Statewide Standard Treatment Protocol

Delaware Basic Life Support Protocols, Guidelines and Standing Orders

For

Prehospital and Interfacility Patients

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State of Delaware
Department of Health and Social Services
Division of Public Health
Office of Emergency Medical Services

2022 Statewide Standard Treatment Protocols and
Basic Life Support Standing Orders

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This treatment protocol for basic life support has been adopted and is enacted by the State Fire Prevention Commission pursuant to Delaware Code, Title 16, Chapter 98, Section 9802 (24).

Ronald Marvel
Chairman
State Fire Prevention Commission
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INTRODUCTION AND EMT STANDARD OF CARE
Delaware Emergency Medical Technician (EMT) Protocols

Issued by the State of Delaware EMS Medical Directors
In cooperation with the Delaware State Fire Prevention Commission and
the Office of Emergency Medical Services

The Delaware Emergency Medical Technician protocols and the standing orders contained
within have been developed as an adjunct to the standards of care as contained in the United
States Department of Transportation Educational Standards and verified through the National
Registry of Emergency Medical Technicians certification process.

All Delaware certified EMS providers administering patient care are doing so under the
provisions of the State EMS Medical Director's medical license in accordance with Delaware
Code Title 16, Chapter 98 Section 9802.

These protocols are not all-inclusive. They address in particular those patients for which EMTs
may assist with previously prescribed medications such as nitroglycerin, invasive procedures
such as automatic external defibrillation, and complex clinical situations such as refusal of
treatment which the EMS medical directors have chosen to address through protocols as
reinforcement to standard EMT training.

Deviation from standing orders may be undertaken only by direct order from an approved base
station physician serving as Medical Control Physician within a Delaware Office of EMS
approved facility or by a State of Delaware EMS medical director directly involved in the care of
the patient.

"Any person, agency, organization or entity who knows or in good faith suspects child abuse or
neglect shall make a report in accordance with § 904 of this title (Title 16 of Delaware Code).
For purposes of this section, "person" shall include, but shall not be limited to, any physician,
any other person in the healing arts including any person licensed to render services in
medicine, osteopathy or dentistry, any intern, resident, nurse, school employee, social worker,
psychologist, medical examiner, hospital, health care institution, the Medical Society of
Delaware or law enforcement agency." Child Abuse Reporting Phone Contact: 1-800-292-
9582 or iseethesigns.org

Any person having reasonable cause to believe that an adult person is infirm or incapacitated
as defined in § 3902 of this title (Title 31 of Delaware Code) and is in need of protective services
as defined in § 3904 of this title shall report such information to the Department of Health and
Social Services. Division of Services for Aging and Adults with Physical Disabilities (DSAAPD): 1-800-223-9074.

If an EMS provider has reasonable cause to suspect that a person is a potential victim of human
trafficking, report the concern. National Human Trafficking Resource Center Hotline 1-888-373-
7888 (24 hours). The NHTRC call takers are trained to assist by discussing a case in a HIPAA
compliant manner.

If an EMS provider has reasonable cause to suspect that a person is a potential victim of
domestic violence, contact the National Domestic Violence Hotline 1-800-799-SAFE (7233).
Please communicate concerns to receiving facility.

All certified EMS providers involved with patient care shall adhere to all federal and state HIPAA
laws and regulations (45 CFR 160, 162, and 164 & DEL CODE 16, CH12 §1212). Providers should use due caution when using social media in order to comply with HIPAA laws and regulations.
All certified EMS providers, involved with patient care, are equally responsible for assuring the patient(s) receives appropriate medical care.

**Patient** - A patient is an individual who is sick, injured, wounded or otherwise incapacitated or helpless and seeks immediate medical attention for whom EMS has been activated. A person that denies the need for medical treatment and/or transport, but any reasonable EMS provider can see that a person(s) has an obvious injury or illness, should be considered a patient and treated as such.

**Patient Priority:**

- **Priority I** Patient suffering from an immediate life or limb threatening injury or illness.
- **Priority II** Patients suffering from an injury or illness that if left untreated could potentially threaten life or limb.
- **Priority III** Patient suffering from an injury or illness that requires medical attention but does not threaten life or limb.

**EMT Minimum skills and procedures:**

1. Patient assessment (primary and secondary surveys)
2. Patient assessment - using the pediatric assessment triangle for general impression
3. Use of body substance isolation (BSI)
4. Obtaining vital signs including temperatures
5. Scene assessment and notification responsibilities in suspected abuse cases
6. Airway control (manual)
7. Use of airway adjuncts (nasopharyngeal, oropharyngeal airways and other devices approved for BLS by the State EMS Medical Director)
8. Spine immobilization/stabilization
9. Cardio-pulmonary resuscitation
10. Bleeding control and shock management
11. Splinting of fractures and dislocations
12. Use of suction equipment
13. Application of oxygen delivery devices
14. Vaginal delivery
15. Use of tourniquets and approved hemostatic agents
16. Assist with nitroglycerin
17. Assist with bronchodilator
18. Assist with Aspirin
19. Assist with patient’s medication auto-injectors
20. Measurement of blood glucose and administration of oral glucose
21. Administration of defibrillation
22. Pulse oximetry and CO-oximetry
23. Monitor IV fluids
24. Use of a length-based color-coded resuscitation tape for age-appropriate treatments (Broselow Tape® or OEMS determined equivalent)
25. VAD support and emergency procedures
Optional EMT Skills and Procedures:

1. Use of approved mechanical chest compression device
2. Application of Junctional Tourniquet (Optional)
3. Application of iTClamp
Considerations for requesting Advanced Life Support (ALS):

If at any time during contact the patient begins to show signs of worsening, an Advanced Life Support (ALS) unit should be considered.

Basic Life Support (BLS) should request an ALS provider when the patient's needs exceed their capabilities unless transport by BLS to an appropriate receiving facility can be accomplished before ALS can initiate care, then the BLS service should transport immediately. These conditions may include but are not limited to:

- Altered level of consciousness
- Allergic reaction with difficulty breathing or swallowing, altered level of consciousness, or known previous reaction; hives within 5 minutes of exposure
- Cardiac symptoms
- Cardiac arrest
- Diabetic problem (not alert and/or abnormal breathing)
- Multi-system trauma or severe single system trauma
- OB/GYN (imminent delivery, 2nd or 3rd trimester bleeding or miscarriage)
- Overdose/poisoning (associated with any other categories on this list)
- Respiratory distress
- Respiratory arrest/failure
- Sudden Unexplained Infant Death (SUID)
- Seizures/convulsions (Status or trauma related)
- Entrapment with injuries that meet trauma triage criteria
- Severe blood loss
- Shock (Hypoperfusion)
- Stroke/CVA symptoms
- Syncope (associated with any other categories on this list or cardiac history)
- Unconsciousness
- Abnormal vital signs for that particular patient

BLS services should not delay patient care or transport while waiting for ALS personnel. If ALS arrival at scene is not anticipated before initiation of transport, arrangements should be made to rendezvous with the ALS service. If the rendezvous will delay transport greater than the transport time to the hospital, continue transport and advise the hospital of patient condition and lack of ALS on board.
Transport Requirements:

Respond to EMS call in accordance with the currently approved Priority Medical Dispatch (PMD) Protocols.

Transport shall be made in a safe manner as to prevent further injury. Utilize lights and sirens as appropriate based on patient condition.

- It is the consensus of the EMS medical directors that during transport to the hospital, the use of lights and sirens is not medically indicated for the majority of EMS patients.
- It is in the best interest of patient care that the highest medically trained on duty practitioner should determine the appropriate mode of transport based on patient condition.

Transfer patient to ambulance using the most appropriate means necessary while not exacerbating the patient(s) symptoms.

Secure patient in ambulance using appropriate equipment per ambulance and stretcher design. Agency standard operating procedures should meet or exceed manufacturers' recommendations and any applicable Delaware State Fire Prevention regulations and Delaware law.

The medical directors encourage providers to use safety restraints while the ambulance is in motion.

Transport patient to the most appropriate medical facility via appropriate mode of transportation without delay.

- Transport to the closest appropriate medical facility is priority; however, when possible, patient care is enhanced by transport to a facility of prior treatment and the patient's, family's, or personal physician's choice should be considered.
- If the patient's wishes are in conflict with existing protocol (e.g., trauma, OB, NICU, or stroke/STEMI) the appropriate destination should be chosen. The Medical Control physician is the final determinant if assistance is needed.
- EMS providers should consider diversion status when determining destination. See the diversion script approved by the Medical Directors in Appendix E. Priority I patients shall be transported to the closest appropriate facility, unless the facility is closed.
- Patient care does not end until transfer of care of the patient to appropriately trained healthcare provider.
- EMT's may use onsite Medical Control and alternate patient care locations during events deemed as Mass Gatherings by the Office of EMS and approved by the State EMS Medical Director.

At the time of patient delivery to an approved healthcare facility, the EMT must give a verbal report to a physician, physician assistant, or nurse at the patient's bedside (a triage desk report is appropriate if patient's disposition is to hallway or waiting room).
Documentation Requirements:

An essential part of prehospital medical care is the completion of a Patient Care Report (PCR). The PCR provides written documentation of patient condition and treatment for medical and legal purposes. EMS personnel shall be responsible for providing clear, concise, complete and accurate documentation.

EMS providers must complete, without exception, a State of Delaware PCR on each patient contact, and shall document all relevant findings, and treatments.

- In the absence of extraordinary circumstances, a PCR should be submitted to the receiving facility within four (4) hours of patient disposition.
- EMS providers must complete and submit a PCR to the receiving facility prior to going off duty.
- EMS calls should only be entered into the PCR system as a “service call” or “public assist” if the person does not meet the definition of a patient after an assessment. A patient is an individual who is sick, injured, wounded or otherwise incapacitated or helpless and seeks immediate medical attention for whom EMS has been activated. A person that denies the need for medical treatment and/or transport, but any reasonable EMS provider can see that a person(s) has an obvious injury or illness, should be considered a patient and treated as such.
- If the reasonable EMS provider on scene determines that a person (s) is a patient, the person (s) should be treated as a patient and a patient refusal is needed.
- A PCR entry is not needed for any Good Samaritan occurrences. Providers are encouraged to leave their information with the transporting agency.

A completed PCR is also necessary to identify EMS providers in the event of a potential infectious disease exposure.

Use of Quality Assurance/Quality Improvement (QA/QI) Requirements:

Quality Assurance/Quality Improvement (QA/QI) measures must be compliant with the established Delaware State Fire Prevention Commission QA/QI Committee detailed in the State of Delaware's Ambulance Regulations and approved by the State BLS EMS Medical Director and State EMS Medical Director.
EMT/TELEPHONE REPORT GUIDELINES

The EMT report to Medical Control should be brief and concise. The goal is to provide enough vital information to Medical Control so that they may provide informed direction for the patient's continued care and plan for the patient's disposition. Reports generally should not exceed thirty (30) seconds in duration in order to provide economical use of time by the EMT, the Medical Control physician, and nursing personnel.

See Appendix F for hospital contact information or click the link below:
https://www.dhss.delaware.gov/dph/ems/ems.html

For Priority I patients call online Medical Control utilizing the following report format:

- BLS unit number
- Specific notification (Trauma, Cardiac Arrest, Stroke, CPAP, etc.)
- Estimated time of arrival.
- Priority.
- Patient age.
- Patient sex.
- Chief complaint and related past medical history (i.e., patient with chest pain, history of MI and CABG or patient with altered mental status and history of insulin dependent diabetes).
- Vital signs.
- Significant physical findings (i.e., patient with shortness of breath found to have wheezing and to be hot to the touch, or the patient complaining of leg pain who has deformity of the mid-thigh without distal pulses).
- Care rendered.
- Response to care.

For hospitals that prefer radio reports regarding BLS patients who are a Priority of II or III and are being treated by standing orders with no anticipated requests for orders, the following brief report format is acceptable:

- BLS unit number.
- Priority.
- Patient age.
- Patient sex.
- Chief complaint
- Standing Order being followed
- Estimated time of arrival

The above information should be more than adequate for most BLS runs. When additional information is felt to be important for patient care or disposition, the Medical Control physician is well within their jurisdiction to request more information.
GENERAL PATIENT CARE (ADULT)

INDICATIONS: Any patient, who is greater than or equal to the age of 15 years, requiring prehospital medical evaluation by a prehospital health care provider in the State of Delaware.

The General Patient Care protocol will be followed in conjunction with all other applicable protocols.

A patient is an individual who is sick, injured, wounded or otherwise incapacitated or helpless and seeks immediate medical attention for whom EMS has been activated. A person that denies the need for medical treatment and/or transport, but any reasonable EMS provider can see that a person(s) has an obvious injury or illness, should be considered a patient and treated as such.

The most current version of the American Heart Association Guidelines for Cardiopulmonary Resuscitation is considered the standard for CPR within these protocols.

- Scene Safety, Observe body substance isolation (BSI) precautions.
- Identify the number of patients; perform Triage if necessary. See Triage Protocol.
- Consider the need for additional resources.
- Manage cervical spine as needed.
- Complete patient assessment: Level of consciousness (AVPU, Determine GCS).
- Assess and manage the airway.
- Assess breathing rate, rhythm, quality and oxygenation.
- Assess and manage circulation.
- Obtain vital signs. Monitor Blood Glucose as appropriate.
- Obtain SAMPLE history and OPQRST history if patient can speak (Onset, Provocation/Palliation, Quality, Rate, Severity, Time)
- Assess pertinent body systems as appropriate.
- Assess and record pain severity, if applicable.
- Assign treatment priority and make a transport decision.
- For transport consider closest appropriate medical facility, keeping in mind patient (family) requests and diversion status.
- Victims of sexual assault should be transported to a facility staffed with a Sexual Assault Nurse Examiner (SANE) / Forensic Nurse Examiner (FNE). If patient has significant trauma transport to appropriate trauma facility.
- On scene direction of medical care is provided by the on-duty Delaware EMS provider.
with the highest level of licensure and/or certification. Rescue operations and control of the scene remains under the direction of the Fire Officer in Charge.

- **Contact Medical Control** as needed.

- Monitor and reassess as appropriate.

- Responsibility of care does not end until transfer of care of the patient to an appropriately trained health care provider is completed.

Any person having reasonable cause to believe that an adult person is infirm or incapacitated as defined in § 3902 of this title (Title 31 of Delaware Code) and is in need of protective services as defined in § 3904 of this title shall report such information to the Department of Health and Social Services.

Division of Services for Aging and Adults with Physical Disabilities (DSAAPD): 1-800-223-9074.
ADULT NAUSEA / VOMITING

INDICATIONS: Any patient who is older than the age of 15 years. Ongoing nausea or vomiting that is not from a toxin or alcohol ingestion.

- Follow General Patient Care protocol
- Provide appropriate supplemental oxygen. Obtain a pulse oximeter reading. Manage the airway appropriately.
- Consider Ondansetron (Zofran) 8 mg sublingual (oral dissolving tablets) for adults.
- Nausea may be improved by allowing patient to smell or sniff an open alcohol prep/wipe.
  - Allow patient to hold and sniff an Isopropyl Alcohol pad, commonly known as prep/alcohol wipe has been demonstrated to reduce nausea
  - Have patient hold the alcohol pad to the nose and breathe in through their nose until symptoms of nausea dissipate.
  - This can be used in conjunction with Ondansetron (Zofran) or independent of Ondansetron (Zofran)
GENERAL PATIENT CARE (PEDIATRIC)

INDICATIONS: Any patient, who is less than the age of 15 years (neonates are defined as a patient 30 days and under), requiring prehospital medical evaluation by a prehospital health care provider in the State of Delaware.

The General Patient Care protocol will be followed in conjunction with all other applicable protocols.

