affected extremity.
   c) Assist patient experiencing moderate to severe allergic reaction symptoms or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed or EMS service’s epinephrine (1:1,000) 0.3 mg in 0.3 mL IM or patient’s prescribed fast-acting bronchodilator.

   IF THE SNAKE IS DEAD, AND IF IT IS PRACTICAL, DELIVER IT WITH ITS HEAD INTACT. DEAD SNAKES STILL BITE!

   d) Immobilize extremity.
   e) Apply cool packs for relief of pain only.
   f) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.
   2 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

   Consider additional doses of naloxone. (NEW ’16)

   g) Establish IV access with LR; administer 20 mL/kg bolus in uninjured extremity. Titrate to a systolic pressure of 100 mmHg.

   h) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.
   0.4–2 mg SLOW IVP/IO (titrate)/IM/IN (If delivery device is available—divide administration of the dose equally between the nares to a maximum of 1 mL per nare.) (NEW ’16)
   Titrate to adequate respiratory effort.
JJ. OVERDOSE/POISONING: INJECTION (Continued)

i) If organophosphate poisoning, consider atropine.
   2–4 mg IVP or IM every 5–10 minutes.

j) Consider antidote to specific agent if available.

k) Consider antibiotic specific to agent in mass casualty incident, if available.

l) Identify markings (insects, bites, needlestick, etc.).
m) Do not apply distal and/or proximal constricting bands for a poisonous snakebite to an extremity. Do remove any jewelry on the affected extremity.

n) Assist patient experiencing moderate to severe allergic reaction symptoms or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed or EMS service’s epinephrine (1:1,000) 0.15 mg in 0.15 mL IM or patient’s prescribed fast-acting bronchodilator.

o) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.

   28 days to 4 years (NEW ’16): Administer naloxone 0.8–1 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

   5 years to adult (NEW ’16): Administer naloxone 2 mg IN atomizer (Divide administration of the dose equally between the nares to a maximum of 1 mL per nare.)

   Consider additional doses of naloxone. (NEW ’16)

p) Establish IV access with LR; administer 20 mL/kg bolus in uninjured extremity. Titrate to a systolic pressure of 100 mmHg.

q) If patient has respiratory depression with decreased LOC, constricted pupils, and provider suspects an opioid/narcotic overdose, Administer naloxone.

   0.1 mg/kg SLOW IVP/IO (titrate)/IN (Divide administration of the IN dose equally between nares to a maximum of 1 mL per nare.) (NEW ’16)

   Maximum single dose 2 mg
   ET dose 0.2–0.25 mg/kg

r) If organophosphate poisoning, consider atropine.

   0.02 mg/kg IV/IO or IM every 5–10 minutes

s) Consider antidote to specific agent if available.

t) Consider antibiotic specific to agent in mass casualty incident, if available.

4. Continue General Patient Care.
KK. OVERDOSE/POISONING: STIMULANT TOXICITY

1. Initiate General Patient Care.

2. Presentation
   a) Moderate toxicity:
      - Patient exhibits chest pain, hypertension, supraventricular tachycardia, moderate anxiety, respiratory distress, and/or hallucinations
   b) Moderate to severe toxicity:
      - Includes the symptomatology described above along with severe agitation, seizures, and hyperthermia

3. Treatment
   a) Ensure scene is secure and safe from paraphernalia.
   b) Initiate patient care.
   c) Identify amount, route, and time the stimulant was introduced into the body if possible.
   d) Establish IV access with LR. Consider blood draw if possible.
   e) Consider 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)
      If IV unavailable, 2 mg IN or 5 mg IM may be administered.
      IN administration max 1 mL per nare
      Larger doses may be needed to treat stimulant toxicity. Additional doses require medical consultation.
   f) Initiate Chest Pain Protocol and treat accordingly with unstable angina or suspected MI.

SUPRAVENTRICULAR TACHYCARDIA (SVT) MAY RESOLVE WITH THE ADMINISTRATION OF MIDAZOLAM. TREATING SVT DUE TO STIMULANT TOXICITY WITH ADENOSINE WILL NOT WORK SINCE THE SUBSTANCE CAUSING THE SVT WILL STILL BE IN THE SYSTEM AND CAUSE REFRACTORY SVT AFTER THE ADENOSINE HAS WORN OFF.
g) Establish IV access with LR.

h) Consider midazolam 0.1 mg/kg in 2 mg increments SLOW IVP over 1–2 minutes 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 maximum single dose of 5 mg.
   IN administration max 1 mL per nare
   Additional doses require medical consultation.

4. Continue General Patient Care.
LL. EXCITED DELIRIUM SYNDROME (ExDS) (NEW ’16)

1. Initiate General Patient Care
2. Presentation:
   a) Excited delirium syndrome (ExDS) is a potentially life-threatening condition in which a person is in a psychotic and extremely agitation state. Mentally, the subject is unable to process rational thoughts or to focus his/her attention. Physically, the body’s systems are functioning at such a high rate that they begin to shut down and fail. When these two factors occur at the same time, a person can act erratically enough that he/she becomes a danger to self and to the public.
   b) History of present illness often includes:
      (1) Ingestion of a stimulant or hallucinogenic drug
      (2) Drug/alcohol withdrawal
      (3) Psychiatric patient who is off of medication
   c) Signs and symptoms: ExDS is characterized as having a minimum of bizarre and aggressive behavior and one of the above history. The more signs and symptoms the patient exhibits, the more likely the patient is to have ExDS and the higher the risk for complications.
      (1) Tachycardia
      (2) Hypertension
      (3) High body temperature
      (4) Dilated pupil
      (5) Incoherent or nonsensical speech
      (6) Rapid or inconsistent breathing patterns
      (7) Paranoia
      (8) Skin changes:
         (a) Hot/dry skin (in the anticholinergic patient)
         (b) Profuse sweating (in the cocaine/MDMA/methamphetamine patient)
      (9) Shivering
      (10) Inappropriate removal of clothing
      (11) Patients who present after receiving multiple TASER or other less lethal energy by law enforcement

MANY LIFE-THREATENING MEDICAL EMERGENCIES PRESENT WITH SIMILAR SIGNS OF EXDS. EXAMPLES INCLUDE HYPOGLYCEMIA, HYPOXIA, SEIZURES, HEAD INJURIES, AND SEPSIS. EMS PROVIDERS MUST ALWAYS ASSESS FOR THE POSSIBILITY OF OTHER EMERGENCY MEDICAL CAUSES FOR THE PATIENT’S PRESENTATION.

ANOTHER KEY SYMPTOM THAT OCCURS JUST PRIOR TO THE ONSET OF SUDDEN DEATH IN A PATIENT EXPERIENCING EXDS IS “INSTANT TRANQUILITY.” THIS SYMPTOM IS NOTED WHEN A PATIENT WHO HAS BEEN VERY VIOLENT AND AGITATED SUDDENLY BECOMES QUIET AND LETHARGIC. THIS IS A SIGN OF IMMINENT CARDIOPULMONARY ARREST. PATIENTS WHO HAVE UNDERGONE PERIODS OF PROLONGED PHYSICAL STRUGGLE WITHOUT SEDATION WITH BENZODIAZEPINES ARE AT HIGH RISK FOR CARDIAC ARREST. ALL EFFORTS MUST BE MADE BY ALS PROVIDERS TO EXPEDITIOUSLY ADMINISTER MIDAZOLAM TO THE AGITATED AND STRUGGLING EXDS PATIENT.
LL. EXCITED DELIRIUM SYNDROME (ExDS) (Continued)

3. Treatment (BLS)
   a) Ensure scene is secure and safe.
   b) Initiate patient care.
      (1) Obtain a measured temperature, as these patients often have severe hyperthermia.
      (2) If possible, attempt to identify the amount, route, and time of any substance ingested.
      (3) Suspected ExDS patients with evidence of head injury or traumatic mechanism of injury should receive Spinal Protection Protocol.
   c) Patients displaying signs of ExDS do not have medical capacity to refuse care.
      (1) If a suspected ExDS patient resists the delivery of care, ALS resources, EMS supervisors (where available), and law enforcement shall be requested to facilitate the treatment and transport of the patient in a safe and effective manner.
      (2) Patients who exhibit violent behavior shall require a police officer to accompany the patient during transport. Appropriate physical restraint procedures should be utilized per Restraint Protocol.

Patients displaying signs and symptoms of ExDS shall be treated and transported at the Advanced Life Support Level. ALS care and treatment will be guided by the signs and symptoms that the patient is exhibiting, as well as possible occult injuries that may have occurred while the individual was being subdued. The appropriate lifesaving treatment for ExDS is the administration of benzodiazepines, fluid resuscitation, and decreasing hyperthermic core body temperature.

Patients who have received multiple rounds of energy from conducted electrical weapons (including T.A.S.E.R) and are displaying signs of ExDS are at heightened risk for sudden cardiac death. These patients should be treated with benzodiazepines and closely monitored for any evidence of hemodynamic collapse.

d) Establish IV/IO access. Consider blood draw if possible.
e) Administer 20 mL/kg IV fluid bolus LR if tachycardiac and/or hyperthermic.
f) Check glucometer and treat accordingly.
g) Administer midazolam.
   (1) Administer midazolam in 2 mg increments (SLOW IV/IO push over 1–2 minutes).
   (2) May be repeated twice to a maximum total IV dose of 6 mg prior to consult.
   (3) Reduce by 50% for patients 69 years or older.
   (4) If IV/IO unavailable or unsafe to obtain, administer 2 mg increments IN (1 mL per nare)
   (5) If IV/IO/IN administration routes are not possible, administer 5 mg IM.
   (6) Multiple doses may be required to achieve therapeutic effect.
       Additional doses require medical consultation.
LL. **EXCITED DELIRIUM SYNDROME (ExDS) (Continued)**

h) Consider the administration of cold packs to the groin, neck, and axilla for patients displaying evidence of hyperthermia.

**PATIENTS DISPLAYING SIGNS AND SYMPTOMS OF EXDS SHOULD NOT RECEIVE HALDOL AND/OR BENADRYL FOR CHEMICAL RESTRAINT. THESE MEDICATIONS MAY WORSEN AN ANTICHOLINERGIC CRISIS. HALDOL MAY INCREASE THE POSSIBILITY OF CARDIAC DYSRHYTHMIA BY PROLONGING THE QT INTERVAL AND MAY ALSO INCREASE THE CHANCES OF A SEIZURE BY LOWERING THE BODY’S SEIZURE THRESHOLD.**

i) Establish IV/IO access. Consider blood draw if possible.

j) Administer 20 mL/kg IV fluid bolus LR if tachycardiac and/or hyperthermic.

k) Check glucometer and treat accordingly.

l) Administer midazolam.
   