A patient is an individual who is sick, injured, wounded or otherwise incapacitated or helpless and seeks immediate medical attention for whom EMS has been activated. A person that denies the need for medical treatment and/or transport, but any reasonable EMS provider can see that a person(s) has an obvious injury or illness, should be considered a patient and treated as such.

The most current version of the American Heart Association Guidelines for Cardiopulmonary Resuscitation is considered the standard for CPR within these protocols.

- Scene Safety, Observe body substance isolation (BSI) precautions.
- Identify the number of patients; perform Triage if necessary. See Triage Protocol.
- Consider the need for additional resources.
- Manage cervical spine as needed.
- General assessment should be done using the pediatric assessment triangle (PAT).
  - Appearance
  - Work of breathing
  - Circulation
- After using the PAT, proceed to a primary assessment:
  - Airway for patency.
    - If patient has stridor, transport sitting straight upright to assist with clearing of respiratory secretions. Do not attempt to examine upper airway or otherwise aggravate the patient.
    - Manage the airway as required.
  - Breathing for respiratory effort and quality; use of a Pulse Oximeter as appropriate.
    - Administer oxygen as appropriate.
  - Circulation for pulse rate, skin temperature and capillary refill.
    - CO-oximetry may be performed as an option by agencies carrying CO monitoring equipment.
• Expose the patient as needed for assessment needs.
  o Keep in mind that pediatric patients are prone to hypothermia faster than their adult counterparts. Maintain a warm environment and keep exposure to a minimum.

• Treat life-threatening conditions as necessary.

• Evaluate blood pressure, pulses, respiratory rate, GCS (Glasgow Coma Scale) and tactile temperature: if available use thermometer to take an accurate temperature. Refer to normal vital signs chart for pediatrics, or a Broselow tape or OEMS determined equivalent.

• Monitor and reassess the patient as appropriate.

• Monitor blood glucose level as appropriate.

• If no life threat has been determined in the primary survey, proceed to a secondary survey that will include a focused medical history using the SAMPLE mnemonic and thorough physical exam.

• Assess and record pain severity, if applicable, using age-appropriate pain scale.

• Assign treatment priority and make transport decision.

• For transport, consider closest appropriate medical facility, keeping in mind patient (family) requests and diversion status.

• If at all possible, do not separate the parent/caregiver and the child.

• Patients should be taken to the approved facility's emergency department, labor and delivery area or to an inpatient bed if arranged prior to arrival at the facility. If there are questions or doubts as to the appropriate facility or point of delivery, the Medical Control physician will be the arbitrator. All unstable patients should be transported directly to an emergency facility.

• Victims of sexual assault should be transported to a facility staffed with a Sexual Assault Nurse Examiner (SANE) / Forensic Nurse Examiner (FNE). If patient has significant trauma, transport to appropriate trauma facility.

• Patients are to be transported to Delaware Office of EMS approved facilities within the EMS agency's usual operations area.

• On scene direction of medical care is provided by the Delaware EMS provider with the highest level of licensure and/or certification.

It should be noted that the protocol above is a guideline to be followed in as much as it aids in providing appropriate and timely medical care. The EMT provider may change the order or omit steps listed above as dictated by sound judgment of the care provider and/or presentation of the patient(s).
"Any person, agency, organization or entity who knows or in good faith suspects child abuse or neglect shall make a report in accordance with § 904 of this title (Title 16 of Delaware Code). For purposes of this section, "person" shall include, but shall not be limited to, any physician, any other person in the healing arts including any person licensed to render services in medicine, osteopathy or dentistry, any intern, resident, nurse, school employee, social worker, psychologist, medical examiner, hospital, health care institution, the Medical Society of Delaware or law enforcement agency."

Child Abuse Reporting Phone Contact: 1-800-292-9582 or iseethesigns.org.
REFUSAL OF SERVICE

INDICATIONS: EMTs respond to various scenes where 911 or emergency services are activated due to a person(s)/patient(s) that appears to be in some sort of distress and may be in need of Emergency Medical assistance. It is important that the EMT obtains the patient’s informed consent before leaving the scene; otherwise, the EMT might be exposed to legal liability for abandonment of the patient.

A patient is an individual who is sick, injured, wounded or otherwise incapacitated or helpless and seeks immediate medical attention for whom EMS has been activated. A person that denies the need for medical treatment and/or transport, but any reasonable EMS provider can see that a person(s) has an obvious injury or illness, should be considered a patient and treated as such.

- Coercing a patient or family into a Refusal of Services may lead to loss of EMS provider privilege by the State Fire Prevention Commission.
- Follow General Patient Care Protocols and any other appropriate protocols that may be required based on the patient condition, complaint, or your assessment.
- Discussion of refusal should be initiated by the patient and/or their guardian.
- If patient and/or patient’s guardian wishes to refuse treatment and/or transport to a medical facility:
  - Inform the patient about the needed treatment and possible outcomes including verbalizing the possibility of disability and death.
  - Every effort should be made to persuade the patient to consent to treatment and/or transport.
  - Consider involving family, Medical Control and law enforcement as needed.
- Contact Medical Control for patients presenting or having originally presented with:
  - Suspicion of intoxication by drugs or alcohol
  - Past medical history or suspicion of dementia
  - Any intervention performed by any other healthcare provider
  - A summons of EMS to a health care facility or call initiated by a healthcare provider
  - Suspicion of acute mental disease or suicidal or homicidal ideation
  - Suspicion of a significant head injury
  - Respiratory distress
  - Abnormal vital signs (normal vital signs are defined as a heart rate between 60-100 bpm, systolic blood pressure greater than 100mmHg, respiratory rate 12-20, and a SpO2 reading greater than 92% on room air)
  - Altered mental status who remain altered
  - An age less than 18 years
  - Any time ALS is dispatched and recalled by BLS prior to ALS arrival unless the patient meets hypoglycemia, opiate overdose or taser barb removal protocol parameters for refusal of service.

Obtain a signed Refusal of Service form and document the informed consent process, concerns, and if applicable the physician number on the appropriate reports.
CHEST PAIN

Non-traumatic - Possible Cardiac Origin

INDICATIONS: The pattern of pain suggestive of cardiac origin is highly variable. Chest or epigastric pain associated with shortness of breath, sweating, nausea, vomiting, radiating or non-radiating pain of the neck, jaw, left arm, or back. Patients with chest pain of suspected cardiac etiology require rapid stabilization and transport.

- Follow General Patient Care Protocol.
- Obtain a Pulse Oximeter reading. Provide appropriate supplemental oxygen titrated to greater than 92%.
- Do not administer aspirin if the patient reports an allergy to aspirin or other NSAIDs.
- Administer/assist the patient with taking uncoated Aspirin up to 325mg. This dose includes any dispatcher directed or patient administered doses prior to EMS arrival.
- Withhold nitrates and **Contact Medical Control** if the patient relates taking sildenafil (Viagra/Revatio) or vardenafil (Levitra) within the last 24 hours or tadalafil (Cialis, Adcirca) for pulmonary hypertension, or any other prescription erectile dysfunction drugs or pulmonary hypertension drugs within the last 48 hours.
- If the patient has their prescribed nitroglycerin and their systolic blood pressure is greater than 90 mm Hg, assist or give the patient nitroglycerin as prescribed. Assess the patient's blood pressure before each dose. The patient should not take nitroglycerin if the systolic blood pressure falls below 90 mm Hg. Do not exceed 3 doses given 3 to 5 minutes apart. Further orders must come from **Medical Control**.
- Make sure that the medication is prescribed to the patient and has not expired.
- If a paramedic unit is not available, radio a report to the emergency department advising of the estimated time of arrival (ETA) and patient status. Consider paramedic unit intercept route. Do not delay transport.
- **Contact Medical Control** directly with questions regarding nitroglycerin therapy or if medication is administered without a paramedic.
ACUTE RESPIRATORY DISTRESS (ADULT)

INDICATIONS: Signs and symptoms of acute exacerbations of asthma, emphysema, reactive airway disease and allergic reactions may include wheezing, cough, shortness of breath, diminished breath sounds, retractions, tachypnea, and/or air hunger.

- Follow General Patient Care Protocol.
  - Be aware of any current recommended restrictions of nebulized medications
- Provide appropriate supplemental oxygen. Obtain a Pulse Oximeter reading. Consider obtaining a carbon monoxide reading, if greater than 5, apply oxygen.
- Assess lung sounds.

**MDI**
- If a patient has a bronchodilator meter dose inhaler prescribed by their physician, give 4 puffs and can repeat in 5 minutes. Hold if pulse is greater than 150 beats per minute. Contact Medical Control before assisting with additional doses or if the patient took more than 4 puffs within one hour of EMS arrival.

**NEBULIZER**
- If patient's heart rate is less than 150 beats per minute, and if appropriate, assist the patient with their own nebulizer as prescribed by the patient's physician, up to a 2-unit dose. Connect nebulizer to an oxygen source at 8 liters per minute.
- If upon arrival patient is currently taking prescribed nebulizer, it is appropriate to transport the patient while finishing the treatment with EMS provided oxygen. Once complete continue appropriate oxygen therapy.

- If a paramedic unit is not available, radio a report to the emergency department advising of the estimated time of arrival (ETA) and patient status. Consider paramedic unit intercept route. Do not delay transport.
- EMS agencies may also carry and utilize Albuterol meter dose inhaler with appropriate spacer attached, in place of nebulized Albuterol. Give 4 puffs and can repeat in 5 minutes. Hold if pulse is greater than 150 beats per minute.
- Contact Medical Control directly with any questions or concerns regarding medication therapy as needed or if medication administered without a paramedic.
ACUTE RESPIRATORY DISTRESS/FAILURE (PEDIATRIC)

INDICATIONS: Signs and symptoms of acute exacerbations of asthma, reactive airway disease and allergic reactions may include wheezing, cough, shortness of breath, diminished breath sounds, retractions, tachypnea, stridor and/or air hunger.

Acute Respiratory Distress: a clinical state characterized by abnormal respiratory rate, and an increased effort represented by nasal flaring, retractions and accessory muscle use. Respiratory distress can be associated with changes in airway sounds, skin color, and mental status.

Acute Respiratory Failure/Arrest: a clinical state of inadequate oxygenation, ventilation, or both. It may be characterized by signs of distress or inadequate respiratory effort.

- Follow Pediatric General Patient Care Protocol.
  - Be aware of any current recommended restrictions of nebulized medications
- Perform an initial assessment using the pediatric assessment triangle.
- Keep patient warm.
- Consider Complete Airway Obstruction (Foreign body).
- Consider Partial Airway Obstruction (Upper airway).
  - Do not attempt an invasive airway maneuver (Oropharyngeal and Nasopharyngeal Airway adjuncts). Avoid any agitation, place child in position of comfort.
- Provide appropriate supplemental oxygen, as appropriate. Obtain a pulse oximeter reading. Consider obtaining a carbon monoxide reading, if greater than 5, apply oxygen via non-rebreather.
- Consider Reactive Airway Disease (lower airway).
  - Wheezing, grunting, retractions, tachypnea, diminished respirations, decreased breath sounds, tachycardia/bradycardia, and/or decreased level of consciousness.
  - Place patient in position of comfort.

  **MDI**
  - If a patient has a bronchodilator meter dose inhaler prescribed by their physician, assist the patient as prescribed. The inhaler may be used again in fifteen minutes for a total of 4 puffs. **Contact Medical Control** before assisting with additional doses or if the patient took more than 4 puffs within one hour of EMS arrival (use spacer if available).
  - If the patient's pulse rate is over 180 beats per minute, **Contact Medical Control** prior to a second dose of bronchodilator.
NEBULIZER

- If patient's heart rate is less than 180 beats per minute, and if appropriate, assist the patient with their own nebulizer as prescribed by the patient's physician. These include albuterol, levalbuterol and Combivent. Connect nebulizer to an oxygen source at 8 liters per minute.

- If upon arrival patient is currently taking his prescribed nebulizer, it is appropriate to transport the patient while finishing the treatment with EMS provided oxygen. Once complete continue appropriate oxygen therapy.

- If a paramedic unit is not available, radio a report to the emergency department advising of the estimated time of arrival (ETA) and patient status. Consider paramedic unit intercept route. Do not delay transport.

- **Contact Medical Control** directly with any questions or concerns regarding medication therapy as needed or if medication administered without a paramedic.

RESPIRATORY FAILURE/ARREST

- Manage Airway as appropriate.

- Nasal airways are contraindicated in infants and small children. Assess the size of the nares and the proper length of the adjunct, as not to occlude the airway.

- Place in the appropriate airway adjunct to assist in patency.

- Administer appropriate oxygen therapy.

- Monitor cardiac status and be prepared to begin CPR.

**Note:** A high-risk infant is an infant who is on an apnea monitor, had an episode of apnea or who has been identified as having an "apparent life-threatening event" (ALTE) or brief resolved unexplained event (BRUE). These infants include those who have experienced periods of apnea (cessation of breathing) or are at risk of prolonged apnea. When you arrive at the scene of an incident involving this type of baby, no matter how well the baby may look, transport to the ED is always advised.
ALBUTEROL

INDICATIONS: Signs and symptoms of acute exacerbations of asthma, emphysema, reactive airway disease and allergic reactions may include wheezing, cough, shortness of breath, diminished breath sounds, retractions, tachypnea, and/or air hunger. Providers will be able to identify the need for Albuterol medication treatments and administer them as appropriate.

- Follow General Patient Care Protocol.
  - Be aware of any current recommended restrictions of nebulized medications

- Consider the administration of 0.5 mg nebulized ipratropium bromide (Atrovent) with Albuterol.

- Consider use of the Allergic Reaction Protocol.

- Request ALS. If a paramedic unit is not available, radio a report to the emergency department advising of the estimated time of arrival (ETA) and patient status. Consider paramedic unit intercept route. Do not delay transport.

- If patient is less than 1 year, **Contact Medical Control** immediately.

- If patient’s pulse is less than 150 beats per minute for adult or 180 beats per minute for pediatric and the patient has a known history of Asthma or COPD; or signs of Asthma, COPD, or Allergic reaction are present, administer Albuterol as follows:
  - For patients 6 years of age and above: administer 5 mg Albuterol via nebulizer with oxygen flow set at 8LPM.
  - For patients 1-5 years of age: administer 2.5 mg Albuterol via nebulizer with oxygen flow set at 8LPM.

- Reassess patient, especially lung sounds, vitals, and oxygen saturation.

- If signs and symptoms of respiratory distress persist, repeat dose as follows:
  - For patients 6 years of age and above: administer 5 mg Albuterol via nebulizer with oxygen flow set at 8LPM*.
  - For patients 1-5 years of age: administer 2.5 mg Albuterol via nebulizer with oxygen flow set at 8LPM*.

- **Contact Medical Control** with any questions or concerns. Document Medical Control physician number and any orders on the patient care report.

- Document on the EMS patient care report the name of the medication, the time(s) of administration, the number of doses, and pulse rate before administration.

**When using an albuterol MDI, please use clean spacer if available.**

* If respiratory status worsens, refer to CPAP protocol with inline neb Utilize BVM for pediatric patients.
CONTINUOUS POSITIVE AIRWAY PRESSURE (CPAP)

INDICATIONS: Respiratory distress or failure, due to cardiogenic pulmonary edema (CHF), asthma, chronic obstructive pulmonary disease (COPD), or emphysema in which the patient demonstrates spontaneous respirations and a patent, self-maintained airway.

VITAL SIGNS AND ASSESSMENT CRITERIA:

- Tachypnea = Respiratory Rate greater than or equal to 24 bpm
- Tachycardia = Heart Rate greater than or equal to 100 bpm
- Hypertension = Systolic Blood Pressure greater than or equal to 120 mmHg
- Hypoxia = Pulse Oximetry reading less than or equal to 90%
- Labored breathing that results in the patient being unable to complete a full sentence

CONTRAINDICATIONS:

- Circumstances in which endotracheal intubation or a surgical airway is preferred or necessary to secure a patent airway
- Circumstances in which the patient does not improve or continues to deteriorate despite CPAP administration
- Patients with respiratory distress secondary to trauma

Follow General Patient Care Protocol.