   (1) 0.1 mg/kg in 2 mg increments (SLOW IV/IO push over 1–2 minutes) with a maximal single dose of 2 mg.
   
   (2) If IV/IO unavailable or unsafe to obtain, administer 2 mg increments IN (1 ml per nare)
   
   (3) If IV/IO/IN administration routes are not possible, administer 2 mg IM.
   
   (4) Multiple doses may be required to achieve therapeutic effect with a maximum total dose of 5 mg. Additional doses require medical consultation via a medical consult center.

m) Consider the administration of cold packs to the groin, neck, and axilla for patients displaying evidence of hyperthermia.

4. Continue General Patient Care.
1. Initiate General Patient Care.

2. Presentation
   Pain may be present in many different conditions. Management of pain in the field can help to reduce suffering, make transport easier, and allow the emergency department personnel to initiate specific treatment sooner.

3. Treatment Indications
   a) Measure level of pain. Ask adults 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.png)
MM. PAIN MANAGEMENT (Continued)

b) Allow patient to remain in position of comfort unless contraindicated.
c) Monitor airway and vitals signs every 5 minutes for unstable patients.
d) Mild pain
   (1) Indications for pain management
      (a) Isolated musculoskeletal injuries such as sprains and strains
      (b) Pain related to childhood illnesses such as headache, ear infection, and pharyngitis
   (2) Contraindications for pain management with acetaminophen
      (a) Head injury
      (b) Hypotension
      (c) Administration of acetaminophen or medications containing acetaminophen within the previous four hours
      (d) Inability to swallow or take medications by mouth
      (e) Respiratory distress
      (f) Persistent vomiting
      (g) Known or suspected liver disease
      (h) Allergy to acetaminophen
   (3) Administer acetaminophen to patients ages 3 years and above judged to be in mild to moderate discomfort.
      (2–5 on FACES scale) by child or parent.
      (a) Standard unit dosing of liquid preparation (NEW ’16):
         (i) Less than 2 years of age: Not indicated
         (ii) 2–4 years: Unit dose 160 mg/5 mL
         (iii) 5–12 years: TWO unit doses of 160 mg/5 mL each for a total of 320 mg/10 mL
         (iv) 13–Older: 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 X 2 for a total of 650 mg with sips of water as tolerated by the patient.

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 HIS/HER PAIN AND APPROPRIATE DEFINITIVE TREATMENT.

e) Moderate to severe pain
   (1) Indications for pain management
      (a) The patient reports moderate to severe pain.
      (b) In the provider’s judgment, the patient will benefit from treatment with an opioid analgesic, including patients who are MOLST and/or EMS/DNR patients.
MM. PAIN MANAGEMENT (Continued)

(2) Contraindications for pain management
   (a) Hypersensitivity or known allergy to the medication (morphine or fentanyl)
   (b) Uncorrected respiratory distress or hypoxemia refractory to supplemental oxygen
   (c) Uncorrected hypotension, defined as a persistent systolic pressure less than 90 mmHg

(3) Administer agent
   (a) Morphine IV/IM
      (i) Administer 0.1 mg/kg maximum single dose of 20 mg.
      (ii) 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.
      (iii) Obtain on-line medical direction for additional doses, if required.

   OR

   (b) Fentanyl IV/IN/IM. IN administration max 1 mL per nare
      (i) Administer 1 mcg/kg to a maximum initial dose of 200 mcg.
      (ii) 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.
      (iii) Obtain on-line medical direction for additional doses, if required.

   (c) Morphine IV/IM
      (i) Administer 0.1 mg/kg to a maximum initial dose of 20 mg.
      (ii) 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.
      (iii) Obtain on-line medical direction for additional doses, if required.

   OR

   (d) Fentanyl IV/IN/IM. IN administration max 1 mL per nare
      (i) Administer 1 mcg/kg to a maximum initial dose of 200 mcg. Administer at a rate of 0.5 mcg/kg/min.
      (ii) 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.
      (iii) Obtain on-line medical direction for additional doses, if required.
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 AS NEEDED WITH FLUID BOLUSES. REASSESS VITAL SIGNS AFTER ADMINISTRATION OF THE MEDICATION.

USE OPIOID ANALGESIA WITH CAUTION IN THE MANAGEMENT OF PATIENTS WITH ALTERED MENTAL STATUS. OBSERVE FOR RESPIRATORY DEPRESSION AND TAKE STEPS AS NEEDED TO ENSURE A STABLE AIRWAY.

4. Repeat - Measure level of pain and monitor the patient’s level of pain during subsequent treatment and transport.

5. Transport

PATIENTS RECEIVING A NEW OPIOID (EITHER WITHIN 1 HOUR OR GREATER THAN 1 DOSE WITHIN ANY TIME FRAME) FROM ALS OR BY THE SENDING FACILITY MUST BE TRANSPORTED BY ALS.

6. Continue General Patient Care.
NN. RESPIRATORY DISTRESS: ALLERGIC REACTION (NEW '16)

1. Initiate General Patient Care.

2. Presentation
   a) An allergic reaction is an exaggerated response of the body’s immune system to any substance.
   b) Allergic reactions may range from mild to severe life-threatening anaphylactic reactions.

   (1) MILD: Local swelling and itching at the site

   (2) MODERATE: Hives and/or mild wheezing

   (3) SEVERE: Diffuse wheezing, pharyngeal swelling, dyspnea, hypoperfusion, abnormal skin color, stridor, and/or loss of peripheral pulses

3. Treatment
   a) Assist patient experiencing moderate symptoms or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed or EMS service’s epinephrine (1:1,000) 0.5 mg in 0.5 mL IM or patient’s prescribed fast-acting bronchodilator. (NEW ’16)

   b) Albuterol inhaler (2 puffs) may be repeated once within 30 minutes.

   c) Consider additional doses of epinephrine (1:1,000) 0.5 mg in 0.5 mL IM or prescribed fast-acting bronchodilator. (NEW ’16)

   d) Moderate Distress
      (1) Administer epinephrine 1:1,000.
         0.01 mg/kg IM
         Maximum single dose 0.5 mg
         May repeat every 5 minutes for total of 3 doses for severe reactions.
         Additional doses of epinephrine require medical consultation.

      (2) Establish IV access with LR; administer 20 mL/kg bolus.
         Titrate to a systolic pressure of 100 mmHg.

      (3) Administer diphenhydramine.
         50 mg SLOW IVP or IM
         Additional doses of diphenhydramine require medical consultation.

      (4) Administer a combination of albuterol/Atrovent via nebulizer.
         Albuterol 2.5 mg and Atrovent 500 mcg

      (5) If further treatments are indicated, an additional albuterol-only nebulizer may be given.
NN. RESPIRATORY DISTRESS: ALLERGIC REACTION
(Continued)

e) **Mild Allergic Reaction**

(1) Consider diphenhydramine.
   25 mg SLOW IVP or IM  
   **OR**  
   Consider epinephrine 1:1,000.
   0.01 mg/kg IM  
   Maximum single dose 0.5 mg

(2) Consider additional fluid administration.
   Maximum dose 2,000 mL without medical consultation

f) Assist patient experiencing moderate or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed or EMS service’s epinephrine (1:1,000).
   Less than 5 years of age 0.15 mg and 0.15 mL IM  
   Greater than 5 years of age 0.5 mg and 0.5 mL IM  
   or patient’s prescribed fast-acting bronchodilator. *(NEW ’16)*

  g) Albuterol inhaler (2 puffs) may be repeated once within 30 minutes.

  h) Consider additional doses of epinephrine (1:1,000)
     Less than 5 years of age 0.15 mg and 0.15 mL IM  
     Greater than 5 years of age 0.5 mg and 0.5 mL IM  
     or fast-acting bronchodilator. *(NEW ’16)*

  i) **Moderate Distress**
     Less than 5 years of age 0.15 mg and 0.15 mL IM.
     Greater than 5 years of age 0.5 mg and 0.5 mL IM.
     May repeat every 5 minutes for total of 3 doses for severe reactions.
     Additional doses of epinephrine require medical consultation. *(NEW ’16)*

     (1) Establish IV/IO access with LR.
NN. RESPIRATORY DISTRESS: ALLERGIC REACTION
(Continued)

(2) 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 IV/IO.

(3) Administer diphenhydramine.
   1 mg/kg SLOW IVP/IO or IM
   Maximum single dose 50 mg
   Additional doses of diphenhydramine require medical consultation

(4) a combination of albuterol/Atrovent via nebulizer:
   • For an infant less than 1 year of age, administer albuterol 1.25 mg via nebulizer; Atrovent is contraindicated.
   • For a child 1 year of age or greater, but less than 2 years of age, administer albuterol 1.25 mg and Atrovent 250 mcg. For a patient 2 years of age or greater, administer albuterol 2.5 mg and Atrovent 500 mcg.