Assure a patent airway.

Administer 100% O2 via appropriate delivery system.

Perform appropriate patient assessment including obtaining vital signs, pulse oximeter (SpO2) reading, and cardiac rhythm (regular or irregular).

Apply CPAP device per manufacturer's instructions. *

Monitor continuous pulse oximetry.

Albuterol nebulizers may be kept in-line with CPAP. This may require the nebulizer flow rate to increase based on CPAP manufacturer's recommendations.

Contact Medical Control as soon as possible to allow for prompt availability of hospital CPAP / BiPAP equipment and respiratory personnel.

*For circumstances in which the patient does not improve or continues to deteriorate despite CPAP and/or medical therapy, terminate CPAP administration and perform BVM ventilation
ALLERGIC REACTION

INDICATIONS: Generalized allergic manifestations such as urticaria, swelling, respiratory distress, or a known allergen exposure.

Severe Allergic Reactions include: Airway obstruction (partial or complete), swelling of the tongue, face, or neck areas, clinical evidence of shock including altered mental status, confusion, hypotension (systolic less than 90mmHg), delayed capillary refill, and cool, clammy, or mottled skin.

- Follow General Patient Care Protocol.
- Assess lung sounds.
- Consider Respiratory Distress Protocol.

SEVERE ALLERGIC REACTIONS (ANAPHYLAXIS)

- If a patient has epinephrine via auto-injector prescribed by their physician, assist the patient with their epinephrine auto-injector. Otherwise, utilize the BLS agency OEMS approved auto-injector if available.
- Administer one dose of epinephrine via auto-injector as indicated.
- Epinephrine auto-injector (patients greater than 30kg / 66lbs) delivers a single 0.3 mg epinephrine dose
- Pediatric epinephrine auto-injector (children 15-30kg / 33-66lbs) delivers a single 0.15 mg epinephrine dose
- Check the auto-injector to ensure the medication has not expired, has not become discolored, does not contain particulates, or sediments.
- Medical Control should be contacted before an additional dose of epinephrine via auto-injector is administered if symptoms continue after 10 minutes.
- Consider Albuterol protocol.
- If a paramedic unit is not available, radio a report to the emergency department advising of the estimated time of arrival (ETA) and patient status. Consider paramedic unit intercept route. Do not delay transport.
- Contact Medical Control directly with any questions or concerns regarding medication therapy as needed or if medication administered without a paramedic.
Optional IM Draw Dose for Epinephrine

- OEMS approved Restricted Draw Syringe
- 0.5mg epinephrine dose IM for adult patients greater than 30kg / 66lbs
- 0.25mg epinephrine dose IM for pediatric patients between 15-30kg / 33-66lbs
- Medical Control should be contacted before an additional dose of epinephrine via auto-injector is administered if symptoms continue after 10 minutes.

**MODERATE ALLERGIC REACTIONS**

- Administer Diphenhydramine (Benadryl®) 25-50 mg orally for adults
- Administer 25 mg liquid Diphenhydramine (Benadryl) to pediatric patients greater than 2 years of age.
ALTERED MENTAL STATUS

INDICATIONS: Incomprehensible speech, inappropriate verbal responses, inability to follow verbal commands, decreased responsiveness, or unresponsiveness. If a patient is known to have Diabetes Mellitus and has altered mental status, the cause of the altered mental status may be low blood sugar.

- Follow General Patient Care Protocol.
- Determine the appropriate response of the patient based on the developmental expectations of each age group. Enlist the assistance of the parent/caregiver or family member to determine what is "normal" for this patient. *
- **Contact Medical Control** if the patient is less than the age of 15 years of age for guidance with appropriate blood sugar level and medication dosages.
- Provide appropriate supplemental oxygen. Obtain a pulse oximeter reading. Manage the airway appropriately.
- Obtain a blood sugar level. If the blood sugar is less than 60mg/dl and the patient is alert and able to protect their airway, use oral glucose 15-24 grams. Make sure that the oral glucose has not expired.
- If the patient is unresponsive or not alert enough to protect their own airway, paramedics or hospital personnel will need to administer intravenous glucose in order to avoid aspiration.
- Monitor and record vital signs.
- If patient exhibits signs of shock, treat for shock.
- If after 10 minutes the patient continues to be symptomatic, re-determine blood glucose level and administer a second dose of oral glucose 15-24 grams if glucose is still below 60mg/dl and patient is alert and able to protect their own airway.
- If a paramedic unit is not available, initiate transportation to a CT capable, stroke certified** medical facility and provide a radio report to the emergency department advising them of the ETA and patient's condition. Do not delay transport.
- **Contact Medical Control** directly with any questions or concerns regarding medication therapy as needed.
- Consider alternative causes of altered mental status.
- If an insulin dependent diabetic with documented hypoglycemia due to missed meal or increased physical activity, who is not on any oral hypoglycemics and is with family/friends with the capability to consume a meal, fully awakens with EMS treatment, he (she) may refuse transport to a medical facility.
- For suspected narcotic overdose:
  - Refer to Suspected Opiate Overdose Protocol.
**Special Considerations for causes of Altered Mental Status:**

A  Alcohol and abuse
E  Epilepsy, electrolytes, encephalopathy
I  Insulin
O  Opiates, overdose
U  Uremia
T  Trauma, temperature
I  Infection
P  Poison, Psychogenic
S  Shock, seizure, stroke, space occupying lesions, SAH

**Stroke Certified by the State of Delaware or The Joint Commission (TJC), formerly the Joint Commission on Accreditation of Healthcare Organizations (JCAHO). Refer to the county EMS Medical Director’s current list of stroke certified medical facilities.**
SUSPECTED OPIATE OVERDOSE

INDICATIONS: Incomprehensible speech, inappropriate verbal responses, inability to follow verbal commands, decreased responsiveness, or unresponsiveness, respiratory distress or apnea. This protocol will allow BLS to treat patients with a history based on bystanders, provider’s prior knowledge of the patient, or suspicion of potential narcotic overdose as evidenced by nearby medications or drug paraphernalia.

If patient is pulseless start CPR. With suspected opiate overdose, naloxone (Narcan) should be given after CPR is initiated.

- Follow General Patient Care Protocol with an emphasis on airway management.
- In unresponsive patients initiate BVM ventilation with appropriate airway management techniques and CPR as appropriate.
- Obtain a pulse oximeter reading. Provide appropriate supplemental oxygen titrated to greater than 92%.
- If patient is less than 1 years old, Contact Medical Control immediately for guidance.

**INTRANASAL**

- Administer 1mg to 2 mg Naloxone (Narcan) IN (OEMS approved intranasal device) to provide for a patent, self-maintained airway and adequate respirations. If no improvement in the patient’s respiratory status after four (4) minutes, a second dose of 1 mg to 2 mg of Naloxone may be given in the opposite nare.

**INTRAMUSCULAR (AUTO-INJECTOR)**

- Administer 0.4 mg naloxone (Narcan) IM (intramuscular via the auto- injector device) to provide for a patent, self-maintained airway and adequate respirations. If no improvement in the patient’s respiratory status after four (4) minutes, a second dose of 0.4 mg of Naloxone IM may be given in the opposite thigh.

**Caution:** patients with near complete reversal of a narcotic overdose may become very agitated and combative.

- Continue to manage the patient’s airway until they are breathing adequately and are able to protect their airway from aspiration.
- Do not delay safe transport to await results of treatment.
- If there is no response to Naloxone (Narcan) within five minutes, consider other causes of altered mental status and proceed to alternative standing orders.
- If a paramedic unit is not available, radio a report to the emergency department advising of the estimated time of arrival (ETA) and patient status. Consider paramedic unit intercept route. Do not delay transport.
- Contact Medical Control directly with any questions or concerns regarding medication therapy.
Considerations for refusal:
If the patient regains full consciousness after treatment, they should be encouraged to be transported to the hospital to monitor for relapse and receive counseling about substance abuse treatment.

If the patient wishes to refuse treatment, medical control contact is not necessary if the patient meets the following criteria:

- Alert and oriented
- Suspicion of opioid ingestion through injection or inhalation
- It is preferred that the patient be left with an adult who can monitor their safety.
- Has been given information about resources for treatment and/or opioid rescue kit (if available).
- If there are any patient care concerns, call medical control for assistance.
SEPSIS (Adult)

INDICATIONS: Sepsis is the life-threatening manifestation of severe infection.

- Follow General Patient Care Protocol.
- Provide appropriate supplemental oxygen. Obtain a pulse oximeter reading.
- Monitor blood glucose level as appropriate.
- Consider sepsis in patients presenting with:
  - Suspicion of infection*
  - 2 or more systemic inflammatory response syndrome (SIRS) criteria:
    - Temperature greater than 38 C (100.4 F) or less than 36 C (96.8 F)
    - Heart rate greater than 90
    - Respiratory rate greater than 20
    - Hypotension (Systolic BP less than 90)
- Consider requesting paramedics
- **Contact Medical Control** directly if 2 or more SIRS criteria exist, and request SEPSIS alert.
- Notify receiving hospital upon arrival of potentially septic patient

*Risk factors for infection:

- Elderly patients with altered mental status from baseline
- Nursing home patients
- Chronic disease (e.g., diabetes, renal failure/dialysis)
- Immunosuppression (e.g., cancer with chemotherapy, HIV+, transplant)
- Indwelling catheters and central lines
SUSPECTED STROKE

INDICATIONS: Abnormalities in VAN Stroke Assessment, altered mental status, seizure, speech deficit, facial droop, headache, paresthesia, and hemiparesis in the absence of trauma, weakness, ataxia, visual disturbances, nausea, vomiting, general malaise, abnormal pupillary function, or other symptoms of suspected cerebral ischemia or hemorrhage.

- Follow General Patient Care Protocol.
- Administer oxygen via nasal cannula, to maintain oxygen saturation equal to 92%.
- Place patient flat if tolerated (to augment cerebral perfusion), or in a low to semi-Fowler’s position if airway or secretion concerns.
- Obtain a blood sugar.
- Consider ALS.
- Determine onset of symptoms. Onset is defined as last time the patient was verified as not having a neurological deficit or Last Known Well (LKW).
- Obtain family contact information, preferably a cell phone number.

Perform VAN Stroke Assessment:

- Determine if patient has arm and/or leg drift, unilateral weakness, or paralysis. If no weakness noted, assessment ends.
  - Patient is **VAN negative**
- If arm and/or leg drift, unilateral weakness, or paralysis is present, continue with the VAN Assessment.
  - **V** – **Visual Disturbance**: Does the patient have double-vision, visual field cut, or new loss of vision?
  - **A** – **Aphasia**: Does the patient have difficulty forming words, or difficulty understanding/following commands? Does the patient have difficulty recognizing objects correctly?
  - **N** – **Neglect**: Refers to patient’s senses and gaze. Does the patient present with a gaze deviation or inability to cross midline? Is the patient unable to feel both sides at the same time when touched, unable to recognize their own arm, or ignoring one side?

- If patient has arm and/or leg weakness AND any visual/aphasia/neglect symptoms, patient is **VAN positive**.
- If patient has arm and/or leg weakness and NO visual/aphasia/neglect symptoms, patient is **VAN negative**.
Contact Medical Control for suspected stroke patients. Early notification and prehospital stroke alert activation for receiving hospitals is paramount with acute stroke patients.

Transport to the nearest appropriate State of Delaware or Joint Commission certified Stroke Center without delays and request prehospital stroke alert for the following categories:

- VAN negative and LKW less than 4.5 hours → go to nearest certified Stroke Center.
- VAN positive and LKW less than 4.5 hours → contact local medical control to discuss destination.
- VAN positive and LKW greater than 4.5 hours (including wake up stroke or unknown LKW) or considering hemorrhagic stroke → consider transport to certified Thrombectomy Capable or Comprehensive Stroke Center.

Communicate to receiving facility the use of anticoagulants.

See Stroke Assessment Tool (Appendix G).

Changes in hospital fibrinolytic protocols and the addition of interventional therapy may occur which could result in an interval change to this standing order.

*Certified Stroke Center by the State of Delaware or The Joint Commission (TJC), formerly the Joint Commission on Accreditation of Healthcare Organizations (JCAHO)

**The Office of Emergency Medical Services (OEMS) will periodically compile and publish a list of approved receiving facilities based on the receiving facilities level of certification and available types of care. This list should be considered when determining the most appropriate destination for patients.
INITIATION OF RESUSCITATIVE EFFORTS

INDICATIONS: For initiation of cardiopulmonary resuscitation (CPR) for patients in cardiac arrest.

- Follow General Patient Care Protocol.
- Witnessed or suspected recent arrest will get immediate CPR.
- For patients with Ventricular Assist Devices (VAD’s) reference VAD protocol.
- CPR (use of mechanical chest compression device is recommended as per manufacturer’s recommendations) shall be initiated for all patients unless one or more of the following criteria apply:
  - Resuscitation would place the rescuer at significant risk of physical injury.
  - Patient is pulseless and apneic (without vital signs), cold in a warm environment, along with rigor mortis and/or dependent lividity.
  - Injuries which are obviously incompatible with life. *
    - decapitation
    - body fragmentation
    - severe crush injury to head (without vital signs)
    - severe crush injury to chest (without vital signs)
    - severe thermal burns (without vital signs)
    - gunshot wounds to the head with lateral entrance wound and an opposite side exit wound (without vital signs)
  - Decomposition of the body
    - skeletalization
    - severe bloating (without vital signs)
    - skin slough (without vital signs)
- Confirmation that a patient is without vital signs should be done by looking, listening, and feeling for breathing, along with checking for a carotid pulse and one additional pulse point (i.e., femoral, radial).
- It is preferable these steps be performed by two EMTs. Both providers must agree with the decision not to begin CPR. If there is any disagreement, resuscitation is to begin immediately.
- Presentation of any legal document to withhold life-saving efforts refer to DNR Orders protocol. Contact with Medical Control as required. (Example: DMOST)
- For patients who do not meet the criteria for initiation of cardiopulmonary resuscitation, withhold resuscitation and have paramedics continue in non-emergent for a death pronouncement.

*At no time should BLS cancel paramedics. ALS must make pronouncement in the field.
CARDIAC ARREST (ADULT)

INDICATIONS: Current AHA guidelines reflect the importance of compressions for survival from cardiac arrest. EMS practice must evolve to address this important change.

- Compressions should begin as soon as possible following EMS arrival.
- Treating the patient where they are found allows compressions to be started without delay. Only provider safety issues should prompt patient movement off the scene.
- High Quality CPR
  - Crews should perform continuous compression PIT CREW HIGH PERFORMANCE CPR,
    - Switch compressors every 2 minutes
    - No pauses for ventilations
    - Ventilations on the upstroke of CPR
- No procedure should slow or stop compressions
- Compressions should be FAST, HARD, and DEEP at a rate of 100-120 compressions per minute and to a depth of at least 2 inches.
  - Ensure complete recoil of the chest wall prior to the next compression
- Ventilations
  - Ventilate at 8-10 breaths per minute to decrease intra-thoracic pressure
  - Ventilations should be just enough to see chest rise
- Interruption for defibrillation should be minimal and compressions should resume AS SOON AS shock delivery is complete.
- Consider mechanical CPR device if adequate PIT CREW HIGH PERFORMANCE CPR cannot be maintained or transport is indicated,
- Mechanical chest compression device*, if utilized, should be set to continuous.
- Complete a minimum of 20 minutes of high-quality CPR performed by initial arriving EMS professional on scene before moving patients or initiating transport, unless the use of a mechanical chest compression device has been established and is providing effective compressions.
  - Patient movement on stretchers prevents effective CPR
  - Effective CPR cannot be safely performed in a moving ambulance**

*CPR assist device must be an FDA approved device approved for use by the Delaware Office of Emergency Medical Services and coordinated with the county EMS medical director and county paramedic service.