(5) If further treatments are indicated, an additional albuterol-only nebulizer may be given.

j) Mild Allergic Reaction

   Consider diphenhydramine.
   1 mg/kg SLOW IVP or IM
   Maximum single dose 25 mg
   OR
   Consider epinephrine 1:1,000.
   0.01 mg/kg IM
   Maximum single dose 0.5 mg

4. Continue General Patient Care.
RESPIRATORY DISTRESS: ANAPHYLAXIS (NEW ’16)

1. Initiate general patient care.

2. Presentation
   a) Anaphylaxis is a condition defined by respiratory and/or cardiovascular collapse resulting from an exaggerated response of the body’s immune system to any substance.
   b) Anaphylaxis is likely to present with one or more of the following:
      (1) Acute onset of illness after exposure to a known allergen with two or more of the following:
         (a) urticaria of skin and/or mucosa or acute swelling/edema (eg, tongue, airway, stridor, lips)
         (b) respiratory compromise
         (c) hypotension
         (d) persistent GI symptoms of vomiting, abdominal pain, or diarrhea
      (2) Acute onset of illness after exposure to a known allergen with hypotension

3. Treatment
   a) Assist patient experiencing moderate to severe symptoms or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed or EMS service’s epinephrine auto-injector or manual (1:1,000) 0.5 mg in 0.5 mL IM or patient’s prescribed fast-acting bronchodilator.
   b) Consider additional doses of epinephrine (1:1,000) 0.5 mg in 0.5 mL IM.
   c) Additional treatments to consider AFTER administration of epinephrine
      (1) Albuterol inhaler (2 puffs) may be repeated once within 30 minutes.
   d) Administer epinephrine
      (1) 0.3 mg IM in the lateral thigh via epinephrine auto-injector or epinephrine (1:1,000) 0.5 mg in 0.5 mL IM
      (2) May repeat every 5 minutes for a total of 3 doses for severe reactions.
      (3) For patients who are in extremis with severe hypotension or impending respiratory failure, consider initiating an epinephrine drip after having administered 3 doses of IM epinephrine.
         (a) Mix 1 mg of epinephrine (either 1:1,000 or 1:10,000) in a 1 liter bag of LR IV/IO. Initiate an infusion with a wide open macro drip titrating to a systolic pressure of greater than 90 mmHg. When drip administered, this will be reported as an exceptional call.
e) Additional treatments to consider AFTER administration of epinephrine
   (1) Albuterol/atrovent via nebulizer: Albuterol 2.5 mg and Atrovent 500 mcg; may repeat albuterol neb 2.5 mg as needed every 5–10 minutes
   (2) Diphenhydramine 50 mg SLOW IVP or IM
   (3) Establish IV access with LR
   (4) Administer 20 mL/kg bolus for hypotension
   (5) Dexamethasone 10 mg IV/IO

f) Assist patient experiencing severe symptoms with the patient’s prescribed or EMS service’s epinephrine
   (1) Less than 5 years of age: 0.15 mg IM in the lateral thigh via epinephrine auto-injector or manual administration 0.15 mg in 0.15 mL IM
   (2) 5 and greater: administer 0.3 mg IM in the lateral thigh via epinephrine auto-injector or manual administration 0.5 mg in 0.5 mL IM
   (3) Consider additional doses of epinephrine (1:1,000) 0.5 mg in 0.5 mL IM.
   (4) Additional treatments to consider AFTER administration of epinephrine
      (a) Albuterol MDI inhaler (2 puffs) may be repeated once within 30 minutes.

   (5) Less than 5 years of age: 0.15 mg IM in the lateral thigh via epinephrine auto-injector or manual administration 0.15 mg in 0.15 mL IM
   (6) 5 and greater: administer 0.3 mg IM in the lateral thigh via epinephrine auto-injector or manual administration 0.5 mg in 0.5 mL IM
   (7) May repeat every 5 minutes for a total of 3 doses for severe reactions.

(5) Dexamethasone 0.5 mg/kg to a maximum 10 mg IV/IO

4. Continue General Patient Care.
PP. RESPIRATORY DISTRESS: ASTHMA/COPD

1. Initiate General Patient Care.

2. Presentation
   Patient may exhibit any of the following: wheezing and/or crackles, abnormal respiratory rate, rapid heart rate, stridor, grunting, cyanosis, mottled skin, altered mental status, nasal flaring, retractions, accessory muscle use, dyspnea, diminished or absent breath sounds, and/or tripod positioning.

3. Treatment
   CONSIDER MEDICAL CONSULTATION FOR PATIENTS GREATER THAN 45 YEARS OF AGE OR PATIENTS WITH A CARDIAC HISTORY.
   a) Assist patient experiencing moderate to severe symptoms or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed fast-acting bronchodilator or prescribed epinephrine auto-injector.
   b) Use of the EMS service’s manual epinephrine (1:1,000) 0.5 mg in 0.5 mL or 0.3 mg via epinephrine auto-injector IM requires medical consultation.
   c) Albuterol inhaler (2 puffs) may be repeated once within 30 minutes.
   d) Consider additional doses of patient’s prescribed fast-acting bronchodilator or manual epinephrine (1:1,000) 0.5 mg in 0.5 mL or 0.3 mg via epinephrine auto-injector IM.
   e) Establish IV access with LR on all Priority 1 or 2 patients and all patients with a history of cardiac disease.
   f) Patients with moderate to severe respiratory distress may require high flow oxygen via non-rebreather mask, continuous positive airway pressure (CPAP), or BVM while receiving medication via nebulizer.
   g) Administer a combination of albuterol/Atrovent via nebulizer. Albuterol 2.5 mg and Atrovent 500 mcg
   h) If further treatments are indicated, an additional albuterol-only nebulizer may be given.
   i) Consider CPAP if patient continues to deteriorate in spite of above nebulized treatments. Continue inline nebulizations.
   j) Consider the administration of epinephrine 1:1,000. 0.3 mg IM in the lateral thigh via epinephrine auto-injector or 0.5 mg in 0.5 mL IM
      May repeat every 5 minutes for a total of 3 doses for severe reactions. OR
   k) Consider the administration of terbutaline. 0.25 mg IM
   l) For moderate to severe exacerbations, consider the administration of dexamethasone 10 mg IV/PO.
m) For moderate to severe exacerbations, consider the administration of magnesium sulfate 1–2 grams in 50–100 mL Lactated Ringer’s or D5W IV/IO over 10–20 minutes.

n) Consider additional doses of epinephrine, albuterol, or terbutaline.

o) Assist patient(s) experiencing moderate to severe symptoms or mild symptoms with a history of life-threatening allergic reaction with the patient’s prescribed or EMS service’s epinephrine (1:1,000) 0.15 mg in 0.15 mL IM or patient’s prescribed fast-acting bronchodilator.

**MEDICAL CONSULTATION IS REQUIRED IF THE PATIENT HAS CONGENITAL HEART OR CHRONIC LUNG DISEASE.**

p) Fast-acting bronchodilator (2 puffs) may be repeated once within 30 minutes.

q) Consider additional doses of patient’s prescribed fast-acting bronchodilator or epinephrine (1:1,000) 0.15 mg in 0.15 mL IM.

r) Patients with moderate to severe respiratory distress may require high flow oxygen via non-rebreather mask, CPAP, or BVM while receiving medication via nebulizer.

s) Administer a combination of albuterol/Atrovent via nebulizer:

   1. For an infant less than 1 year of age, administer albuterol 1.25 mg via nebulizer; Atrovent is contraindicated.
   2. For a child 1 year of age or greater, but less than 2 years of age, administer albuterol 1.25 mg and Atrovent 250 mcg.
   3. For a patient 2 years of age or greater, administer albuterol 2.5 mg and Atrovent 500 mcg.

   t) If further treatments are indicated, an additional albuterol-only nebulizer may be given.

**AND/OR MEDICAL CONSULTATION IS REQUIRED IF THE PATIENT HAS CONGENITAL HEART OR CHRONIC LUNG DISEASE.**

u) Administer epinephrine 1:1,000.

   Less than 5 years of age: 0.15 mg IM in the lateral thigh via epinephrine auto-injector or manual administration 0.15 mg in 0.5 mL IM
   5 years and greater: administer 0.3 mg IM in the lateral thigh via epinephrine auto-injector or manual administration 0.5 mg in 0.5 mL IM
   May repeat every 5 minutes for a total of 3 doses for severe reactions.

v) Consider magnesium sulfate 50 mg/kg IV/IO to a 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.**

w) For moderate to severe exacerbations, consider the administration of dexamethasone 0.5 mg/kg PO/IV up to a maximum dose of 10 mg.

x) Consider additional doses of albuterol or epinephrine.

y) Establish IV/IO access with LR in a clean area, if appropriate.

4. Continue General Patient Care.
1. Initiate General Patient Care.

2. Presentation
   Forms of Croup:
   - **Mild** - Barky cough exhibited 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)

   **IF EPIGLOTTITIS IS SUSPECTED, I.E., DROOLING WITH ABOVE SIGNS AND SYMPTOMS, DO NOT INITIATE THIS PROTOCOL WITHOUT APPROPRIATE MEDICAL DIRECTION.**

3. Treatment
   a) 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).

   b) Place patient on cardiac monitor and record vital signs. (This may be done concurrently with medication administration if patient is unstable.)

   c) MILD: For children exhibiting symptoms of a mild croup presentation, administer dexamethasone 0.5 mg/kg PO up to a maximum dose of 10 mg.

   d) MODERATE: For children who exhibit symptoms of a moderate croup presentation, 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 1:1,000 via nebulizer.

   e) SEVERE: If respiratory distress is so severe that respiratory arrest is imminent:
      (1) First, administer 0.01 mg/kg of epinephrine 1:1,000 IM (max single dose of 0.5 mg).
      (2) Then administer dexamethasone 0.5 mg/kg IV up to a maximum dose of 10 mg AND 2.5 mL of epinephrine 1:1,000 via nebulizer. If IV not established, give IM dexamethasone.

   f) Establish communications with the appropriate facility and obtain medical direction if patient is less than 1 year of age, if additional nebulized epinephrine is needed due to level of distress, or if other interventions or directions are needed.

   **ALL PATIENTS WHO RECEIVE NEBULIZED EPINEPHRINE MUST BE TRANSPORTED BY AN ADVANCED LIFE SUPPORT UNIT TO THE APPROPRIATE MEDICAL FACILITY.**

4. Continue General Patient Care.
RR. RESPIRATORY DISTRESS: PULMONARY EDEMA/CONGESTIVE HEART FAILURE

1. Initiate General Patient Care.

2. Presentation
   Accurate diagnosis of congestive heart failure (CHF)/acute pulmonary edema (APE) as the cause of respiratory distress can be challenging. The most accurate identification of CHF/APE is made using the medical history, risk factors, medications, and physical exam with interpretation of blood pressure.
   CHF/APE is difficult to distinguish, at times, from other respiratory causes. Factors most associated with a short-of-breath patient having CHF include: a history of CHF, exam features of jugular venous distension and EKG evidence of Atrial Fibrillation. CHF patients are commonly on anti-hypertensive and cardiac medicines. Orthopnea (use of additional pillows to prop the head up during sleep), Dyspnea on Exertion and Paroxysmal Nocturnal Dyspnea (PND) are symptoms associated with CHF/APE. Blood pressure is frequently elevated, usually greater than 160/100 but not uncommonly greater than 180/120.

   EMS providers should strongly consider CHF/APE in patients possessing the factors above, presenting with acute respiratory distress, tachypnea, hypoxia, rales, or wheezing and marked hypertension, even in the absence of peripheral edema.

   **ALERT**

   GERIATRIC PATIENTS DEMONSTRATING MARKED HYPERTENSION IN ASSOCIATION WITH SHORTNESS OF BREATH/RESPIRATORY DISTRESS AND WHEEZING (IN THE ABSENCE OF ASTHMA OR INFECTION) STRONGLY SUGGESTS CHF/APE.