**For patient care and provider safety, the EMS medical directors advocate the use of an optional mechanical chest compression device.
VENTRICULAR ASSIST DEVICE (VAD OR LVAD)

INDICATIONS: A ventricular assist device is a surgically implanted mechanical pump that is used to support blood flow and heart function in people who have weakened hearts. The device pumps blood from the lower chamber of the heart to the body and vital organs. This protocol applies to any medical emergency where the patient has one of these devices.

- Follow General Patient Care Protocol.
  - Unresponsive and not breathing, initiate compressions.
    - LVAD is not a contraindication to CPR
- The patient and family are trained on this device. Listen to and document their guidance.
- Listen to heart sounds. In a functioning device, you should hear a continuous whirling sound.
- Locate the device, usually found at the patient’s waist. Look at the controller and identify which device is being used. Contact the emergency number provided if your patient is unstable, alarms are activated, or the device is off.
- If patient has altered mental status, refer to the Altered Mental Status Protocol.
- Consider Respiratory Distress Protocol.
- Find their backup bag and keep it with the patient.
- Vitals
  - Pulse may not be palpable in these patients.
  - Manual blood pressure may not be obtainable. Utilize an automated cuff to determine blood pressure.
  - Pulse oximetry may not be accurate due to continuous flow from VAD.
- Refer to color code on the VAD to the color on the guidance chart.
- Consider closest appropriate VAD/LVAD facility within geographic operational area.
- Questions or concerns, Contact Medical Control.
<table>
<thead>
<tr>
<th>Question</th>
<th>HeartMate II</th>
<th>HeartWare</th>
<th>Jarvik 2000 Flowmaker</th>
<th>HeartMate XVE</th>
<th>Thoratec PVAD/IVAD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Can I perform CPR?</td>
<td>Only if absolutely necessary</td>
<td>Yes, but risk of dislodging device is high</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Is there a hand pump or external device?</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes Pump @ 60-90 bpm Blue &amp; Red bulbs</td>
</tr>
<tr>
<td>If the device slows down, will an alarm go off?</td>
<td>Yes Audible &amp; flashing light</td>
<td>Yes Audible &amp; flashing light</td>
<td>Yes Audible &amp; solid red light</td>
<td>Yes Audible &amp; solid red light</td>
<td>Yes</td>
</tr>
<tr>
<td>Can I speed up the rate of the device?</td>
<td>No</td>
<td>No</td>
<td>Indicator dial</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Can I use an AED?</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Does the patient have a pulse with this device?</td>
<td>May be weak or non-palpable</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>What is an acceptable Mean arterial pressure?</td>
<td>70-90 mmHg</td>
<td>75-90 mmHg</td>
<td>65-75 mmHg</td>
<td>Normal BP 110/80-140/80</td>
<td>Normal BP readings</td>
</tr>
<tr>
<td>What color is the battery alarm?</td>
<td>Yellow to red</td>
<td>Solid yellow to flashing red</td>
<td>Battery light yellow to red</td>
<td>Solid Yellow - Solid Red battery light</td>
<td>Blinking yellow light</td>
</tr>
<tr>
<td>What color is the low flow hazard alarm?</td>
<td>Red heart flashing</td>
<td>Yellow triangle flashing</td>
<td>Red Solid Red Stop Sign w/ bell on the interior</td>
<td>Solid Red Heart</td>
<td>Solid Red Light</td>
</tr>
<tr>
<td>Are there any transport limitations?</td>
<td>No</td>
<td>Ground transport only</td>
<td>No</td>
<td>10,000 feet elevation maximum</td>
<td>10,000 feet elevation maximum</td>
</tr>
<tr>
<td>How long do batteries last?</td>
<td>Black-3 Hours</td>
<td>4-6 Hours Charge indicator light present on battery</td>
<td>10 Hours</td>
<td>8 Hours</td>
<td>2 1/2 Hours</td>
</tr>
</tbody>
</table>

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CARDIAC ARREST (PEDIATRIC)

INDICATIONS: Current AHA guidelines reflect the importance of compressions for survival from cardiac arrest. EMS practice must evolve to address this important change.

- Compressions should begin as soon as possible following EMS arrival.
- Treating the patient where they are found allows compressions to be started without delay. Only provider safety issues should prompt patient movement.

**High-quality CPR**

- Crews should perform continuous compression PIT CREW HIGH PERFORMANCE CPR. Switch compressors every 2 minutes
  - No pauses for ventilations
  - Ventilations on the upstroke of CPR
- No procedure should slow or stop compressions
- Interruption for defibrillation should be minimal and compressions should resume AS SOON AS shock delivery is complete.
- Frequently switch providers performing chest compressions to maintain peak performance

Compressions should be FAST and HARD at a rate of 100-120 compressions per minute and with sufficient force to depress at least one third the anterior posterior (AP) diameter of the chest, as follows:

- 1 ½ inches (4 cm) in infants
- 2 inches (5 cm) in children
- Ensure complete recoil of the chest wall prior to the next compression

**Ventilations**

- Ventilate at 20-30 breaths per minute with low volume to decrease intrathoracic pressure
- Patients should be bagged using one-hand squeeze
- Ventilations should be just enough to see chest rise
- Avoid excessive ventilation

**Complete a minimum of 20 minutes of high-quality CPR performed by initial arriving EMS professional on scene before moving patients off scene or initiating transport.**

- Patient movement on stretchers prevents effective CPR
- Effective CPR cannot be safely performed in a moving ambulance
DO NOT RESUSCITATE ORDERS

INDICATIONS: Current guidelines for do not resuscitate orders. If any issues arise with orders listed below, Contact Medical Control Immediately.

Living Will*
- Living wills do not apply to out-of-hospital care.
- A living will has no impact on the decision of whether or not to initiate or continue resuscitative efforts or any other care.

Do Not Resuscitate Order (DNR)*
- Contact Medical Control immediately.

Prehospital Advance Care Directive (PACD)*
- Contact Medical Control immediately.

Delaware Medical Orders for Life-Sustaining Treatment (DMOST)*
- A DMOST form is a medical order sheet based on the person’s current medical condition and wishes. Refer to Appendix H.
- The DMOST form will clearly indicate the patient’s wishes concerning life-sustaining treatment and CPR.
- Any section not complete implies full treatment for that section.
- DMOST is used for patients with serious illness or fraility whose death within the next year would not be considered an unexpected event.
- The DMOST form may be voided at any time by a patient with decision making capacity or by their designated representative in cases where the patient lacks this capacity.

Treatment Guidelines: No matter what is chosen, the patient will be treated with dignity and respect, and health care providers will offer comfort measures.

Section A: Goals of Care
- Outlines specific goals to be achieved through the use of this treatment plan

Section B: Cardiopulmonary Resuscitation (CPR)
- If the patient does not have a pulse and is not breathing
  - “Attempt resuscitation/CPR” – provide full care
  - “Do not attempt resuscitation/CPR” – no CPR, no defibrillation/AED, no ACLS

Section C: Medical Interventions
- If the patient has a pulse and is breathing
  - “Full Treatment” – provide all appropriate medical interventions including intubation and mechanical ventilation. Transfer to a hospital if necessary.
  - “Limited Treatment” – use appropriate medical treatment such as IV fluids and oxygen. Do not use intubation or mechanical ventilation. May use noninvasive positive airway pressure (CPAP)
    - Under the “Limited Treatment” option the patient may also elect to indicate
their desire for transport

- **Transfer to hospital for medical interventions**
- **Transfer to hospital only if comfort need cannot be met in current setting**
  - “Treatment of Symptoms Only/Comfort Measures” - use medications, including pain medications, by any route, positioning, wound care, and other measures to keep clean, warm, dry, and comfortable. Use oxygen, suctioning, and manual treatment of airway obstruction as needed for comfort. Transfer only if comfort needs cannot be met in current setting.
  - “Other Orders” – outlines any other specific treatment wishes. Patients already receiving long-term mechanical ventilation or other interventions may indicate treatment limitations here.

**Section D: Artificially Administered Fluids and Nutrition**
- Generally, does not impact pre-hospital care

**Section E: Review of Orders with Patient**
- Documents that orders were reviewed with patient or their representative
- Identifies who is authorized to act as a representative for patients lacking capacity to do so on their own behalf
- Specifies if the authorized representative may not change or void the DMOST form

**Section F: Signatures**
- EMS provider must review this section to ensure it is signed by the patient (or their authorized representative) and the healthcare provider

*If a question should arise regarding DNR’s, PACDs, DMOST or living wills at any time during treatment, **Contact Medical Control.**
PEDIATRIC AND ADULT TRAUMA

INDICATIONS: This Trauma Protocol applies to patients with any of the following field triage criteria. If any of the conditions in abnormal vital signs, obvious injury are present or mechanism of injury/ evidence of high energy impact, transport to a Trauma Center. Consider air medical transport.

Vital Signs:

Adults: Glasgow Coma Scale less than 13.
Systolic BP less than 90 mmHg.
Respiratory rate less than 10 or greater than 29.

Pediatrics: Pediatric Glasgow Coma Scale less than 13
Refer to the Vital Signs in Appendix B.

Patients with abnormal vital signs should be transported to the closest appropriate Trauma Center.

When in question, Contact Medical Control.

Patients with GCS less than 13 or exhibiting new onset paralysis or paresis: consider direct transport to a Level I or II Trauma Center (Maryland Level III) with neurosurgical capabilities.

If NO for all elements in Vital Signs, proceed to Obvious Injury.

Obvious injury:

Penetrating injury to the torso, axilla, abdomen, head, neck, proximal extremities or groin.
Major burns, inhalation injury, or trauma with burns.
More than one proximal long bone fracture.
Pelvic fracture (suspected on clinical grounds).
Flail chest or other major chest injury
Limb paralysis
Major external hemorrhage.
Amputation above wrist or ankle
Crushed, degloved or mangled extremity
Open or depressed skull fracture
AVPU scale: does not respond to voice

Patients with obvious injury should be transported preferentially to the closest appropriate Trauma Center.

After evaluation of injuries proceed to the Mechanism.

Mechanism:

Patient ejection (partial or complete) from vehicle
Motorcycle crash greater than 20 mph or rider thrown
Death of passenger in same vehicle compartment
Falls greater than 20 feet (adult)
Falls greater than 10 feet (child) or 2-3 times the height of the child
Auto-pedestrian/ auto-bicycle injury-thrown, run over or with significant (greater than 20 mph) impact
Vehicle telemetry consistent with high risk injury
High risk auto crash: inner intrusion greater than 12” occupant greater than 18” anywhere
Patients with any of the above mechanisms should be transported preferentially to the highest level trauma center practical.

If NO for all elements in Mechanism, proceed to Extenuating Circumstances.

**Extenuating Circumstances:** *(Not stand alone criteria for the initiation of trauma protocol or helicopter transport.)*

- Pregnancy greater than 20 weeks
- Renal dialysis
- Age less than 15 or greater than 55 years
- Other significant medical conditions - discuss with Medical Control
- Time Sensitive extremity injuries
- Required by patient condition in the judgment of the prehospital provider
- Anticoagulation medications and bleeding disorders (Factor deficiencies, ITP).

If YES to extenuating circumstances, Contact Medical Control and transport to the closest appropriate Trauma Center with necessary resources.

If NO to all above, routine transport. When in doubt, transport to the closest appropriate Trauma Center.

Consider Pediatric and Adult airway management protocol.

For suspected unstable pelvic fractures, apply pelvic compression device per manufacturer instructions. If an appropriate pelvic compression device is not available, use a sheet to apply compression.

For burns, refer to Burn protocol.

In cases of severe hemorrhage:
- Apply direct pressure to the hemorrhaging wound
- If direct pressure is not adequate to control hemorrhage, a provider may use a tourniquet for hemorrhage that is anatomically amenable to tourniquet application and note time of application.
- The iTClamp may be utilized when available for hemorrhage control when appropriate.
- If sucking chest wound, cover wound with a commercial chest seal. If none are available, cover wound with an occlusive dressing sealed on 3 sides. Release dressing if worsened shortness of breath.
- If evisceration, cover with sterile dressing moistened with sterile saline or water; cover the area with an occlusive material (plastic wrap). Cover the area with a towel or blanket to keep warm. DO NOT PUSH VISCERA BACK INTO ABDOMEN. Transport with knees slightly bent.

For hemorrhage that cannot be controlled with above, apply approved hemostatic agent with direct pressure, or through packing of the wound with gauze either impregnated with hemostatic agent or not. If packing the wound, gauze must be inserted deeply and fully and can include multiple rolls of gauze. Consider application of an Abdominal Aorta Junctional Tourniquet (AAJT).

- Patients with hemorrhagic shock should be taken to the closest appropriate Trauma Center, without delay.
- Head or spinal trauma patients with GCS less than 13 or exhibiting new onset paralysis or paresis; consider direct transport to a Level I or II trauma Center (or Level III Maryland Trauma Center).
- Patients who are less than 15 years of age should be transported to a pediatric trauma center when patient condition, time and distance allow.
• Burn patients should be evaluated at the closest appropriate Trauma Center.
• Consider helicopter transport if ground transport to the appropriate Trauma Center is expected to exceed 10 minutes.
• Unstable penetrating trauma patients should go directly to the closest appropriate Trauma Center.
• Patients in shock with deteriorating vital signs or ongoing airway compromise should be transported to the closest appropriate Trauma Center.

Trauma scene times should be less than 10 minutes unless there are extenuating circumstances. Reasons for scene times over ten minutes should be documented in the chart. Appropriate reasons for prolonged trauma scene times include extrication, awaiting BLS, securing scene safety, presence of multiple victims, awaiting helicopter touch down for transport to a higher level trauma center, etc.

**Penetrating Trauma**  
*Gunshot, knife or impaling injury to the neck, torso or proximal extremity.*

• Scene time: goal less than 5 minutes.

• Scene activities:
  o Hemorrhage control with tourniquets, hemostatic agents, wound packing, Junctional tourniquets and iTClamps.
  o Open airway, apply nasal or oral airways as indicated
  o Rapid movement to transport unit

• Enroute activities:
  o Expose and continue with trauma assessment
  o Call medical control for rapid short report:
    ▪ Arriving BLS only
    ▪ Age & sex
    ▪ Mechanism
    ▪ Vital signs (HR & BP)
    ▪ No Airway & IV management
    ▪ ETA
    ▪ Expect room assignment
PAIN MANAGEMENT (ADULT)

INDICATIONS: For the management of acute pain caused by musculoskeletal injury (fractures, dislocations, sprains, strains) who are conscious and alert, able to follow commands and maintain the patency of their own airway.

CONTRAINDICATIONS:

- Liver disease.
- Allergy to acetaminophen.
- Patient age less than 15 years.
- Use of Acetaminophen containing medication in the last 4 hours.

- Follow General Patient Care Protocol.
- Obtain a Pulse Oximeter reading. Provide appropriate supplemental oxygen titrated to greater than 92%.
- Assess and record pain severity using 0-10 scale.
- If appropriate, consider the use of non-pharmacological pain management techniques:
  - Place patient in a position of comfort.
  - Apply ice packs and/or splints for suspected fractures or dislocations.
  - Provide verbal reassurance to control anxiety.
- If pain management efforts are unsuccessful, consider the administration of acetaminophen 1000 mg PO.
- Reassess patient including changes to their pain severity score.
- In cases of severe pain uncontrolled through the above measures, consider requesting ALS for further pain management therapy.
- For patients experiencing nausea with their pain, consider Ondansetron (Zofran) 8mg sublingual (oral dissolving tablets) unless otherwise contraindicated.
- Transport in a position of comfort and continue re-assessment enroute.
SELECTIVE SPINAL MOTION RESTRICTION

INDICATIONS: Apply this guideline to all patients involved in known or suspected blunt trauma.