   Acute Respiratory Distress from CHF may range from mild to severe, life-threatening cases of Acute Pulmonary Edema. This classification is for patients with Systolic BP greater than 110 mmHg.

   a) Asymptomatic – dyspnea on exertion but no symptoms at rest.
   b) Mild – mild dyspnea at rest, despite O₂ treatment. Able to speak in full sentences.
   c) Moderate – moderate dyspnea. O₂ saturation less than 93% on oxygen. Systolic BP usually greater than 150. Unable to speak in full sentences. Normal mental status.
   d) Severe – severe dyspnea, respiratory failure, hypoxia (O₂ saturation less than 90% on oxygen), diaphoresis, Systolic BP commonly greater than 180. One word sentences, altered consciousness.

   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 (NTG).
RR. RESPIRATORY DISTRESS: PULMONARY EDEMA/CONGESTIVE HEART FAILURE  (Continued)

3. Treatment
   a) Position patient in high Fowler’s position.
   b) Rate the patient’s difficulty breathing on a scale where 0 is “no trouble breathing” and 10 is “the worst trouble breathing.”
   c) Continuous positive airway pressure (CPAP) should be considered for moderate dyspnea and must be implemented in severe dyspnea. (Use early; attempt to administer 3 doses of NTG while setting up, acclimatizing the patient, and applying CPAP.)
   d) Establish IV access with LR.
   e) Identify rhythm and treat according to appropriate algorithm.
   f) For patients with hypertension and moderate to severe symptoms, administer NTG (does not require IV before administration). If SBP drops below 90 mmHg, treat with medical fluid bolus: initial bolus 250–500 mL, may repeat once.
      (1) Asymptomatic - apply oxygen per GPC to maintain O₂ saturation greater than 93%.
      (2) Mild - administer low dose NTG 0.4 mg SL at 3–5 minute intervals to a maximum dose of 1.2 mg.
      (3) Moderate and severe - CPAP is preferred therapy. Until CPAP is applied, administer high dose NTG. Assess BP before each administration.
   g) Consider dopamine 2–20 mcg/kg/min. Titrate to SBP 100 mmHg or medical-consultation-directed BP. IV infusion pump preferred.
MEDICAL CONSULTATION IS REQUIRED IF THE PATIENT HAS CONGENITAL HEART OR CHRONIC LUNG DISEASE.

h) Position patient in semi-Fowler’s position.

i) Establish IV access with LR.

j) Identify rhythm and treat according to appropriate algorithm.

k) Patients with moderate to severe respiratory distress may require high flow oxygen via non-rebreather mask, CPAP, or BVM while receiving medication via nebulizer.

l) Consider albuterol.
   For children less than 2 years, albuterol 1.25 mg
   For children greater than or equal to 2 years, albuterol 2.5 mg

m) Consider morphine.
   0.1 mg/kg SLOW IVP/IO/IM (1–2 mg/min)
   Maximum dose 5 mg

n) Consider dopamine.
   2–20 mcg/kg/min
   Titrate to pediatric medical consultation directed BP.
   IV infusion pump preferred.

4. Continue General Patient Care.

5. Consider transport to the pediatric specialty center that follows patient.
6. UNIVERSAL ALGORITHM FOR PEDIATRIC RESPIRATORY DISTRESS FOR BLS

Assess Responsiveness

Not Responsive
   Assess ABCs
   Go to Universal Algorithm for Pediatric Emergency Cardiac Care for BLS

Responsive
   Assess Breathing
   If respiratory with adequate rate and effort (b): Oxygen 90–100% via nonrebreather 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. 2. 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, 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.
7. **UNIVERSAL ALGORITHM FOR PEDIATRIC RESPIRATORY DISTRESS FOR ALS**

- **Assess Responsiveness**
  - Not Responsive: Assess ABCs
  -Responsive: Assess Breathing
    - If respiratory with adequate rate and effort (b): Oxygen 90–100% via nonrebreather 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
  - 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: Wheezing, 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.
SS.  SEPSIS: ADULT (NEW ’16)

1. Initiate General Patient Care

2. Presentation
   a) 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.
   b) The following patient populations are considered especially high risk for sepsis and should have their temperature measured:
      (1) Altered mental status
      (2) Patients in long term care facilities (nursing home)
      (3) Indwelling catheters
      (4) Oncology patients
      (5) Solid organ transplant
      (6) Bed ridden
   c) For an adult patient, 18 years of age and older, to qualify for this protocol, he/she must have a suspected source of infection AND also present with at least two of the following criteria:
      (1) Temp greater than 100.4°F (38°C) or less than 95.9°F (35.5°C)
      (2) HR greater than 100 bpm
      (3) RR greater than 25 (or EtCO₂ less than or equal to 32 mmHg)
      (4) Hypotension (systolic BP less than 90 mmHg)
      (5) Point of care lactate reading greater than or equal to 4 mmol/L (if available)
   d) Patients with hypotension or altered mental status should be considered to have septic shock and treated and transported rapidly. Patients may be treated under this protocol if they do not meet the above criteria with medical consultation.


3. Treatment
   a) Place patient in position of comfort, or supine if hypotension is present.
   b) Carefully monitor airway and respiratory status, manage as required using the appropriate respiratory distress protocol (especially for patients with suspected pneumonia).
   c) Initiate large bore IV. If large bore IV not available, consider a second peripheral IV with the intention of not causing delay in transport and reserve the use of IO for priority 1 patient. If transport time is greater than 20 minutes and IV access is unsuccessful, consider placement of an IO (especially for septic shock). Consider performing a blood draw if time permits.
SS. SEPSIS: ADULT (Continued)

d) If lungs are clear, and patient does not have a history of CHF or end stage renal failure, provide 2 L of LR wide open. Reassess every 500 mL for shortness of breath, blood pressure, and SpO₂ saturation changes.

OR

e) 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).

f) If available, perform point of care lactate testing (Jurisdictional Pilot Program only).

FLUID LIMITS OR DOSES MAY BE MODIFIED WITH CONSULTATION.

g) Place patient on cardiac monitor and perform 12-lead (do not delay IV therapy or fluid bolus).

h) If hypotension persists after 2 L of LR are provided, consider an additional 2 L of LR (up to a maximum of 30 mL/kg total, including the first 2 L bolus) and/or dopamine 5–20 mcg/kg/min (paramedic only). Titrate to a Mean Arterial Pressure of 65 mmHg or systolic BP of 90 mmHg.
1. Initiate General Patient Care

2. Presentation
   a) 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.
   b) The pediatric septic patient may be difficult to identify due to a poor history or providers may have difficulty identifying an obvious source of infection, as many pediatric sepsis patients are very young children or infants.
   c) The following pediatric patients are at greater risk for sepsis and should have their temperature measured:
      (1) Altered mental status
      (2) Asplenia (spleen removed from treatment of trauma or illness)
      (3) Bone marrow or solid organ transplant
      (4) Cancer patients
      (5) Cerebral Palsy
      (6) Sickle Cell Disease
      (7) Central or indwelling catheters
      (8) Immunodeficiency or immunosuppression
      (9) Bed ridden
      (10) Severe mental delay
   d) For a pediatric patient, who has not reached his/her 18th birthday, to qualify for this protocol, he/she must have a known or suspected infection AND also present with at least three of the Pediatric Sepsis Rule-In Criteria by Age.
   e) A patient not meeting three or more Pediatric Sepsis Rule-In Criteria by Age may be treated under this protocol with Pediatric Base Station approval if sepsis is suspected by the prehospital provider.

**ALERT**

ALTERED MENTAL STATUS REQUIRES GLUCOSE CHECK.

f) 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.
### 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 °C or greater than 100.4 °F</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.

---

**IF A PEDIATRIC PATIENT MEETS THE ABOVE PEDIATRIC SEPSIS RULE-IN CRITERIA BY AGE, THIS PATIENT IS A PRIORITY 1 OR 2 PATIENT AND REQUIRES NOTIFICATION AS “SEPSIS ALERT” TO THE NEAREST APPROPRIATE FACILITY PRIOR TO ARRIVAL.**

**IF A PEDIATRIC PATIENT MEETS ANY OF THE SEPSIS RULE-IN PLUS ONE OR MORE OF THE SHADED AREAS IN THE CHART, CONSULTATION WITH A DESIGNATED PEDIATRIC BASE STATION IS REQUIRED AND SHOULD BE COMBINED WITH LOCAL BASE STATION CONSULTATION.**

3. **Treatment**

   a) Carefully monitor airway and respiratory status. Manage as required using the appropriate respiratory distress protocol (especially for patients with suspected pneumonia).

   b) Place patient on cardiac monitor.

   c) If patient meets the pediatric sepsis rule-in criteria and meets one of the high risk criteria (shaded), initiate IV/IO access and provide a 20 mL/kg bolus of LR IV/IO over 5–20 min. Maximum single dose of 2L.

   d) 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.

   e) For volume-sensitive children administer initial fluid bolus of 10 mL/kg LR IV/IO (max of 250 mL). (Volume-sensitive children are children who need smaller fluid bolus volumes due to special needs including neonates (birth to 28 days), congenital heart diseases, chronic lung disease, or chronic renal failure.)
TT. SEPSIS: PEDIATRIC (Continued)

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

FLUID LIMITS OR DOSES MAY BE MODIFIED WITH CONSULTATION.

g) Dopamine 5–20 mcg/kg/min IV/IO. Titrate to age-specific vital signs.

h) Consider initiation of a second IV. Initiation of second IV shall not delay transport.

i) Patients with fever or known or suspected infection and hypotension or altered mental status should be considered to have septic shock and treated and transported rapidly.

4. Continue General Patient Care.
Support ABCs and provide any needed BLS/ALS interventions

Determine presence of stroke severity using Cincinnati Prehospital Stroke Scale

New onset and positive stroke assessment?

Treat and transport per pt presentation

Determine time patient last known well
Check Glucose
Checklist
Fibrinolytic Therapy
LAMS Assessment

Signs and symptoms consistent with stroke AND onset less than 3.5 hrs.

Transport to nearest Primary Stroke Center

Transport to nearest Stroke Center as Priority 1 and Stroke Alert
UU. STROKE: NEUROLOGICAL EMERGENCIES (Continued)

1. Initiate General Patient Care.

2. Presentation
   Patient may present with numbness or weakness (often on one side only), difficulty speaking, blurred vision, dizziness, or a severe, unexplained headache. May be accompanied by seizures or altered mental status.

The Cincinnati Prehospital Stroke Scale
(Kothari R, et al. Acad Emerg Med 1997; 4:9866-990.)

Facial Droop (have patient show teeth or smile):
- Normal – both sides of face move equally
- Abnormal – one side of 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 (other findings, such as strength of grip, may be helpful)
- Abnormal – one arm does not move or one arm drifts down compared with the other

Abnormal Speech (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 the wrong words, or is unable to speak

If Cincinnati Prehospital Stroke Scale is positive, perform the Los Angeles Motor Scale (LAMS). Relay LAMS score to the receiving hospital during Stroke Alert notification. (NEW ’16)

The Los Angeles Motor Scale (LAMS)

<table>
<thead>
<tr>
<th>Facial droop</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Absent</td>
<td>0</td>
</tr>
<tr>
<td>Present</td>
<td>1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Arm drift</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Absent</td>
<td>0</td>
</tr>
<tr>
<td>Drifts down</td>
<td>1</td>
</tr>
<tr>
<td>Falls rapidly</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Grip strength</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>0</td>
</tr>
<tr>
<td>Weak grip</td>
<td>1</td>
</tr>
<tr>
<td>No grip</td>
<td>2</td>
</tr>
</tbody>
</table>

3. Treatment
   a) Position patient with head elevated at 30 degrees.
   b) Complete the Fibrinolytic Therapy Checklist for Ischemic Stroke.
   c) If the patient is a candidate for fibrinolytic therapy AND can be delivered to the hospital within 3.5 hours* of when patient was last known well, transport the patient to the closest Designated Stroke Center. If there is not one within 30 minutes, then go to the nearest hospital.
UU. STROKE: NEUROLOGICAL EMERGENCIES (Continued)


*STROKE TREATMENTS ARE TIME SENSITIVE. REDUCTION IN TIME OF SYMPTOM ONSET TO TREATMENT IMPROVES OUTCOMES

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.

d) Use glucometer and treat if glucose less than 70 mg/dl.
e) Establish IV access with LR.
f) If the patient is hypotensive, obtain medical consultation.
g) Consider obtaining blood sample using closed system.
h) Do not treat hypertension in the field.

THE CAUSES OF STROKES IN CHILDREN ARE DIFFERENT FROM ADULTS. WHILE STROKES ARE UNCOMMON IN CHILDREN, THEY DO OCCUR AND ARE MOST OFTEN CAUSED BY ONE OF THE FOLLOWING CONDITIONS: CONGENITAL HEART DEFECTS, INFECTIONS (INCLUDING CHICKEN POX, MENINGITIS, OR ENCEPHALITIS), BRAIN INJURY, OR BLOOD DISORDERS (SUCH AS SICKLE CELL DISEASE). STROKES IN CHILDREN ARE MOST OFTEN SEEN IN INFANTS BUT DO OCCUR IN CHILDREN OF ANY AGE.

CHILDREN WITH STROKE SYMPTOMS WHO HAVE NOT REACHED THEIR 18TH BIRTHDAY SHALL BE TREATED UNDER THE PEDIATRIC PROTOCOL. CONSULT WITH A LOCAL BASE STATION AND A PEDIATRIC BASE STATION TO ARRANGE TRANSPORT TO A MARYLAND PEDIATRIC TRAUMA CENTER

i) Administer oxygen at 2–6 liters via nasal cannula (unless hypoxic or in respiratory distress).
j) Position patient with head elevated at 30 degrees.
k) If a child presents with a SUSPECTED stroke (e.g., sickle cell patient), consult with the nearest Pediatric Base Station and local Base Station.
l) Use glucometer and treat accordingly. (See Section IV, Glucometer Protocol.)
UU. STROKE: NEUROLOGICAL EMERGENCIES (Continued)

m) Establish IV access with LR.

n) If the patient is hypotensive, obtain medical consultation.

o) Consider obtaining blood sample using closed system.

p) Do not treat hypertension in the field.

4. Continue General Patient Care.

<table>
<thead>
<tr>
<th>Fibrinolytic Therapy Checklist for Ischemic Stroke</th>
</tr>
</thead>
<tbody>
<tr>
<td>All of the “YES” boxes and all of the “NO” boxes must be checked before a patient should be transported to a “Designated Stroke Center.”</td>
</tr>
</tbody>
</table>

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

<table>
<thead>
<tr>
<th>YES</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ 18 years of age or older</td>
<td>☐ Within 14 days of major surgery or serious trauma</td>
</tr>
<tr>
<td>☐ Signs and symptoms of stroke with neurologic deficit (abnormal Cincinnati Stroke Scale)</td>
<td>☐ History of intracranial hemorrhage</td>
</tr>
<tr>
<td>☐ Patient can be delivered to a Stroke Center within 3.5 hours of when patient was last known well</td>
<td>☐ Witnessed seizure at stroke onset</td>
</tr>
</tbody>
</table>

**EXCLUSION CRITERIA**
(All of the “NO” boxes must be checked)

<table>
<thead>
<tr>
<th>NO</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ Active internal bleeding (e.g., GI or urinary bleeding within the last 21 days)</td>
<td>☐ History of cancer of the brain</td>
</tr>
<tr>
<td>☐ Known bleeding disorder</td>
<td></td>
</tr>
<tr>
<td>☐ Within 3 months of intracranial surgery, serious head trauma, or previous stroke</td>
<td></td>
</tr>
</tbody>
</table>
V. TRAUMA PROTOCOL: BURNS

1. Initiate General Patient Care.
2. Presentation
   a) The primary objectives in burn care by EMS providers are to stop the burning process, establish IV access, avoid hypothermia, and transport patients quickly and safely to a burn center. While patients with large burns (greater than 20%), facial burns, and/or significant smoke inhalation often require endotracheal intubation and mechanical ventilation during their resuscitation and care, airway compromise in the first few hours following a burn is uncommon.
   (1) In adults, prehospital tracheal intubation following acute burns is generally unnecessary unless signs of respiratory failure are present (symptomatic airway obstruction, shock, altered mental status, hypoxemia while receiving supplemental oxygen, or dyspnea, etc.).
   (2) Pediatric airways are smaller than adult airways and require frequent and thorough assessment for signs of respiratory distress. Intubate if necessary.
   b) Burns are the body’s response to injuries to the skin, muscles, bone, nerves, and blood vessels caused by thermal, chemical, electrical, radiation, or light source. Patients may exhibit any of the following: reddening of the skin, deep and intense pain, blisters, mottled appearance, and/or charred black or brown areas with severe or no pain.
   c) Indications for Referral to a Burn Center
      (1) All third degree burns (full thickness)
      (2) Second degree burns (partial thickness) greater than 10% total body surface area
      (3) Burns of the face, hands, feet, major joints, genitalia, or perineum
      (4) Electrical burns, including lightning or contact with high voltage (greater than 120 volts)
      (5) Suspected inhalation injury of toxic smoke (Monitor the patients with suspected inhalation injury for delayed airway obstruction, respiratory distress, or oxygen desaturation as the patient may need emergent airway management.)
      (6) Circumferential burns involving the extremities or torso
      (7) 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 referred to the nearest appropriate trauma center for initial care.

Children who meet burn inclusive criteria who have not reached their 15th birthday should be transported to a pediatric burn center.

Patients presenting with altered mental status or nausea with vomiting, seizures, loss of consciousness, or marked dyspnea in the face of suspected carbon monoxide or toxic inhalation with or without minor burns should be considered for transport to the hyperbaric specialty center. Patients in closed space incidents are more likely to manifest these symptoms.

3. Treatment
   a) Extract the patient from burning vehicles or buildings if safe to do so and move patient to a place of relative safety.
b) Do what is necessary to stop the burning process. If water is used to extinguish the fire, remove wet clothing and dry the patient to prevent hypothermia.

c) Administer oxygen in as high a concentration of oxygen as possible (note: pulse oximetry is not reliable in the presence of carbon monoxide or cyanide exposure).

d) Determine percent of body surface area (BSA) burned and depth.

e) Treat associated trauma.

f) For burns greater than 10%, follow Hypothermia Protocol as well.

g) Remove all rings, bracelets, and other jewelry.

h) Cover wounds appropriately (with a clean sheet or Mylar blanket—sterile dressings no longer recommended).

i) For chemical burns, brush off dry chemical, remove clothing, flush with water.

DO NOT GIVE ANYTHING BY MOUTH.

DO NOT PLACE ICE OR ICE PACKS ON ANY PATIENT WITH BURNS GREATER THAN 5% TOTAL BODY SURFACE AREA.

CONSIDER UTILIZING AEROMEDICAL RESOURCE IF PATIENT IS MORE THAN 30 MINUTES FROM A BURN CENTER/HYPERBARIC MEDICINE SPECIALTY CENTER BY GROUND.

j) Establish IV access with LR, if appropriate.
   (1) 10 mL/kg bolus.
   (2) For shock patients, administer a fluid bolus of 20 mL/kg LR followed by a second 20 mL/kg LR if needed. Titrate to a systolic pressure of 100 mmHg.


l) Consider additional fluid administration. Maximum dose 2,000 mL without medical consultation.

m) Establish IV access with LR, if appropriate.
   (1) 10 mL/kg bolus.
   (2) 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 IV/IO.

n) Third and subsequent fluid boluses at 20 mL/kg LR IV/IO.


4. Continue General Patient Care.
Note: The surface of the patient’s palm equals 1% of his/her body surface area.
1. Initiate General Patient Care.

2. Presentation
   The patient may present with profuse bleeding, avulsions, lacerations, foreign objects, impaled objects, and/or soft tissue damage to the eye(s) and/or surrounding facial areas.

3. Treatment

   NEVER APPLY PRESSURE TO THE EYEBALL OR GLOBE!

   IF THE PATIENT HAS OTHER ASSOCIATED TRAUMA OR BURNS, TRANSPORT THE PATIENT TO THE APPROPRIATE TRAUMA OR BURN CENTER; OTHERWISE, TRANSPORT THE PATIENT TO THE NEAREST EYE TRAUMA CENTER, IF APPROPRIATE.

   DO NOT USE CHEMICAL COLD PACKS ON THE FACE.

   a) **Foreign objects NOT embedded in the eye(s):** Flush with copious amounts of water (preferably sterile), normal saline, or LR from the bridge of the nose outward.

   b) **Injury to orbits (area around the eye):** Consider head stabilization and Spinal Protection Protocol.

   c) **Lacerations/injuries to the eyeball or globe:** Shield affected eyeball 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.

   d) **Impaled objects:** Stabilize object, shield affected eyeball, and dress other eye to reduce movement.

   e) Establish IV access with LR, if appropriate.

g) **Foreign objects NOT embedded in the eye(s):** Flush with copious amounts of water (preferably sterile), normal saline, or LR from the bridge of the nose outward.

h) **Injury to orbits (area around the eye):** Consider head stabilization and Spinal Protection Protocol.

i) **Lacerations/injuries to the eyeball or globe:** Shield affected eyeball 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.

j) **Impaled objects:** Stabilize object, shield affected eyeball, and dress other eye to reduce movement.

k) Establish IV/IO access with LR, if appropriate.