Implement spinal motion restriction (rigid collar) in the following circumstances:

- Significant multiple system trauma.
- Severe head or face trauma.
- If altered mental status (including drugs, alcohol and trauma) and:
  - No history available
  - Found in setting of possible trauma (e.g., lying at the bottom of stairs or in street)
- Loss of consciousness after trauma.
- Any fall with evidence of striking head.
- Spinal pain or tenderness, including any neck pain with a history of trauma.
- Numbness or weakness in any extremity after trauma
- Patient with significantly painful distracting injury.

For Patient transport:

- If ambulatory, allow patient to move to stretcher mattress with minimal spinal motion
- If non-ambulatory, consider using backboard, scoop/orthopedic stretcher, vacuum mattress, or other device for transfer of patient to stretcher with minimal spinal motion
- In certain situations, the long backboard will still be used as an extrication/moving device but plays no significant role in restricting spinal motion. If a backboard is utilized during extrication, the EMS crew may, at its discretion, remove the board prior to transport.
- Transport on stretcher mattress without backboard if patient is ambulatory or if scoop/orthopedic stretcher and be removed with minimal patient motion.
- Use of a scoop/orthopedic stretcher, backboard or Reeves stretcher is required for patients being transported by pre-hospital aviation.
- New onset neurological deficit, consider aviation for transport to Level 1 trauma center.

Note: Penetrating trauma to the extremities or core (below the clavicles) without neurologic deficit does not require a rigid collar.

See Selective Spinal Motion Restriction Algorithm below.
Neuro Exam: Any focal deficit?

- **NO**

Significant mechanism of Injury? ****
High-energy events such as ejection, high falls, and abrupt deceleration crashes

- **NO**

Alertness: Alteration in mental status?

- **NO**

Intoxication: Any evidence?
Includes Drugs and / or Alcohol

- **NO**

Distracting Injury:
Any painful injury that might distract the patient from the pain of a c-spine injury?

- **NO**

Spinal Exam:
Point tenderness over the spinous process(es) or pain to ROM?

- **NO**

Spinal Motion Restriction
Not Required

**YES**

Apply rigid collar

Restrict Spinal Movement

**AMBULATORY**
Bring stretcher to patient, assist patient onto stretcher with minimal spinal movement, and then secure patient to stretcher.

**NONAMBULATORY**
Use Long Spine Board (OR any of the multiple equivalent devices) to TRANSFER patient to stretcher with minimal spinal movement, consider removing the device, then secure to stretcher.

May use multiple providers to transfer patient to stretcher using in-line spinal techniques such as log roll / straddle slide to maintain spinal precautions without a device, then secure to stretcher.

**NOTE:** For patients greater than 65 years of age, consider a lower threshold.
BURNS

INDICATIONS: A patient who has been exposed to radiation, thermal, electrical, environmental, or chemical reactions that cause burns

- Follow Pediatric and Adult Trauma Protocols.
- Stop the burning process. If safe to do so:
  - Decontaminate if chemical or radiological burn.
    - Obtain SDS or other specific chemical data, if possible and timely
    - Stay on scene to irrigate if required
  - Ensure patient is not energized if electrical or lightning burn prior to entering the area.
- If patient is in cardiac arrest, utilize cardiac arrest protocol.
- Consider ALS for pain management.
- Monitor for airway burns; if any of the following signs or symptoms are present, request ALS:
  - Singed facial or nasal hairs
  - Hoarse voice or stridor
  - Difficulty breathing
  - Carbonaceous sputum
  - Burns on face.
- Remove jewelry and clothing; completely expose burned area.
- Assess burn percentage using the "Rule of Nines." Refer to Appendix B.
- Bandage burned areas using clean, dry sheets.
- DO NOT pop/ lance any blisters that form.
- Cover the patient and provide for an appropriate warm environment to prevent heat loss.
- Friction burns “Road Rash” type injuries
  - Cover with clean, dry sheets
  - Document injuries
TRIAGE FOR MASS CASUALTY

**INDICATIONS:** If a victim appears to be a young adult, use START. If a victim appears to be a child, use Jump START.

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**Simple Triage And Rapid Treatment (START)**

**Step 1:**
Triage officer announces that all patients that can walk should get up and walk to a designated area for eventual secondary triage. All ambulatory patients are initially tagged as Green.

**Step 2:**
- Triage officer assesses patients in the order in which they are encountered
- Assess for presence or absence of spontaneous respirations
- If breathing, move to Step 3
- If apneic, open airway
- If patient remains apneic, tag as Black
- If patient starts breathing, tag as Red

**Step 3:**
- Assess respiratory rate
- If less than 30, proceed to Step 4
- If greater 30, tag patient as Red

**Step 4:**
- Assess capillary refill
- If less than 2 seconds, move to Step 5
- If greater than 2 seconds, tag as Red

**Step 5:**
- Assess mental status
- If able to obey commands, tag as Yellow
- If unable to obey commands, tag as Red

**Jump START**

**Step 1:**
Identify and direct all ambulatory patients to designated Green area for secondary triage and treatment. Begin assessment of non-ambulatory patients as you come to them.

**Step 2:**
- If breathing spontaneously, go on to the next step, assessing respiratory rate.
- If apneic or with very irregular breathing, open the airway using standard positioning techniques.
- If positioning results in resumption of spontaneous respirations, tag the patient immediate and move on.

**Step 3:** Jump
If no breathing after airway opening, check for peripheral pulse. If no pulse, tag patient deceased/non-salvageable and move on.
If there is a peripheral pulse, give 5 mouth to barrier ventilations. If apnea persists, tag patient deceased/non-salvageable and move on.
• If breathing resumes after the “jumpstart”, tag patient immediate and move on.

Step 4:
• If respiratory rate is 15-45/min, proceed to assess perfusion.
• If respiratory rate is less than 15 or greater than 45/min or irregular, tag patient as immediate and move on.

Step 5:
• If peripheral pulse is palpable, proceed to assess mental status.
• If no peripheral pulse is present (in the least injured limb), tag patient immediate and move on.

Step 6:
• Use AVPU scale to assess mental status.
• If Alert, responsive to Verbal, or appropriately responsive to Pain, tag as delayed and move on.
• If inappropriately responsive to Pain or Unresponsive, tag as immediate and move on.

Modification for non-ambulatory children
• Infants who normally can’t walk yet
• Children with developmental delay
• Children with acute injuries preventing them from walking before the incident
• Children with chronic disabilities

Step 7:
• Evaluate using the Jump START algorithm
• If any RED criteria, tag as RED.
• If pt. satisfies YELLOW criteria:
  o YELLOW if significant external signs of injury are found (i.e. deep penetrating wounds, severe bleeding, severe burns, amputations, distended tender abdomen)
  o GREEN if no significant external injury

Step 8:
Unless clearly suffering from injuries incompatible with life, victims tagged in the BLACK category should be reassessed once critical interventions have been completed for RED and YELLOW patients.

See Appendix I: Mass Casualty Triage Chart
TASER PROBE REMOVAL

**INDICATIONS:** Patient with uncomplicated probes from a conducted electrical weapon (Taser®) embedded into non-sensitive areas of the skin.

**CONTRAINDICATIONS:**

Patients with probe penetration in vulnerable areas of the body (female breast, genitalia, or above the level of the clavicles) or suspicion that the probe is imbedded into bone, blood vessels, or other sensitive area; should be transported for evaluation and probe removal at the receiving medical facility.

- Follow General Patient Care Protocol and/or Trauma Care Protocols
- Ensure wires are disconnected from the weapon.
- Stabilize skin around probe using non-dominant hand. Grasp the probe by metal body using your dominant hand. Remove probe in a single and quick motion.
- Dispose of taser probe in sharps container.
- Clean the wound with antiseptic wipe and apply an appropriate dressing.

*Patient may refuse EMS transport if all other refusal requirements are met.*
WATER RELATED EMERGENCIES

INDICATIONS: Victims of non-fatal drowning, history of diving or breathing compressed air. Symptoms include altered mental status, numbness, pain in extremities, joint pain, blurred vision, blood from nose ears and mouth, seizures, unresponsiveness.

- Consider Pediatric and Adult Trauma Protocols.
- Remove wet clothing. Keep patient warm and dry.
- Assess for hypothermia and take actions to maintain warmth.
- Any submersion injury patient should be transported for evaluation regardless of current assessment findings.
- Be alert for vomiting.

Note: Turnout gear is contraindicated for water rescues. Personal flotation devices should be worn within 10 feet of the water’s edge.
EYE INJURIES

INDICATIONS: Injury to eye(s) or sudden vision impairment. Includes foreign body irritants, bright light exposure such as UV or welding, chemical irritation, dislodgement, or penetrating injury.

- Utilize Pediatric and Adult Trauma Protocol

- Penetrating Trauma:
  - Do no palpate or apply pressure to the eye
  - Shield injured eye
  - Patch uninjured eye for adult patients only
  - DO NOT remove impaled objects. Protect the eye from further injury.
  - Avoid unnecessary movement. Advise patient not to cough, sneeze, or move more than absolutely necessary.

- Dislodged Eye:
  - Cover with cup and secure

- Ultraviolet light exposure (arc welder, sun lamp burn)
  - Place cool compress lightly over both eye lids
  - Symptoms may be delayed 3-10 hours

- Sudden loss of vision
  - Assess for trauma and transport.

- Chemical Burn
  - Irrigate eye with normal saline or water (as appropriate for the chemical involved) for 20 minutes
  - If initiating transport will not interrupt eye irrigation, irrigate enroute to the hospital
  - Notify the hospital of the chemical involved
BITES AND ENVENOMATION

INDICATIONS: Bite by a suspected poisonous snake or spider with puncture marks to the skin and accompanied by swelling, pain, warm skin around bite area. A rash, wound, oozing blood, evidence of infection, difficulty breathing, wheezing, shortness of breath, hives, itching, hypotension or other signs of shock may be present. Animal bites or human bites received where the skin is broken.

- Follow General Patient Care Protocol.
- Request Animal Control if needed
- Obtain a description of the insect/animal in question
  o Picture or adequate description
- Do Not transport insect/animal in question with you to the hospital.

Venomous Bites

- Contact Medical Control
- Transport to closest facility for further evaluation
- Note the time, location size of bite/sting
- Mark the area of swelling lightly with a pen or marker to assist with ongoing evaluation
- Remove any constricting clothing or jewelry near the bite area.
- Immobilize the area of injury.
- DO NOT apply ice on the wound
- DO NOT apply any constriction or tourniquet.

Human bites to a provider should be treated as an infectious disease exposure and reported appropriately.
HEAT EMERGENCIES

**INDICATIONS:** Exposure to high temperatures, hot skin, tachycardia, altered mental status, dizziness, nausea, headache, mild hypotension, weakness/fatigue, thirst, or muscle cramps.

- Follow General Patient Care Protocol.
- Remove excess clothing
- Place patient in a cool environment
- If skin is normal to the touch: apply cool compresses. Dampen towel and apply to the head & neck areas.

**Heat Stroke (Life Threatening Emergency)**

- Consider ALS
- Perform active cooling of patient.
  - Cold Water Immersion on scene if available
  - Pouring cool water over the patient is also acceptable
  - Ambulance AC on coldest setting, fans on highest setting
- If patient begins to shiver, slow the cooling process

**Heat Exhaustion**

- Allow oral intake of cool fluids (water or sports drink only) if the patient is alert and oriented and has patent airway.

**Heat Cramps**

- Allow oral intake of cool fluids (water or sports drink only) if the patient is alert and oriented and has patent airway.
COLD EMERGENCIES

INDICATIONS: Patient exposed to cold environmental conditions. This may be acute or over a period of time. Body temperature less than 96.8° F (36° C), frostbite, submersion in cold water.

- Follow General Patient Care Protocol.
- Remove wet clothing and replace with warm blankets.
- Utilize heat packs on the exterior of blankets in the neck, arm pit and groin areas.
- DO NOT place heat packs, hot water bottles, IV bags, or other heat-retaining devices directly on the patient’s skin.
- Utilize cardiac arrest protocol if needed. Avoid prolonged pulse checks.
- If patient is in cardiac arrest from drowning or hypothermia; ensure the patient is dry and packaged with blankets.

Frostbite

- Keep patient warm while exposing the affected part.
- DO NOT rub the affected part, permit the patient to use the affected part, puncture blisters, expose part to dry heat, or immerse part in hot water.
- Remove jewelry if near the affected area.
- Apply loose clean dressing to the affected area.
OBSTETRICS

INDICATIONS: Patients for whom delivery is imminent as evidenced by crowning, bloody show, breech or limb presentation. OB patients not to full gestation (40 weeks) should be transported to the closest appropriate facility.

- Follow General Patient Care Protocol.
- Gather obstetrical history
  - number of times pregnant (gravid)
  - number of living children (para)
  - estimated due date
  - any known problems, # of children in current pregnancy (twins, triplets, etc.)
  - assess length of contractions and amount of time between contractions
- If delivery is not imminent, transport patient in a left lateral recumbent position to prevent undue pressure from being exerted on the lower vena cava by the gravid uterus.
- If delivery is imminent, remove clothing below the mother's waist; check for crowning as appropriate.
- Create a clean field for the delivery area utilizing OB kit.
- If patient is experiencing urge to push or to have a bowel movement, prepare to deliver.
- Assist with delivery.
- If complications arise, request ALS and Contact Medical Control immediately.
- If meconium staining is present, request ALS.
- Only suction mouth with bulb syringe in the presence of airway compromise.
- **DO NOT** pull on infant or put traction on umbilical cord. Guide & control the infant out.
- Maintain newborn’s body temperature. If mother capable, place infant skin to skin to preserve body temperature. Consider emergency blanket, silver swaddler, or other heat retaining device. If mother not capable of skin-to-skin contact use the above blankets to preserve warmth. Ensure the head is covered, but do not cover the newborn’s face.
- Dry the newborn with a clean towel.
- Stimulate the infant to cry by rubbing the back.
- Clamp the umbilical cord after newborn is dry; 6 inches away from the newborn’s navel area and second clamp 2 inches farther. Cut cord between the clamps.
Monitor newborn's heart rate, respirations, and oxygen saturation. Consider blow-by oxygen 5-6LPM. Normal pulse is 120-160/minute and normal respiratory rate is 40-60/minute.

If heart rate less than 100 beats per minute, gasping or apnea, provide BVM ventilation with oxygen at a rate of 40-60 breaths/minute; Request ALS.

If heart rate less than 100 beats per minute and the newborn is cyanotic or has labored breathing, without gasping or apnea, clear the airway, check for anatomical issues to BVM compliance.

If heart rate is less than 60 beats/ minute perform CPR; Request ALS

Obtain APGAR scores at 1 minute and 5 minutes after delivery

Re-assess newborn every 30 seconds for change in heart rate or respiratory status. If no change at 1 minute, provide BVM ventilation and request ALS.

Obtain a blood glucose reading via heel stick from the newborn. If reading is less than 40mg/dl, request ALS for hypoglycemia.

Transport to the hospital. Consider safe transport methods.

If multiples or cardiac arrest/respiratory situation, request additional resources.

Treating the Mother after Delivery

Place mother in a position of comfort following the delivery. Prepare for placenta.

Apply direct pressure to any external bleeding only.

If placenta delivers, place in a red bag, seal it, and keep it with the mother.

Perform fundal massage if cramping or bleeding persists.

Utilize bulky gauze dressing to absorb any bleeding. Do not insert these into the vagina or anus.

If the mother’s BP drops below 90mmHg systolic during the delivery, place her in a left lateral recumbent position if possible and re-assess BP.

Limb Presentation

If a limb presents, treat as a life-threatening emergency.

Request ALS

Place the mother in a knee to chest position on the stretcher

Transport emergently to the hospital.
Breech Delivery

Deliver newborn as above with the following change: After chest has delivered, place a gloved hand in the vaginal opening to create an airway for the newborn, providing a V-shaped area around the infant’s mouth. Ensure the cord is not wrapped around the newborn’s head.

Cord Prolapse

- Place the mother in a knee to chest position on the stretcher.
- Keep cord moist with saline soaked gauze.
- Attempt to relieve pressure on the cord by gently lifting the presenting part off the cord.
- Ensure a pulse can still be felt in the cord.

Nuchal Cord

If the cord is around the newborn’s neck, it must be released from the neck immediately. Slip cord over newborn’s delivered head or shoulder. If unable to slip the cord free, clamp and cut the cord.

*It is normal for a newborn to have an oxygen saturation of 60-65% in the first minute of life; 65-70% in the second minute of life; the oxygen saturation at 5 minutes is 80-85%; at ten minutes oxygen saturation is 85-95%.*
PATIENT RESTRAINT

INDICATIONS: Patient care remains the primary responsibility of the EMS provider. The method of restraint shall not restrict the adequate monitoring of vital signs, ability to protect the patient’s airway, compromise peripheral neurovascular status or otherwise prevent appropriate and necessary therapeutic measures. It is recognized that evaluation of many patient parameters requires patient cooperation and thus may be difficult or impossible.

Soft restraints are to be used only when necessary, in situations where the patient is potentially violent and may be of danger to themselves or others. Patients who have clinical capacity retain a right to refuse transport. Soft restraints are padded or leather wrist or ankle straps. Plastic ties, ropes, and handcuffs are not considered soft restraints. EMS providers must remember that aggressive violent behavior may be a symptom of medical conditions such as but not limited to:

- Head trauma
- Alcohol/drug related problems
- Metabolic disorders (i.e., hypoglycemia, hypoxia, etc.)
- Psychiatric/stress related disorders

- All restraints should have the ability to be quickly released, if necessary, in an emergency.
- If a spit shield/hood is in place prior to EMS arrival, it may remain in place unless medically necessary to remove.
  - Ensure no obstruction to airway or breathing
- Any patient in handcuffs shall be considered in law enforcement custody and require law enforcement presence to maintain custody of that individual. It is medically acceptable to have a law enforcement officer follow a restrained patient’s ambulance to the hospital in their law enforcement vehicle, as long as they maintain a position and contact with the transporting ambulance that will allow the officer to quickly release any restraining device that requires a key or special releasing device that they have applied in the event of a sudden deterioration in a restrained patient’s condition.
- Patients should be transported in the supine position to ensure adequate respiratory and circulatory monitoring and management.
- The prone position shall not be utilized and is not permitted under this protocol. This position carries a higher risk of patient injury or death.
- All restrained patients should be placed on a stretcher with adequate foam padding particularly underneath the head. Extremity restraints should be secured to the stationary portion of the stretcher frame.
- Stretcher straps should still be placed on all patients as these are similar to seatbelts during transport.
- Restraints that use multiple knots or that may restrict chest wall motion are unacceptable.
- Restrained extremities should be monitored for color, sensory and motor function, pulse quality, and capillary refill at the time of application and at least every 15 minutes thereafter. The patient's respiratory status and pulse oximetry should be monitored during transport.

- Consider requesting ALS for chemical sedation if patient is in continued agitated state.

- Restraint documentation on the EMS PCR shall include:
  - Reason for restraint
  - Agency responsible for restraint application (i.e., EMS, Police)
  - Documentation of vital signs, pulse ox, capillary refill and peripheral neurovascular status (motor/sensory).

- Medical Control must be contacted if a patient is deemed too violent or uncooperative to be safely transported using the restraint methods and devices permitted by their prehospital protocols.

*This policy is not intended for the Interfacility transport of medically cleared involuntarily committed psychiatric patients.*
FIRE GROUND REHABILITATION

INDICATIONS: The intent of rehabilitation (Rehab) is to provide a structured, consistent method for the evaluation and remediation of common ailments associated with the activities at fire/hazardous materials and incident scenes; including but not limited to overexertion, dehydration, metabolic disturbances, and exposure to temperature extremes. This protocol shall be used in a rehabilitation area/sector or during medical monitoring of personnel at an incident scene.

- Remove member from climactic conditions.
  - Rest area should be free of smoke and sheltered from extreme heat or cold.

- Provide rest and recovery period.
  - Member should be afforded the ability to rest for at least 10 minutes or as long as needed to recover work capacity.

- Provide cooling or rewarming as required.
  - Remove PPE if warm. Provide blankets if cold. Assess for environmental emergencies.

- Provide fluid replacement.
  - On scene, potable fluids should be provided so members can satisfy thirst.

- Provide calorie and electrolyte replacement when appropriate (extended operations).
  - Whenever food is available, means for members to wash their hands and face must be provided.

- Provide medical monitoring. Evaluation should include, but not limited to:
  - Presence of chest pain, dizziness, shortness of breath, weakness, nausea or headache.
  - General complaints such as cramps or aches and pains.
  - Symptoms of heat or cold related stress.
  - Changes in gait, speech or behavior.
  - Alertness and orientation to person, place and time.

  - Baseline vital signs should be established when the member first enters the rehabilitation area. Vital signs should include pulse, respirations, blood pressure, pulse oximetry and carbon monoxide assessment. Prior to exiting the rehabilitation area, another set of vital signs should be obtained. These vital signs should show trending evidence of improvement over the baseline set of vital signs.

- Treat abnormal findings in accordance with local protocols.
  - When EMS treatment or transport is provided, a patient care report must be generated.

- Member accountability.
  - Accountability should be maintained to track members as they enter and leave rehabilitation.

- Release
  - Prior to leaving rehabilitation, the member, as well as EMS, should agree that the member is adequately rested and able to safely perform full duty if released.
NON-EMERGENCY / INTERFACILITY / DEVICE DEPENDENT TRANSPORT

INDICATIONS: To provide emergency medical services safely and without delay to the patients requiring transfer from one medical facility to another medical facility, for patients being discharged from a medical facility to a home residence, patients that are direct admissions to the hospital that bypass the Emergency Department, or for patients being transported to and from a routine medical appointment.

These protocols are not intended to indemnify the medical sender or receiving facilities from their obligations under the EMTALA statutes.

This protocol is only intended for use for patients that otherwise meet basic life support transport criteria. Transport personnel are not authorized and will not provide services beyond their scope of care. Should services beyond scope be required, a person qualified in its performance shall accompany the patient during transport.

Providers dispatched for a non-emergency transport that encounter a patient experiencing an acute emergency are responsible for providing patient care or transferring the patient to a higher-level provider, if needed, and transporting the patient to an emergency department.

Temporary intravenous medications like antibiotics, intravenous drip medication that require frequent monitoring and maintenance, or intravenous pumps that are not part of the patient’s long-term care plan are excluded from this protocol. These excluded medications require advanced personnel for transport.

- BLS personnel may transport patients who meet the criteria of this protocol.
- Non-emergency Interfacility transports shall not compromise the local 911/EMS resources of the community. It is the responsibility of the ambulance service to determine whether adequate resources are available to maintain appropriate EMS coverage to their community before committing to such transport.
- All Interfacility transports will be documented using the approved prehospital EMS PCR.

Intravenous Fluid Transport

- All patients with an established intravenous (IV) access.
- The destination facility shall be an inpatient facility no more than 60 minutes from the facility of origin if IV fluids are hanging. If a saline/heplock is in place, time constraints do not apply *
- Patients with IV fluids shall have only standard IV fluids (normal saline, ½ normal saline, ringers lactate, or dextrose 5% and water) hanging at the time of transport. The fluids will be set at a Keep Vein Open (KVO) rate by the sending facility and will not have medications or supplements added to the fluid.
- EMT’s may transport IV fluids in place only (no medications).
- The EMT shall not alter the flow rate of the IV fluids unless it is to shut them off in the event of an emergency. IV fluids need to be shut down for the following reasons:

- Swelling, redness, or increased pain at the site of the IV insertion.
- Fluids in the bag have emptied.
- The IV catheter is inadvertently dislodged from the site.
- If complications arise

- Paramedic intervention considered if the patient's condition deteriorates en route as evidenced by unstable vital signs, change in mental status or onset respiratory distress, chest pain, or neurological changes. The EMT is encouraged to **Contact Medical Control** any time questions or concerns arise.

- The goal is to not have the bag of IV fluids empty prior to arrival at the destination facility. In the event this happens, the IV will be shut off for the remainder of the transport.

**Definitions:**

Pumps used by home-bound patients are considered patient administered medication. Home-bound is defined as a residence, rehabilitation unit, or nursing home. These patients may be transported providing the EMT does not have to manipulate or operate the pump and the administration route is through an intravenous line or parenteral nutrition line. If a malfunction arises with the pump or administration line, unplug the pump, turn the power off, and **Contact Medical Control**.

Interfacility transport of patients on medication must be accompanied by an ALS provider (Paramedic, RN, RRT, NP, PA, or MD). This does not apply to home-bound patients receiving patient or family administered medication through an IV or parenteral nutrition line.

**Implanted/ Invasive Device List**

**AICD** (Automatic Internal Cardiac Defibrillator): approved for transport only and may not be manipulated by BLS personnel.

**Completely Implantable Venous Access Port** (Porta-cath, PICC): Used for infusion or long-term medication therapy (antibiotic, chemotherapy, etc.). These may not be accessed by EMT’s. Transport of patients receiving medication requires ALS to accompany the patient.

**Epidural Catheters:** are approved for transport only and may not be manipulated by EMS personnel. If it dislodges apply sterile pressure dressing **Contact Medical Control**.

**Foley Catheter:** are approved for transport only and may not be manipulated by EMS personnel. If the catheter dislodges **Contact Medical Control**.

**Gastrointestinal Tubes:** Approved for transport only and may not be manipulated by EMS personnel. If it dislodges, apply dressing over the site and **Contact Medical Control**.

**Implantable Central Venous Catheters** (Hickman, Broviac): These are surgically implanted for patients requiring long term venous access for meds or dialysis. These may have more than one lumen.

**KVO rate:** 1 drop per minute.

**Percutaneous Drainage Tubes or Surgical Drains** (Vacuum Drains): are approved for transport only and may not be manipulated by EMS personnel. If it dislodges, apply dressing over the site and contact Medical Control.
**Peritoneal Dialysis Catheters**: approved for transport only. If it dislodges, apply sterile pressure dressing and contact Medical Control.

**Ventricular Assist Device (VAD, LVAD)**: are approved for transport and may not be manipulated by EMS personnel except in the event of a device failure. EMS providers should be briefed on the procedures of the specific device before transporting. Request ALS in the event of a device failure and contact the emergency number listed on the device. Follow guidance given and **Contact Medical Control**.

**Ventilators**: Mechanical ventilation is a method to mechanically assist or replace spontaneous breathing. A vent patient is anyone who is dependent on a ventilator to sustain life.

- A vent dependent patient must have someone trained in the operation of the ventilator in use, who is also familiar with the monitoring and management of a patient with ventilator failure. This person may be an ALS provider (Paramedic, RN, NP, RRT, PA, MD) or a family member that has been trained on the device. BLS services are encouraged to check with their insurance agent before providing a transport with a family member in the patient compartment of the ambulance.

- In the event of equipment failure or respiratory emergency, provide appropriate airway management.

- **Medical Control** shall be contacted, and ALS intervention considered if the patient’s condition deteriorates during transport. Deterioration is evidenced by unstable vital signs, change in mental status, or onset of respiratory distress, chest pain, or neurologic changes.
MEDICAL TREATMENT FOR CHEMICAL EXPOSURE

INDICATIONS: Touching, breathing, eating, or drinking harmful chemicals. Exposure to chemicals can result in varying symptoms with different degrees of danger. Mild reactions include burning and tearing of the eyes, throat, nose, chest, and skin. Severe reactions include coughing, wheezing, feeling faint, convulsions and even death.

- Ensure proper decontamination has occurred. Remove patient’s clothing. Be cautious of any off-gassing of chemical from under the clothing that may occur in this process.
- Follow recommendations from the manufacturer SDS within scope of practice protocols.
- Any questions or concerns Contact Medical Control.

Organophosphate poisoning/ nerve agents

Individuals exposed to suspected nerve agents or organophosphates that exhibit signs of SLUDGEM or DUMBBELS. Common names for nerve agents include Tabun (GA), Sarin (GB), and Soman (GD), GF and VX. Included in the organophosphate group are disulfoton, phorate, dimethoate, ciodrin, dichlorvos, dioxathion, ruelene, carbophenothon, supona, TEPP, EPN, HETP, parathion, malathion, ronnel, coumaphos, diazinon, trichlorfon, paraoxon, potasan, dimefox, mipafox, schradan, sevin, chlorpyrifos and dimeton.

| S | Salivation |
| L | Lacrimation |
| U | Urination |
| D | Defecation (diarrhea) |
| G | Gastric emptying (vomiting) |
| E | Emesis |
| M | Miosis |
| D | Diarrhea |
| U | Urination |
| M | Miosis |
| B | Bronchospasm |
| B | Bradycardia |
| E | Emesis |
| L | Lacrimation |
| S | Salivation |

- Administer Duo Dote Auto-Injector Intra-Muscular, if available
- For Severe Symptoms (unresponsiveness, seizure, severe respiratory distress) administer 3 Duo Dote Auto-Injector Intra-Muscular, if available.

Hydrocarbon Exposure

Indications: Anyone doused in gasoline, diesel fuel, kerosene, paint thinner, or other ignitable liquid. These chemicals are absorbed through the skin and harm the liver. The largest hazard is fire.

- Decontaminate the patient. Remove contaminated clothing. In stable patients consider scrubbing skin with soap and water.
- Consider utilizing a hazardous materials decontamination unit.
- Notify the receiving hospital of the exposure and request direction on their decontamination area prior to entering the hospital.

Biological Agents

- Ensure all providers wear N95 masks to protect from inhalation exposure.
- Refer to Medical Control for guidance.
SUSPECTED EMERGING INFECTIOUS DISEASE (EID)

Modified Patient Care Guidelines for COVID-19

INTRODUCTION: An emerging infectious disease (EID) may pose challenges that impact standard medical treatment protocols. COVID-19 is an example of an EID that forces often dynamic changes to the way care is provided to the patients. The following are some suggested treatment modifications during a declared epidemic of a viral illness. (Note: infectious diseases may be dynamic, and providers are encouraged to frequently check the Delaware Division of Public Health website for the most up-to-date information including specifics on presentations, protection methods and epidemiology.)

Personal Protective Equipment (PPE):
- All providers will utilize personal protective equipment (PPE) appropriate to the risks presented – N95 respirator or surgical mask, eye protection / face shield, gown, gloves.
- Don appropriate PPE before proceeding with patient care following current guidance. This includes providing care for all levels of patient acuity (e.g., don PPE before beginning resuscitation of a cardiac arrest with suspected EID).