4. **Continue General Patient Care.**
XX. TRAUMA PROTOCOL: HAND/UPPER/LOWER EXTREMITY TRAUMA

1. Initiate General Patient Care.

2. Presentation
   a) Patient may exhibit injuries to skeletal or soft tissue components of the hand or upper extremity at or below the level of the mid-humerus, including complete or incomplete amputations of the elements of the hand or upper extremity, crush or degloving injuries, and other trauma resulting in loss of perfusion or suspected nerve injury (e.g., compartment syndrome).

   Upper Extremity
   b) Indications for:
      Referral of adult patients to the Curtis National Hand Center at Union Memorial Hospital or
      Referral of pediatric patients to the nearest Pediatric Trauma Center (children who have not reached their 15th birthday)
      Stable patients with an isolated upper extremity injury at or below the mid-humerus

   (Hand Center and/or nearest appropriate trauma center)
   (1) Complete or incomplete hand or upper extremity amputation
   (2) Partial or complete finger or thumb amputation
   (3) Degloving, crushing, or devascularization injuries of hand or upper extremity
   (4) High-pressure injection injuries to hand or upper extremity
   (5) 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
   c) Indications for Referral to Pediatric or Adult Trauma Center: Patient may exhibit injuries to skeletal or soft tissue components with complete or incomplete amputation of ankle/foot lower extremity, complicated nerve, vessel, or compartment syndrome (excessive swelling and pain of extremity with possible evolving nerve deficit injury).

   LIFE BEFORE LIMB.

   TOE INJURIES FROM LAWN MOWER ARE NOT CANDIDATES FOR REIMPLANTATION AND PATIENTS SHOULD GO TO THEIR LOCAL MEDICAL FACILITY.

   d) Contraindications for referral to a Hand Center
      (1) Patients with unstable or abnormal vital signs
      (2) Patients with major and/or multiple system trauma

   e) Contraindication for referral to Pediatric or Adult Trauma Center
      Patients with toe amputation (partial or complete)
XX. TRAUMA PROTOCOL: HAND/UPPER/LOWER EXTREMITY TRAUMA (Continued)

3. Treatment

a) Package amputated extremity in sealed plastic bag (keep dry) and place on top of ice to keep cool. DO NOT FREEZE.

DO NOT SUBMERGE IN WATER OR FREEZE AMPUTATED PART.

USE TIME, DISTANCE, WEATHER, AND PROXIMITY TO DESIGNATED TRAUMA CENTER TO DETERMINE MODE OF TRANSPORT. IF ESTIMATED TRANSPORT TIME TO DESIGNATED HAND CENTER IS LESS THAN 30 MINUTES, USE GROUND TRANSPORT.

b) Establish IV access with LR, if appropriate.

c) Administer fluid bolus, if appropriate.
   20 mL/kg of LR IV
   Titrate to a systolic pressure of 100 mmHg.


e) Consider additional fluid administration.
   Maximum dose 2,000 mL without medical consultation

f) Establish IV/IO access with LR, if appropriate.

g) 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 IV/IO.

h) Third and subsequent fluid boluses at 20 mL/kg LR IV/IO.


4. Continue General Patient Care.
YY. TRAUMA PROTOCOL: MULTIPLE/SEVERE TRAUMA

1. Initiate General Patient Care.

2. Presentation
   The patient may present with hypovolemic or neurogenic shock, hypotension, hypertension, rapid or slow heart rate, unequal pupils, shallow or absent respirations, decreased distal pulses, decreased motor and sensory function in extremities, internal or external bleeding, fractures, or lacerations.

   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.

   CHILDREN WHO MEET INCLUSION BASED ON THE TRAUMA DECISION TREE AND WHO HAVE NOT REACHED THEIR 15TH BIRTHDAY SHOULD BE TRANSPORTED TO A PEDIATRIC TRAUMA CENTER.

3. Treatment
   a) Apply Spinal Protection Protocol for blunt trauma patients. Patients with isolated penetrating trauma should not have spinal immobilization performed.

   b) Control bleeding and immobilize patient, if blunt mechanism indicates. Spinal immobilization should not be performed on patients with isolated penetrating mechanism. If mechanism includes both blunt and penetrating trauma, apply Spinal Protection Protocol. Backboard may be used for patient transfer maneuvers.

   c) Hyperventilate the head-injured patient as follows (NEW '16):
      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
      (1) Who has signs of herniation such as unequal pupils, posturing, or paralysis or
      (2) Who is manifesting a rapidly decreasing GCS or
      (3) With on-line medical consultation

   d) Consider pelvic stabilization technique if indicated.

   e) Establish IV access with LR; administer 20 mL/kg bolus. Titrate to a systolic pressure of 100 mmHg.

   f) Consider additional fluid administration.
      Maximum dose 2,000 mL without medical consultation
g) Apply Spinal Protection Protocol for blunt trauma patients. Patients with isolated penetrating trauma should not have spinal immobilization performed.

h) Control bleeding and immobilize patient, if blunt mechanism indicates. Spinal immobilization should not be performed on patients with isolated penetrating mechanism. If mechanism includes both blunt and penetrating trauma, apply Spinal Protection Protocol. Backboard may be used for patient transfer maneuvers.

i) Hyperventilate the head-injured patient as follows (NEW ’16):
   - 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

j) Who has signs of herniation such as unequal pupils, posturing, or paralysis or
   - (1) Who is manifesting a rapidly decreasing GCS or
   - (2) With on-line medical consultation

k) Establish IV/IO access with LR.

l) 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.

m) Third and subsequent fluid boluses at 20 mL/kg LR IV/IO.

4. Continue General Patient Care.
### GLASGOW COMA SCALE

#### Eye Opening

| Spontaneously | 4 |
| To Voice | 3 |
| To Pain | 2 |
| No Response | 1 |

#### Motor Response

| To Verbal Command | Obeys | 6 |
| To Painful Stimulus | Localizes Pain | 5 |
| Flexion - Withdraw | 4 |
| Flexion - Abnormal | 3 |
| Extension | 2 |
| No Response | 1 |

#### Verbal Response

| Less than 2 years old | 5 |
| 2–5 years old | APPROPRIATE WORDS |
| Greater than 5 years old | ORIENTED AND CONVERSES |
| 5 |
| 4 |
| 3 |
| 2 |
| 1 |

| SMILES/COOS/cries | CRIES/SCREAMS |
| CRIES | INAPPROPRIATE WORDS |
| INAPPROPRIATE CRIES/SCREAMS | DISORIENTED AND CONVERSES |
| GRUNTS | INAPPROPRIATE WORDS |
| NO RESPONSE | INCOMPREHENSIBLE SOUNDS |
| NO RESPONSE | 2 |
| NO RESPONSE | 1 |

#### Glasgow Coma Score

| Total (3–15) | 165 |
| Edition Date July 1, 2016 | 165 |
ZZ.  TRAUMA PROTOCOL: SEXUAL ASSAULT

1. Initiate General Patient Care.

2. Presentation
   Patient may present with no overt evidence of trauma, or may present with bruising, bleeding, or associated physical and/or emotional trauma.

ALL HEALTH CARE PROVIDERS 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.

3. Treatment
   a) Patient may feel more comfortable talking to someone of the same gender.
   b) Maintain non-judgmental, but caring attitude.
   c) Preserve crime scene and clothing articles, if practical.
   d) Maintain strict confidentiality.
   e) Do not perform a genital examination.
   f) Dress wounds (do not attempt to clean).
   g) Discourage any self-treatment (shower, washing, changing clothes).
   h) Treat injuries according to presentation.

4. Continue General Patient Care.
AAA. TRAUMA PROTOCOL: SPINAL PROTECTION

1. Initiate General Patient Care.

2. Presentation
   Indications for initiating spinal protection:
   “Spinal protection” refers to the act of protecting the spinal cord from further injury.
   “Spinal immobilization” is 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.
   a) 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 protection.
      (1) Midline spinal pain, tenderness, or deformity
      (2) Signs and symptoms of new paraplegia or quadriplegia
      (3) Focal neurological deficit
      (4) Altered mental status or disorientation
      (5) 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 Specialty Spinal Center.
   (6) 15 years of age or older AND
   (7) Signs and symptoms of new paraplegia or quadriplegia in the presence of trauma AND
   (8) Patent airway AND
   (9) Hemodynamically stable
      If considering referral to Adult Specialty Spinal Center, consult with both the nearest Trauma Center and the Adult Spinal Specialty Center, when possible.

3. Treatment
   a) Initiate General Patient Care.
   b) All patients meeting the Spinal Protection Protocol shall have manual in-line cervical spine stabilization and application of a correctly sized cervical collar.
   c) Minimize flexion, extension, and rotation of the spinal column.
   d) Patients meeting the Spinal Protection Protocol who are with neurological deficit, or not able to ambulate on their own accord, shall be immobilized with cervical collar and a backboard.
AAA. TRAUMA PROTOCOL: SPINAL PROTECTION (Continued)

e) The following patients only need application of a cervical collar and do not need to be placed in full immobilization with a backboard:
   (1) Patients who are found by EMS providers to be standing or ambulatory,
   (2) 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, or rotation) of the spinal column, and
   (3) Patients who do not have evidence of a neurological deficit.

f) Patients who are placed in a cervical collar without a need for immobilization on a backboard should be assisted in minimal movement to the EMS stretcher and allowed to lie down supine on their own accord.

g) Patients meeting Spinal Protection Protocol and not requiring immobilization with a backboard should be secured to the EMS stretcher in a supine position with the head elevated at 30 degrees.

h) Backboards may be used for patient extrication and patient transfer for patients not meeting Spinal Protection Protocol; however, other devices are preferred (e.g., sheet, Reeves sleeve, or scoop stretcher).

i) If the backboard is used for extrication from the scene to an ambulance, the patient should be removed from the backboard as soon as possible. The stretcher mattress will provide support in place of the backboard.

j) Interfacility transport patients who have already been removed from a backboard should not be placed back on the backboard prior to transport.

k) Helmet Removal
   (1) 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.
   (2) If patient is wearing helmet and no shoulder pads, removal of the helmet is indicated.
   (3) 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.

l) Patients found with backboard applied before EMS arrival
   (1) If EMS providers find patient immobilized on a backboard applied prior to arrival, the principles of the Spinal Protection Protocol still apply.
m) Establish IV/IO access with LR, if appropriate.

n) Administer fluid bolus, if appropriate.
   20 mL/kg of LR IV
   Titrate to a systolic blood pressure of 100 mmHg.

o) Consider dopamine.
   2–20 mcg/kg/min IV/IO
   Titrate to a systolic blood pressure of 100 mmHg.

p) Consider additional fluid administration.
   Maximum dose 2,000 mL without medical consultation.

q) 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 pro-
   tection.
   (1) Midline spinal pain, tenderness, or deformity
   (2) Signs and symptoms of new paraplegia or quadriplegia
   (3) Focal neurological deficit
   (4) Altered mental status or disorientation
   (5) Distracting injury
   (6) Neck pain or torticollis
   (7) High impact diving incident or high risk motor vehicle crash (i.e., head on col-
       lision, rollover, ejected from the vehicle, death in the same crash, or speed
       greater than 55 mph)
   (8) Substantial torso injury
   (9) Conditions predisposing to spine injury

Indications for referral to a Pediatric Trauma Center:
   (10) Patient is less than 15 years of age AND
   (11) Signs and symptoms of new paraplegia or quadriplegia in the presence of
        trauma AND
   (12) Patent airway AND
   (13) Hemodynamically stable

Consult with nearest Trauma Center and, when possible, the nearest Pediatric
Trauma Center.

r) Initiate General Patient Care.

s) All patients meeting the Spinal Protection Protocol shall have manual in-line cer-
   vical spine stabilization and application of a correctly sized cervical collar.
AAA. TRAUMA PROTOCOL: SPINAL PROTECTION (Continued)

t) Minimize flexion, extension, and rotation of the spinal column.

u) Patients meeting the Spinal Protection Protocol who are with neurological deficit, not able to ambulate on their own accord, or who are unable to respond during assessment shall be immobilized with cervical collar and a backboard.

v) The following patients only need application of a cervical collar and do not need to be placed in full immobilization with a backboard:
   (1) Patients who are found by EMS providers to be standing or ambulatory
   (2) 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, or rotation) of the spinal column, and
   (3) Patients who do not have evidence of a neurological deficit.

w) Patients who are placed in a cervical collar without a need for immobilization on a backboard should be assisted in minimal movement to the EMS stretcher and allowed to lie down supine on their own accord.

x) Patients meeting Spinal Protection Protocol and not requiring immobilization with a backboard should be secured to the EMS stretcher in a supine position with the head elevated at 30 degrees.

y) Backboards may be used for patient extrication and patient transfer for patients not meeting Spinal Protection Protocol; however, other devices are preferred (e.g., sheet, Reeves sleeve, or scoop stretcher).

z) If the backboard is used for extrication from the scene to an ambulance, the patient should be removed from the backboard as soon as possible. The stretcher mattress will provide support in place of the backboard.

aa) Interfacility transport patients who have already been removed from a backboard should not be placed back on the backboard prior to transport.

bb) Helmet Removal
   (1) 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.
   (2) If patient is wearing helmet and no shoulder pads, removal of the helmet is indicated.
   (3) 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.
AAA. TRAUMA PROTOCOL: SPINAL PROTECTION (Continued)

cc) Patients found with backboard applied before EMS arrival
   (1) If EMS providers find patient immobilized on a backboard applied prior to
       arrival, the principles of the Spinal Protection Protocol still apply.

dd) Establish IV/IO access with LR, if appropriate.

ee) Administer fluid bolus, if appropriate.
   20 mL/kg of LR IV
   Titrate to a systolic blood pressure of 100 mmHg.

ff) Consider dopamine.
   2–20 mcg/kg/min IV/IO
   Titrate to a systolic blood pressure of 100 mmHg.

gg) Consider additional fluid administration.
   Maximum dose 2,000 mL without medical consultation

4. Continue General Patient Care.
BBB. TRAUMA PROTOCOL: TRAUMA ARREST

1. Initiate General Patient Care.

2. Presentation
   Early cardiac arrest secondary to trauma is usually due to severe hypoxia, neurologic injury, or massive hemorrhage. The patient is unresponsive, pulseless, and apneic.

3. Treatment
   a) Rapid assessment and extrication
   b) Determine if patient meets the criteria for termination of resuscitation for a patient in traumatic arrest. If patient meets criteria, discontinue resuscitation. If criteria are not met, continue resuscitation.
   c) Perform spinal immobilization for blunt trauma patients only. Patients with isolated penetrating trauma should not have spinal immobilization performed. If mechanism includes both blunt and penetrating trauma, perform spinal immobilization.
   d) CPR
   e) Consider AED if arrest is believed to be medical in nature and the patient meets the criteria.

   IF TRAUMATIC ARREST IS SECONDARY TO PENETRATING TRAUMA, PATIENT IS IN A RHYTHM OTHER THAN ASYSTOLE, AND THE TRAUMA CENTER IS WITHIN 15 MINUTES, TRANSPORT THE PATIENT. IF TRANSPORT TIME EXCEEDS 15 MINUTES, CONSULT.

   f) Establish IV access with LR, if appropriate.
   g) Administer fluid bolus, if appropriate.
      20 mL/kg of LR IV
      Titrate to a systolic pressure of 100 mmHg.
   h) Identify rhythm and refer to appropriate algorithm.
   i) If traumatic arrest is suspected due to multi-system blunt trauma, or due to penetrating neck, chest, or abdominal trauma, bilateral needle decompressions should be performed. Once catheters are placed do not remove.
j) Rapid assessment and extrication

k) Perform spinal immobilization for blunt trauma patients only. Patients with isolated penetrating trauma should not have spinal immobilization performed. If mechanism includes both blunt and penetrating trauma, perform spinal immobilization.

l) CPR

m) Consider AED if arrest is believed to be medical in nature. (See Section IV, AED.)

A PATIENT IN CARDIOPULMONARY ARREST SECONDARY TO TRAUMA SHOULD BE TAKEN TO THE NEAREST APPROPRIATE PEDIATRIC TRAUMA CENTER. CONSIDERATION SHOULD BE GIVEN TO TRANSPORTING THE PATIENT TO THE NEAREST EMERGENCY DEPARTMENT OR ADULT TRAUMA CENTER IF THE PEDIATRIC TRAUMA CENTER IS MORE THAN 10 MINUTES ADDITIONAL TRANSPORT TIME!

n) Establish IV/IO access with LR.

o) 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 IV/IO.

p) If traumatic arrest is suspected due to multi-system blunt trauma, or due to penetrating neck, chest, or abdominal trauma, bilateral needle decompressions should be performed. Once catheters are placed do not remove.

4. Continue General Patient Care.
### CCC. TRAUMA DECISION TREE

When in doubt, take patient to an appropriate Trauma Center

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

<table>
<thead>
<tr>
<th>Category Alpha</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>❑ GCS less than or equal to 13</td>
<td></td>
<td></td>
</tr>
<tr>
<td>❑ Systolic BP less than 90 mmHg (Adult) less than 60 mmHg (Peds)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>❑ Respiratory rate less than 10 or greater than 29 (less than 20 in infant age less than one year) or need for ventilatory support</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**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.

<table>
<thead>
<tr>
<th>Category Bravo</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>❑ 2 or more proximal long-bone fractures</td>
<td>❑ Crushed, degloved, mangled, or pulseless extremity</td>
<td>❑ Pelvic fracture</td>
</tr>
<tr>
<td>❑ Amputation proximal to wrist or ankle</td>
<td>❑ Open or depressed skull fracture</td>
<td>❑ Paralysis (spine)</td>
</tr>
<tr>
<td>❑ Chest wall instability or deformity (e.g., flail chest)</td>
<td>❑ Penetrating injuries to head, neck, torso, or extremities proximal to elbow and knee</td>
<td></td>
</tr>
</tbody>
</table>

**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.

<table>
<thead>
<tr>
<th>Category Charlie</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>❑ High Risk Auto Crash</td>
<td>❑ Rollover without restraint</td>
<td>❑ Auto v. pedestrian/bicyclist thrown, run over, or with significant (greater than 20 mph) impact</td>
</tr>
<tr>
<td>❑ Intrusion (including roof) greater than 12 in. occupant site; greater than 18 in. any site</td>
<td>❑ Auto v. pedestrian/bicyclist thrown, run over, or with significant (greater than 20 mph) impact</td>
<td>❑ Motorcycle crash greater than 20 mph</td>
</tr>
<tr>
<td>❑ Ejection (partial or complete) from vehicle</td>
<td>❑ Death in same passenger compartment</td>
<td>❑ Vehicle telemetry data consistent with high risk of injury</td>
</tr>
<tr>
<td>❑ Death in same passenger compartment</td>
<td>❑ Vehicle telemetry data consistent with high risk of injury</td>
<td></td>
</tr>
<tr>
<td>❑ Vehicle telemetry data consistent with high risk of injury</td>
<td>❑ Exposure to blast or explosion</td>
<td></td>
</tr>
</tbody>
</table>

**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.

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

**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.
2. 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 (NEW '16)
   (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 (NEW '16)
   (a) Adult/Adolescent (greater than 13 years of age) Heart rate less than 60
   (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 provider 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.
      Providers may manually decompress the stomach and/or open an existing gastric tube or button.

f) SUGGESTED SIZES FOR RESUSCITATION MASKS (NEW ’16)

<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 (NEW ’16)

<table>
<thead>
<tr>
<th>Age/Weight</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>Adult/Adolescent</td>
<td>Adult 1,000–1,200 mL</td>
</tr>
</tbody>
</table>
14. ELECTRICAL THERAPY: AUTOMATED EXTERNAL DEFIBRILLATION (AED)

a) INDICATIONS

Sudden cardiac arrest (patients with no pulse and not breathing).
Birth - 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 - 8 years of age AED with pediatric capability using the pediatric pad is preferred over an adult AED.
Child 8 years of age or greater Adult AED

b) CONTRAINDICATIONS

Patient exhibiting signs of life

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 patient regains a pulse, the AED prompt states “no shock advised,” or ALS arrives.
(3) No more than 3 stacked shocks (9) or 4 single new device shocks via AED without medical consultation.

(4) 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 and transport.

(5) If shock is not indicated and patient regains pulse, treat per Altered Mental Status 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 provider contact, and the triggering event.

(2) If using an AED with EKG strip recorder, generate 2 recordings.