Airway and Respiratory Considerations:
- Many airway management procedures involve aerosol-producing processes and thus pose potential exposure risks to providers.
- For hypoxic patients, the first attempts to improve oxygenation should be by providing supplemental oxygen.
- Oxygen therapy may be delivered underneath surgical masks.
  - Place a mask over a patient receiving oxygen via nasal cannula or non-rebreather masks.
  - May use higher than normal flow rates (up to 6 lpm) with nasal cannula under surgical mask.
- Advance further into respiratory management (bronchodilators, CPAP, intubation, etc.) only if initial attempts at improving oxygenation are not working.
- If the patient has a prescribed meter-dose-inhaler (MDI) containing a beta-adrenergic agonist medication (Albuterol, Combivent, etc.), administer the patient’s MDI in lieu of a nebulizer treatment.
  - Consider doses of 4-6 puffs repeated every 5 minutes as needed
  - Use a spacer device if available.
- Limit the use of nebulized medications.
  - Use as indicated for patients in significant respiratory distress secondary to bronchoconstriction who do not respond to initial oxygen administration.
  - Use nebulizers via facepiece masks instead of T-pieces.
  - Perform required nebulizer treatments in well-ventilated areas (outside if appropriate).
- Limit the use of CPAP to patients in significant distress or to those who do not respond to initial oxygen administration methods.
  - Use CPAP only in well-ventilated areas.
- Consider skipping CPAP and proceeding directly to intubation for patients in extremis.
- For patients with impending respiratory failure, consider epinephrine (1 mg/mL) 0.3 mg (0.3 mL) IM. Consider reducing epinephrine dose to 0.15 mg (0.15 mL) IM for patients greater than 50 years or age or those with known history of coronary disease.

Advanced Airway Management / Cardiac Arrest Considerations:
- Use a viral/bacterial filter between the patient and the ventilation device.
  - Insert between with face mask and BVM if patient has not had an advanced airway placed.
o Insert between endotracheal tube/supraglottic airway and BVM in cases where an advanced airway has been placed.

- Video laryngoscopy (VL) may allow providers to maintain a safer distance away from the patient’s airway during intubation.
- If appropriate, avoid a more invasive procedure such as intubation by using an alternative supraglottic airway.
- Consider the use of passive oxygenation instead of positive pressure ventilation for the initial management of the cardiac arrest patient.
  o While performing uninterrupted chest compressions, apply a non-rebreather mask to the patient. Provide oxygen at 15 lpm.
  o Consider withholding BVM ventilations until a definitive airway has been established.

Transport Considerations:
- If patient condition permits, limit EMS provider presence in the rear passenger compartment.
- Separate cab from patient care compartment. This may be accomplished by closing an access door between the two compartments. If vehicle design does not allow for this separation, provide some type of physical barrier – plastic sheeting.
- If cab is isolated from patient care compartment, driver should doff PPE and perform hand hygiene prior to beginning transport.
- If the cab cannot be isolated from the patient care compartment, the driver should wear an appropriate mask during transport. Consider opening up driver compartment windows to provide airflow.
- Engage ventilation system in rear patient care compartment. If vehicle design permits, open vent windows to provide adequate air flow and ventilation.
- Limit or prohibit family members from accompanying patient in ambulance. Note: transport of special populations (i.e., parents with children) may be necessary.

Patient Delivery:
- Notify receiving facility early with suspected EID cases.
- When medically appropriate and available, patients should be wearing a face mask on delivery to a medical facility.
- Receiving facility may direct EMS to deliver patient to an alternate site (i.e., triage tent set up at ED door).
- Upon arrival, if possible, discontinue aerosol-generating procedure prior to entering ED.
  o If patient’s clinical presentation prevents this, the driver or another provider should enter the ED, contact staff, and receive direction on how to best conduct patient transfer of care.
- Limit number of EMS personnel in close proximity to patient while providing appropriate care and transfer.

Return to Service:
- EMS providers should don appropriate PPE before decontaminating the ambulance.
- Leave all doors of the ambulance open to allow for adequate ventilation.
- Decontaminate all surfaces in the ambulance following standard decontamination procedures.
- Decontaminate all equipment (i.e., cardiac monitors, pulse oximeters, oxygen administration equipment, gear bags, and stretchers) before replacing in ambulance.
- After delivery of patient and vehicle/equipment decontamination, EMS providers should enter a clean area and doff PPE following agency and facility policies.
- Doff all PPE in a manner that limits or prevents secondary contamination.
- Following removal of PPE, EMS provider must conduct hand hygiene.
Note:

- Remember, as the EID incident progresses, in addition to seeing patients suffering from the effects of the infectious disease, providers will also be treating their “standard” patient volume.
- For patients not suspected as being infected, standard care following the appropriate patient care standing order is indicated.
- Providers must always exercise a common-sense approach to patient treatment. Provide a level of care that is appropriate to the patient’s condition as well as fitting the operational situation that presents. For example, not every patient requires applying a cardiac monitor and risking a secondary exposure through such a contaminated device.
Appendix A

Guidance for EMS when transporting a patient to a hospital on Divert Status

The following “recommended script” is approved for use by EMS providers to be used for patients requesting transport to a specific destination despite divert status:

Sir/Madam,

The E.R. you are asking to be taken to is currently on Divert status which means they are extremely busy caring for the critically ill or managing very high numbers of patients.

At this time, the ER is asking ambulances to go to other emergency departments in the area.

We would be happy to transport you to another hospital/ ER which can address your needs at this time. We can transport you to the ER you requested even though they are on Divert status but please be aware that you may find longer wait times and more crowded conditions than usual.

Where would you like to go?
ANAPHYLACTIC PRECAUTIONS

Anaphylaxis:
A generalized reaction occurring with dramatic suddenness (usually within a few minutes) after a patient has been exposed to some foreign material.

Cause:
Any drug has the potential to precipitate anaphylaxis. Generally, those administered intravenously or parenterally are more likely to result in life-threatening or fatal anaphylaxis than those ingested or applied to the skin or mucous membranes.

Clinical features:
The patient with anaphylaxis may develop laryngeal edema and bronchospasm which cause respiratory distress and anoxia. The sooner the symptoms develop after the initiating stimulus the more intense the reaction. The symptoms include the following: generalized flush, urticaria, pruritus, anxiety, dyspnea, wheezing, choking, orthopnea, vomiting, cyanosis, paresthesia, shock, and loss of consciousness. Anoxia, shock, and death may occur within 5-10 minutes.

Prevention:
Know the patient's allergy history by asking the patient or family before giving a new medication. Know the precautions listed for each drug.

Treatment:
See Allergic Reaction protocol.

INTRAVENOUS INFILTRATION PRECAUTIONS

Before transporting any patient with an intravenous (IV) access catheter with a solution running, the EMT must check the IV site for patency and signs of infiltration and/or phlebitis. If infiltration occurs, stop the IV fluid do not remove the IV device. Contact the Medical Control physician immediately for orders.

Factors that increase the risk of infiltration:

- Sclerotic vascular disease
- Venous obstruction in the arm (check for edema)
- Radiation treatment near the site of injection
- High drug concentration
- Limited choice for vein selection
- Multiple venipunctures
- Elderly or debilitated
• Superior vena cava syndrome
• Specific characteristics of the drug
• Uncooperative/irrational individual

**Symptoms of an infiltration:**

If pain, burning or stinging occurs at the injection site, evaluate the site for swelling, redness, and inflammation. The presence of a blood return or absence of edema does not negate the possibility of the infusate being spread outside the vein to surrounding tissue. Drug leakage may occur at the site of a previous vessel injury while the needle/catheter is still in the vein.
ACETAMINOPHEN (Liquid)

Pharmacology:
- Analgesic
- Antipyretic

Pharmacokinetics:
- Has little effect on platelet function, no effect of homeostasis, and is not known to produce gastrointestinal bleeding.
- Is not an NSAID – it has no anti-inflammatory properties
- Absorption is rapid
  - Onset of action (PO): < 1 hour
  - Peak effect: 10-60 minutes
  - Duration of action: 4-6 hours
- Metabolized in the liver

Indications:
- Pain of musculoskeletal injury origin

Contraindications:
- Allergy
- Liver disease or injury
- Reduced hepatic function
- Heavy alcohol abuse

Precautions:
- Pregnancy.
- Thrombocytopenia

Side Effects:
- Nausea / vomiting
- Abdominal pain
ALBUTEROL SULFATE/IPRATROPIUM BROMIDE (COMBIVENT)

Pharmacology:
- Combination of ipratropium (an anticholinergic bronchodilator) and Albuterol (a beta-2 adrenergic bronchodilator)
- Ipratropium antagonizes the actions of the neurotransmitter acetylcholine, especially at the muscarinic receptor sites in bronchial smooth muscle
- Albuterol stimulates beta-2 adrenergic receptors of the bronchioles

Pharmacokinetics:
- Bronchodilation
- Onset of action approximately 15 minutes
- Peak effect attained within 1 hour
- Duration of action 4-5 hours

Indications:
- To reverse bronchospasm (wheezing)

Adverse Effects:
- Tachycardia, palpitations, peripheral vasodilation, tremor, headache and nervousness may be seen infrequently

Precautions:
- Paradoxical bronchospasm often with firsts use of new canister
- Use with caution in patients with cardiovascular disease

Contraindications:
- Known hypersensitivity

Dosage:
- Adults: 2 puffs by metered dose inhaler
ALBUTEROL SULFATE PROVENTIL, VENTOLIN, AEROLIN, VENTORUN, ASTHAUN, ProAIR)

Pharmacology:
- Synthetic sympathomimetic amine (a type of stimulant)
- Stimulates beta-2 adrenergic receptors of the bronchioles
- Little effect on blood pressure
- Little cardiac effects
- Main effect is bronchodilation
- It may cause some vasodilation as evidenced by headache or flushing

Pharmacokinetics:
- Bronchodilation begins within 5 to 15 minutes after inhalation
- Peak effect occurs in 30 minutes to 2 hours
- Duration of action is usually 3-4 hours

Indications:
- To reverse bronchospasm (wheezing)

Adverse Effects:
- Tachycardia, palpitations, peripheral vasodilation, cough, headache, dizziness, tremor, and nervousness may be seen infrequently

Precautions:
- Bronchospasm may worsen in rare situations due to patient tolerance or hypersensitivity
- If respirations worsen, discontinue use.
- Should be used with caution in patients with hyperthyroidism or coronary artery disease

Contraindications:
- Known hypersensitivity

Dosage:
- **Adults:** 2.5-5.0 mg by nebulized aerosol or 2 puffs by metered dose inhaler.
- **Children:** Age 1-5 2.5 mg, Age 6 and higher 5.0 mg by nebulized aerosol or 2 puffs by metered dose inhaler.
ASPIRIN

Pharmacology:
- Inhibits platelet aggregation and prostaglandin synthesis

Pharmacokinetics:
- Inhibits platelet aggregation by irreversibly inhibiting prostaglandin cycle-oxygenase for the life of the platelet
- This prevents the formation of the platelet aggregating factor thromboxane A2.
- Onset of action is 1-2 hours
- Duration of action is 6 hours

Indications:
- Acute coronary syndrome- acute myocardial infarction, angina pectoris

Adverse Effects:
- Adverse reactions may include anaphylaxis, bronchospasm, dysrhythmias, hypotension, tachycardia, agitation, cerebral edema, intracranial hemorrhage, dehydration, hyperkalemia and renal failure.

Precautions:
- By inhibiting platelet function, aspirin may lead to an increase in bleeding for patients with bleeding disorders

Contraindications:
- Known allergy to aspirin or non-steroidal, anti-inflammatory drugs (NSAIDS) (i.e. Motrin, Aleve, Ibuprofen, etc.)
- Active GI ulcerations or bleeding, hemophilia or other bleeding disorders
- Pregnancy
- Children under 15 years of age

Dosage:
- Up to 325 mg uncoated PO even if the patient is pain-free

How supplied:
- Aspirin: chewable - 81 mg / tablet or Aspirin - 325 mg / tablet
DIPHENHYDRAMINE HYDROCHLORIDE (BENADRYL®)

Pharmacology:
- An antihistamine
- Blocks H1 and H2 receptor sites

Pharmacokinetics:
- Onset of action occurs within 15-60 minutes
- Peak effect at 2-4 hours
- Duration of action is 6-12 hours

Indications:
- Moderate to severe allergic reactions

Adverse Effects:
- Drowsiness, thickening of bronchial secretions, hypotension, tachycardia, bradycardia, and dry mouth

Precautions:
- Used with caution in patients with severe vomiting, asthma, and alcohol intoxication
- MAO inhibitors may prolong and potentiate diphenhydramine

Contraindications:
- Known hypersensitivity, CNS depression

Dosage:
- Adults: 25-50 mg PO
- Children: Contact Medical Control for pediatric dosage
DUODOTE

Pharmacology:

- Duo Dote is an auto injector that provides a single intramuscular dose of atropine and pralidoxime chloride. It is used as a self-administered therapy for symptomatic exposure to anticholinergic nerve agents and organophosphate pesticides.

Pharmacokinetics Atropine:

- Competitively blocks the effects of acetylcholine at muscarinic receptors on smooth muscle, cardiac muscle and secretory gland cells

Pharmacokinetics Pralidoxime:

- Reactivates acetylcholinesterase which has been inactivated by phosphorylation due to some organophosphorus nerve agents or pesticides.
- Does not reactivate phosphorylated acetylcholinesterase that has undergone the aging process

Indications:

- Poisoning by organophosphorus nerve agents and pesticides

Adverse Effects:

- Temporary headache caused by pralidoxime

Precautions:

- Pralidoxime is excreted in the urine – impaired renal function may result in higher blood levels

Contraindications:

- None in the presence of life-threatening organophosphorus poisoning

Dosage:

- Moderate symptoms: Administer 1 Duo Dote IM
- Severe symptoms: Administer 3 Duo Dotes IM

How supplied:

Auto-injector containing 2.1 mg. of Atropine Sulfate and 600 mg. of Pralidoxime Chloride
**EPINEPHRINE**

**Pharmacology:**
The administration of epinephrine causes increases in:

- Systemic vascular resistance
- Systemic arterial pressure
- Heart rate
- Contractile state
- Myocardial oxygen requirement
- Cardiac automaticity

**Pharmacokinetics:**

- Is rapidly inactivated by the liver.
- Intramuscular administration of epinephrine results in slower absorption due to local vasoconstriction.
- Local massage will hasten absorption

**Indications:**

- Epinephrine selectively improves regional blood flow to the heart and brain
- Patients suffering from severe allergic reactions may be given intramuscular epinephrine

**Precautions:**

- Epinephrine causes a dramatic increase in myocardial oxygen consumption
- Its use in the setting of an acute MI should be restricted to cardiac arrest

**Side Effects:**

- The individual may complain of increased heart rate, pale skin (pallor), dizziness, chest pain, headache, nausea, vomiting, excitability and anxiousness after administration of epinephrine

**Dosage:**

Anaphylactic shock:

- **Adults:** 0.5 mg Intramuscular (EpiPen 0.3 mg), or 0.5 mg draw dose

- **Children:** 0.01 mg/kg Intramuscular (EpiPen Jr)
  - Greater than 30 kg: 0.3 mg IM (EpiPen)
  - Less than 30 kg: 0.15 mg IM (EpiPen Jr)
GLUCOSE, ORAL (Insta-Glucose, Glutose)

Pharmacology:
- Carbohydrate gel

Pharmacokinetics Atropine:
- Provides source of carbohydrate for cellular metabolism

Indications:
- Altered mental status with a history of medication-controlled diabetes
- Hypoglycemia

Adverse Effects:
- Transient increase in blood glucose level

Precautions:
- Patient must be able to maintain the patency of their own airway and effectively swallow the medication

Contraindications:
- Unresponsive patient
- Inability to swallow

Dosage:
- 1 tube = 15 - 24 grams of glucose

How supplied:
- Carbohydrate gel
- Tube contains 15-24 grams of glucose (note; check tube labeling for exact amount, may vary slightly between manufacturers)
HEMOSTATIC AGENT

Indications:

- An agent used to reduce bleeding from minute vessels by hastening the clotting of blood or by the formation of an artificial clot.

Usage:

- Life-threatening bleeding may need to apply tourniquets and pressure points to slow such bleeding enough to apply the agent. Once you slow the high-pressure blood loss, you must still get the agent into contact or close proximity to the source of the bleeding. Some hemostatic agents, such as bandages and sponges, may prove difficult to insert deeply enough to contact the affected artery or organ. Once applied, most hemostatic agents require you to maintain direct pressure on the wound for 2-5 minutes, giving the agent the opportunity to work.