(3) Give one to the ALS provider 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.
18. GO-TEAM ACTIVATION

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 providers 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.
21. HIGH PERFORMANCE CPR (HPCPR) (NEW '16)

a) PURPOSE
To improve the overall survival rate of sudden out-of-hospital cardiac arrest patients within the State of 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 is based on research and practices being used in many other high performance EMS systems across the country.

b) INDICATIONS
Patients in cardiac arrest who have reached their 8th birthday.

c) CONTRAINDICATIONS
Patients meeting the criteria for Pronouncement of Death in the Field Protocol
Patients who have not reached their 8th birthday (NEW '16)

d) POTENTIAL ADVERSE EFFECTS/COMPLICATIONS
None

e) PRECAUTIONS
None

f) PROCEDURE FOR HIGH PERFORMANCE CPR
The first provider at the patient's side will assess and initiate compressions.

(1) 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 two inches allowing for complete recoil of the chest after each compression. Compressions should be accomplished with equal time given for the down and up motion and achieve a rate of 100–120/min.

(2) 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 phase to assess pulses and/or defibrillate will be limited to less than 10 seconds.
(3) **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 defibrillator 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 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.

(4) **Ventilations** - Ventilations will be performed without stopping chest compressions. One ventilation will be given every 10th compression during recoil (upstroke). Once an advanced airway is in place, ventilations will be asynchronous with compressions (1 ventilation every 6 to 8 seconds). High performance, continuous compressions remain the priority. Ensure ventilations are adequate with BVM attached to 100% oxygen. Providers will not interrupt compressions to obtain an advanced airway.

(5) **Advanced Life Support** - ALS providers 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 a 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.
(6) **Return of Spontaneous Circulation (ROSC)** - Implement the Return of Spontaneous Circulation and Neuroprotective Induced Hypothermia Protocol as indicated and transport to the closest Cardiac Interventional Center. Following stabilization, post-ROSC, obtain a 12-lead EKG.

g) **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):

![Diagram]

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 provider crew:**
Provider 1 – Chest compressions
Provider 2 – Ventilate, attach/operate AED/defibrillator, assume crew leader responsibilities (providers rotate positions every two minutes)
*Roles remain the same even if providers are ALS equipped*

**3 provider crew:**
Provider 1 – Chest compressions
Provider 2 – Ventilate
Provider 3 – Crew Leader, attach/operate AED/defibrillator
(Providers 1 and 2 rotate every two minutes)
*Roles remain the same even if providers are ALS equipped*
4 provider crew:
Provider 1 – Chest compressions
Provider 2 – Ventilate
Provider 3 – Attach/operate AED/defibrillator
Provider 4 – Crew leader
(Providers 1, 2, and 3 rotate every two minutes)

** Once first two roles have begun treatment, ALS providers will establish IV/IO and administer medications.

Greater than 4 providers - Utilize the same initial assignments as the four provider 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 providers 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.
23. INTRAVENOUS MAINTENANCE THERAPY FOR EMT

a) Provider-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) (NEW ’16)

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 PROVIDER.

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 provider.
UNDER NO CIRCUMSTANCES SHALL THE EMT PROVIDER ATTEMPT TO MAKE ANY ADJUSTMENTS TO IV INFUSION PUMPS, NOR SHOULD THE EMT PROVIDER 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 provider 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.

(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 provider-controlled IV solution

(b) Type of patient-controlled IV solution

(c) Type of patient-controlled IV medication

(d) Volume administered

(e) Complications encountered
25. PATIENT-INITIATED REFUSAL OF EMS

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 provider-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. Providers 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 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 PROVIDER TO EXAMINE A MINOR PATIENT TO DETERMINE THE SEVERITY OF THE ILLNESS OR INJURY, THEN CONSIDER CONTACTING LAW ENFORCEMENT FOR ASSISTANCE. CONSIDER 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
   (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 consciousness, 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 Patient-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 providers 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 his/her care.

(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 provider 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 his/her condition and accept transportation.

(f) If the EMS provider is unsure whether the patient has adequate ability to make medical decisions, he/she should seek medical consultation.

(g) At any time the EMS provider identifies patient conditions that indicate that the patient should be transported to a hospital, and the patient is refusing transport, then the provider 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) 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, “III E. Behavioral Emergencies” 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 provider is unsure if the patient is medically capable of refusing transport.
2. The provider disagrees with the patient’s decision to refuse transport due to unstable vital signs, clinical factors uncovered by the assessment, or the provider’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.
4. Minor patients: No parent, guardian, or authorized decision maker is available or the provider 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, providers 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 provider 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 jurisdiction’s documentation (Medical Incident Report, MAIS form, or jurisdictional equivalent.)
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 his/her 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?
   - Place?
   - Time?
   - Situation?
   
2. Altered level of consciousness?
3. Alcohol or drug ingestion by history or exam with:
   - a. Slurred speech?
   - b. Unsteady gait?
4. Patient does not understand the nature of illness and potential for bad outcome?
5. Abnormal vital signs
   - For Adults
     - Pulse greater than 120 or less than 60?
     - Systolic BP less than 90?
     - Respiration greater than 30 or less than 10?
   - For minor/pediatric patients
     - Age inappropriate HR or
     - Age inappropriate RR or
     - Age inappropriate BP
6. Serious chief complaint (chest pain, SOB, syncope)
7. Head Injury with history of loss of consciousness?
8. Significant MOI or high suspicion of injury
9. For minor/pediatric patients: ALTE, significant past medical history, or suspected intentional injury
10. Provider impression is that the patient requires hospital evaluation

Section Two:
For providers: Following your evaluation, document information and care below:

1. Did you perform an assessment (including exam) on this patient? 
   - If yes to #1, skip to #3
2. If unable to examine, did you attempt vital signs?
3. Did you attempt to convince the patient or guardian to accept transport?
4. Did you contact medical direction for patient still refusing service?
Patient Refusal of EMS

I, ____________________________, have been offered the following by ___________________ (EMS Operational Program) but refuse (check all that apply):

- Examination
- Treatment
- Transport

Patient Name: ___________________________ Phone: ________________
Patient Address: ____________________________________________________
Signature: __________________________________ Witness: ___________________

- Patient
- Parent
- Guardian
- Authorized Decision Maker (ADM)

If you experience new symptoms or return of symptoms after this encounter, we recommend that you seek medical attention promptly.

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 he/she refused the above care/service:

_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________
_________________________________________________________________________________

Jurisdiction ___________________________ Incident: ___________________________ Date: _____________
Unit #: ___________________________ Provider Name/EID: ___________________________ Time: ____________
30. ACUPRESSURE FOR NAUSEA

a) PURPOSE
Acupressure on the P6 point can be used to reduce the intensity of nausea for patients where ondansetron is not preferable or available. It may be helpful as adjunct therapy for patients who have received ondansetron.

b) INDICATION
(1) Patients with active nausea and vomiting
(2) As adjunct therapy to patients receiving ondansetron
(3) To prevent or reduce motion sickness

c) CONTRAINDICATION
None

d) ADVERSE EFFECTS
Redness, swelling, discomfort at site if commercial wrist bands are used

e) PRECAUTIONS
Patients experiencing nausea should receive a complete assessment, especially if cardiac risk factors are present.

f) PROCEDURE
(1) Identify P6 point.
   (a) Place three of the patient’s fingers on the patient’s opposite forearm at the wrist crease.
   (b) Mark the space between the two tendons on the forearm as the P6 point.
(2) Apply pressure at this point for several seconds and encourage the patient to take over care, or apply a commercial device per manufacturer’s instructions. Have patient or parent maintain firm pressure. Onset of relief is between 30 seconds and 5 minutes.
(3) Reassess patient, rescore on BARF Scale at 5 minutes, and document response to therapy.

BARF Nausea Scale
31. MULTIPLE CASUALTY INCIDENT/UNUSUAL EVENT

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.

ALERT: THIS PROTOCOL IS SIMPLY A LIST OF REQUIRED TASKS IN THE EVENT OF AN UNUSUAL EVENT. IT IS NOT ALL-INCLUSIVE. ALL PROVIDERS ARE ENCOURAGED TO REVIEW LOCAL EMERGENCY RESPONSE PLANS, THE MARYLAND TRIAGE SYSTEM TRAINING PROGRAM, START/JUMPSTART, AND NIMS PRACTICES AND PROCEDURES ON AT LEAST AN ANNUAL BASIS.

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 (Section II.G.6)

e) Triage patients using the START/JumpSTART methods (Section II.D.7.e).

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 and/or the state electronic patient tracking system (PTS). Patient information should be written on the triage tag and be entered directly into the PTS as it becomes available.

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.
32. POTENTIALLY VOLATILE ENVIRONMENTS 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 providers, 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 providers shall be under LE escort. EMS providers 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 EMS Providers, 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 provider 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 provider.
(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 providers 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 providers 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 providers. 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 providers.

(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 provider 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 – Respiration**
   (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 EMS Providers.**

(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 EMS Providers*.

(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.
33. EMERGING INFECTIOUS DISEASE (EID) (NEW ’16)

1. Initiate General Patient Care.

2. Presentation
   An Emerging Infectious Disease 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.
2. 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.

3. 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.

**Alert** 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 (DHMH 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 (DHMH 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 (DHMH 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 *(NEW ’16)*

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 DHMH if these patients become ill and require transportation by EMS to hospitals and will contact the local jurisdictional or waived 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 waived commercial services are not available. DHMH will determine the destination hospital in these cases.
i) Communication
EMS providers 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 provider 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 provider 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.
G. BLS GLUCOMETER PROTOCOL
(EMT ONLY)

a) PURPOSE
The glucometer should be utilized by BLS providers to determine the blood glucose level in an attempt to determine the etiology of the patient’s condition and provide treatment tailored to the needs of the patient before ALS intervention can be made.

b) INDICATIONS
The glucometer should be utilized for any patient presenting with an altered mental status, seizure activity, unresponsiveness, stroke, combative, suspected cyanide poisoning, reported history of high or low blood sugar, and pediatric bradycardia or cardiac arrest.

c) TREATMENT
Utilize the glucometer to determine the patient’s blood glucose level. If the glucose level is less than 70 mg/dl:

(1) ADULT: Administer glucose paste (10–15 grams) between the gum and cheek. Consider single additional dose of glucose paste if not improved after 10 minutes.

(2) PEDIATRIC: Administer glucose paste (10–15 grams) between the gum and cheek; this may be accomplished through several small administrations. Consider single additional dose of glucose paste if not improved after 10 minutes.

IF THE GLUCOSE LEVEL IS GREATER THAN 100 MG/DL, DO NOT ADMINISTER GLUCOSE PASTE.