Adverse Effects:

- CELOX: Individuals allergic to shellfish might risk an allergic reaction to chitosan based ChitoGauze.
- QuikClot: If incorrectly applied, the zeolite can quickly reach an extremely high temperature, causing burns and tissue damage.

NOTE: No powder, only agents.
LEVALBUTEROL (XOPENEX, LEVOLIN)

Pharmacology:
- Beta adrenergic receptor agonist
- Stimulates beta-2 adrenergic receptors of the bronchioles

Pharmacokinetics:
- Bronchodilation begins within 5 to 15 minutes after inhalation
- Peak effect occurs in 30 minutes to 2 hours
- Duration of action is usually 3-5 hours

Indications:
- To reverse bronchospasm (wheezing)

Adverse Effects:
- Tachycardia, palpitations, peripheral vasodilation, tremor, headache, dizziness, and nervousness may be seen infrequently

Precautions:
- Bronchospasm may worsen in rare situations due to patient tolerance or hypersensitivity
- If respirations worsen, discontinue use.

Contraindications:
- Known hypersensitivity

Dosage
- Adults: 0.63 = 1.25 mg by nebulized aerosol
- Children: Age 6-11 years: 0.31 to 0.63 mg by nebulized aerosol
ONDANSETRON (ZOFRAN®)

Pharmacology:

- An antiemetic that helps to prevent nausea and vomiting by blocking 5-HT3 receptors so that serotonin is not able to bind to the receptor site and initiate a vomiting reflex

Pharmacokinetics:

- Onset of action occurs within 5 minutes for IV and 30 minutes for ODT administration
- Peak effect occurs around 2 hours with a duration of 3-6 hours
- Duration of action is 2-7 hours

Indications:

- Nausea and vomiting

Adverse Effects:

- Diarrhea, headache, and fever
- Rarely seen are angina chest pain, seizures, feeling of inner restlessness, inability to stay still, and acute involuntary muscle contractions

Precautions:

- Hypersensitivity reactions have been reported in patients who have exhibited hypersensitivity to other selective 5-HT3 receptor antagonists

Contraindications:

- Known hypersensitivity to Ondansetron

Dosage:

- Adults: 8 mg ODT
- Children
  - Give 2 mg ODT if the patient is between 8-15 kg
  - 4 mg if the patient is greater than 15 kg
NALOXONE (NARCAN®)

Description:
- Naloxone is an opioid antagonist.

Pharmacology:
- Naloxone is a competitive narcotic antagonist, which reverses all effects of opioids (i.e. morphine), such as respiratory depression and central and peripheral nervous system effects.

Indications:
- Naloxone is indicated to reverse respiratory and central nervous system depression induced by opioids.

Onset/Duration:
- The onset of action is within a few minutes following an intravenous dose, whereas intramuscular and endotracheal/intranasal administration results in a slower onset of action. The duration of action is approximately 30-60 minutes.

Contraindications:
- Naloxone is contraindicated in hypersensitivity.

Warnings:
- Naloxone may induce opiate withdrawal in patients who are physically dependent. Certain drugs such as propoxyphene (Darvon) may require much higher doses of naloxone for reversal than we currently carry.

Drug Interactions:
- Naloxone is incompatible with bisulfate and alkaline solutions.

Adverse Reactions:
- Adverse reactions may include tachycardia, hypotension, dysrhythmias, nausea, vomiting, and diaphoresis.

Dosage and Routes of Administration:
- 2 mg intranasal (IN). Repeat 2mg IN in opposite nare if no respiratory status change after 4 minutes.
- For pediatric patients, **Contact Medical Control** before administration for guidance on dose.

Storage:
NITROGLYCERINE

Pharmacology:

- Vasodilator-effect on veins more than arteries

Pharmacokinetics Nitro paste:

- Absorbed through the skin
- For antianginal effects the onset is 30 minutes, while duration is 3 hours
- For vasodilation the onset is within 1 hour and duration is 3 to 6 hours.
- Half-life is 1-4 minutes.

Pharmacokinetics Nitro tabs and Nitro Spray:

- Absorbed through oral mucosa
- Antianginal and vasodilation effects within minutes
- Duration of action is less than 5 minutes

Indications:

- For treatment of angina
- Congestive heart failure
- Not to be used for asymptomatic hypertension

Adverse Effects:

- Dose-related
- Headache, hypotension, and dizziness

Precautions:

- May cause hypotension

Contraindications:

- Known hypersensitivity

Dosage:

- One-half to one inch every 6-8 hours
- 0.4 mg sublingual every 5 minutes
- DO NOT USE IN CHILDREN

How supplied:

- Nitrol ointment 2%
- Tablets 0.4 mg
**OXYGEN**

**Pharmacology:**
- Elevates oxygen tension in the blood
- Increases oxygen content of the blood
- Improves tissue oxygenation

**Pharmacokinetics:**
- Changing the percentage of inspired oxygen will result in blood and tissue equilibration within 5 to 20 minutes.

**Indications:**
- Acute chest pain
- Suspected hypoxemia of any etiology (a correlating pulse ox of greater than 92%)
- Cardiopulmonary arrest
- Trauma

**Precautions:**
- The main precaution is not administering enough oxygen to patients who need it. Never withhold oxygen from those in obvious need.
- Oxygen should be given with caution to patients with emphysema
### Appendix C

#### Adult GCS

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<tr>
<th></th>
<th>Eyes</th>
<th>Verbal</th>
<th>Motor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>No Response</td>
<td>No Response</td>
<td>No Response</td>
</tr>
<tr>
<td>2</td>
<td>To Pain</td>
<td>Incomprehensible</td>
<td>Extension (Decerebrate)</td>
</tr>
<tr>
<td>3</td>
<td>To Verbal</td>
<td>Inappropriate</td>
<td>Flexion (Decorticate)</td>
</tr>
<tr>
<td>4</td>
<td>Spontaneous</td>
<td>Confused</td>
<td>Withdraws to Pain</td>
</tr>
<tr>
<td>5</td>
<td>Oriented</td>
<td></td>
<td>Localizes Pain</td>
</tr>
<tr>
<td>6</td>
<td></td>
<td></td>
<td>Obeys Commands</td>
</tr>
</tbody>
</table>

Total: Minimum Score: 3/15 Maximum Score 15/15

#### Pediatric GCS

<table>
<thead>
<tr>
<th></th>
<th>Eyes</th>
<th>Verbal</th>
<th>Motor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>No Response</td>
<td>No Response</td>
<td>No Response</td>
</tr>
<tr>
<td>2</td>
<td>To Pain</td>
<td>Inconsolable, Agitated</td>
<td>Extension (Decerebrate)</td>
</tr>
<tr>
<td>3</td>
<td>To Speech</td>
<td>Inconsistently Consolable/Moans</td>
<td>Flexion (Decorticate)</td>
</tr>
<tr>
<td>4</td>
<td>Spontaneous</td>
<td>Consolable Cry</td>
<td>Withdraws to Pain</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>Coos, Babbles</td>
<td>Withdraws to touch</td>
</tr>
<tr>
<td>6</td>
<td></td>
<td></td>
<td>Normal spontaneous movement</td>
</tr>
</tbody>
</table>

Total: Minimum Score: 3/15 Maximum Score 15/15

#### Normal Vital Signs for Age of Patient

<table>
<thead>
<tr>
<th>Age</th>
<th>Heart Rate</th>
<th>Respiratory Rate</th>
<th>Systolic Blood Pressure (mmHg)</th>
<th>Diastolic Blood Pressure (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Newborn</td>
<td>100-180</td>
<td>25-66</td>
<td>60-90</td>
<td>20-60</td>
</tr>
<tr>
<td>Infant</td>
<td>90-175</td>
<td>22-64</td>
<td>87-105</td>
<td>53-66</td>
</tr>
<tr>
<td>1-5 years</td>
<td>70-156</td>
<td>17-53</td>
<td>90 + 2 x age in years</td>
<td>2/3 systolic</td>
</tr>
<tr>
<td>5-14 years</td>
<td>47-120</td>
<td>12-30</td>
<td>90 + 2 x age in years</td>
<td>2/3 systolic</td>
</tr>
<tr>
<td>14+ years</td>
<td>60-100</td>
<td>8-24</td>
<td>90-140</td>
<td>60-85</td>
</tr>
</tbody>
</table>

The lower limit (5th percentile) of systolic blood pressure can be estimated with this formula: 70 m Hg + (2 x age in years). A low systolic blood pressure should prompt an immediate evaluation for additional signs of inadequate perfusion, such as diminished mental status, prolonged capillary refill, and tachycardia.
**Burn Scale**

Recommendation is to use the Patient's palm as 1% in adults.
In Pediatrics, utilize palm PLUS fingers to estimate 1%

**Modified Lund-Browder chart for reference**
### Pediatric Assessment Triangle (PAT)

![Pediatric Assessment Triangle (PAT)](image)

### Wong-Baker Faces Pain Rating Scale

![Wong-Baker Faces Pain Rating Scale](image)

### Apgar Scoring System

<table>
<thead>
<tr>
<th>Points</th>
<th>0</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Appearance</strong> (skin color)</td>
<td>Blue or pale all over</td>
<td>Pink body, but blue extremities</td>
<td>Pink body and extremities</td>
</tr>
<tr>
<td><strong>Pulse</strong> (heart rate)</td>
<td>Absent</td>
<td>Less than 100</td>
<td>Greater than 100</td>
</tr>
<tr>
<td><strong>Grimace</strong> (reflex/irritability)</td>
<td>No response to stimulation</td>
<td>Grimace/feeble cry when stimulated</td>
<td>Cry or pull away when stimulated</td>
</tr>
<tr>
<td><strong>Activity</strong> (muscle tone)</td>
<td>None</td>
<td>Some flexion</td>
<td>Flexed arms and legs that resist extension</td>
</tr>
<tr>
<td><strong>Respiration</strong> (breathing)</td>
<td>Absent</td>
<td>Weak, irregular, gasping</td>
<td>Strong, lusty cry</td>
</tr>
</tbody>
</table>
Appendix D

INFECTION CONTROL

INDICATIONS: These guidelines should be used whenever contact with patient body substances is anticipated and/or when cleaning areas or equipment contaminated with blood or other body fluids.

These guidelines provide general information related to body substance isolation and the use of universal precautions. These guidelines are not designed to supersede an Emergency Medical Service's Infection Control Policy; but should be a resource for their creation. It will serve as policy in the absence of a service’s policy.

These guidelines do not comprehensively cover all possible situations, and EMS practitioner judgment should be used when the Emergency Medical Service's Infection Control Policy does not provide specific direction.

Nothing in this guideline shall be construed to authorize the disclosure of confidential medical information by the health facility or any of the EMS practitioners except as otherwise authorized by law.

- Wear gloves on all calls where contact with blood or body fluid is anticipated or when handling items or equipment that may be contaminated with blood or other body fluids.

- Wash hands as often as possible and after every call.

- Keep all open cuts and abrasions covered with adhesive bandages that repel liquids.

- Use goggles or glasses when spraying or splashing of body fluids is possible.

- Respiratory precautions should be used when caring for any patient with a known or suspected infectious disease that is transmitted by respiratory droplets or with someone who has a productive cough.

- A mask should be placed upon the patient if his/her respiratory condition permits.

- If an EMS practitioner has a potential exposure to blood, body fluids, or airborne pathogens; the practitioner must follow Delaware Law Title 16 Chapter 12A and the Emergency Medical Service's Infection Control Policy. The incident must be immediately reported to the service’s Infection Control Officer.

- EMS practitioners should clean their wound with soap and water; flush mucous membranes with water/saline; or treat any other wound as dictated by severity of the wound.

- EMS practitioners who have a confirmed exposure (as confirmed by the service's Infection Control Officer or Receiving Medical Facility) should be evaluated at the receiving facility. If the patient is not transported, contact the infection control officer for guidance on a facility to be evaluated.

- A State of Delaware Infectious Control Form "Report of Potential Exposure" should be
filled out at the receiving hospital or forwarded to the Chief Medical Examiner/Coroner as soon as possible.

- EMS practitioners who have been treated for a confirmed exposure should follow through with post-exposure medical care and/or prescribed treatment.

- Thoroughly clean and disinfect equipment after each use following service guidelines that are consistent with the Center for Disease Control recommendations.

- Place all disposable equipment and contaminated trash in a clearly marked plastic biohazard bag and dispose of appropriately.

- Contaminated uniforms and clothing should be removed, placed in an appropriately marked biohazard bag and laundered/decontaminated.

- All needles and sharps must be disposed of in a sharps receptacle unit and disposed of appropriately.
REPORT OF POTENTIAL EXPOSURE FORM
EMERGENCY MEDICAL CARE PROVIDER

**All fields must be completed**

<table>
<thead>
<tr>
<th>Section A: To be completed by the Emergency Medical Care Provider with assistance from the Agency’s DO (please print)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Submitting Agency:</td>
</tr>
<tr>
<td>Submitting Agency’s Designated Officer (DO) Name:</td>
</tr>
<tr>
<td>Submitting Agency’s Address:</td>
</tr>
<tr>
<td>Emergency Medical Care Provider’s Name:</td>
</tr>
<tr>
<td>Source Patient’s Name:</td>
</tr>
<tr>
<td>Location of Incident:</td>
</tr>
<tr>
<td>Date of Exposure:</td>
</tr>
<tr>
<td>Source Patient Transported To:</td>
</tr>
<tr>
<td>Date Form Submitted:</td>
</tr>
</tbody>
</table>

What was the Exposure Route?
- Inhalation
- Ingestion
- Injection
- Direct Contact
- Confined proximity (duration: )
- Inhalation
- Ingestion
- Injection
- Direct Contact
- Confined proximity (duration: )
- Inhalation
- Ingestion
- Injection
- Direct Contact
- Confined proximity (duration: )
- Body Fluid Exposure: 
- Blood
- Urine
- Feces
- Sweat
- Amniotic Fluid
- Personal Protective Equipment (PPE) Used: 
- No
- Gloves
- Eye Protection
- HEPA Mask (N95 or better)
- Surgical Mask
- Other
- Did PPE fail?  
- Yes  
- No  
- Did you receive medical attention?  
- Yes  
- No  

**Describe the incident and extent of the exposure on the backside of this paper. (Be detailed)**

Emergency Medical Care Provider’s Signature
Agency’s Designated Officer’s Signature

<table>
<thead>
<tr>
<th>Section B: To be completed by the receiving medical facility (please print)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Facility Name:</td>
</tr>
<tr>
<td>Health Care Provider’s Name:</td>
</tr>
<tr>
<td>Phone #:</td>
</tr>
<tr>
<td>Facility’s Designated Officer’s Name:</td>
</tr>
</tbody>
</table>
| Source Patient  
- Source has NO known infectious disease.  
- Source has known infectious disease.  
- Source patient NOT tested  
- Confirmed medical record test  
| Emergency Medical Care Provider:  
- Post exposure prophylaxis indicated?  
- Yes  
- No  
- If YES, treatment given:  
- Follow up necessary  
- The Emergency Medical Care Provider has been informed of the results of the evaluation for exposure to bloodborne, airborne, and/or potentially infectious materials  
- Notification made by:  
- Phone  
- Mail  
- Email  
- Fax  
- Other  
- Caller’s Name:  
- Date:  
- Time (24 hr): |
| Facility Notes: |

The information provided on this form is confidential.
**NARRATIVE:** Describe the incident and extent of exposure (include exposed body part, exposure duration, and decontamination).

<table>
<thead>
<tr>
<th>DETAILED NARRATIVE: Describe the incident and extent of exposure (include exposed body part, exposure duration, and decontamination).</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

**INFECTION CONTROL EXPOSURE ALGORITHM**

All forms are located at [http://dhs.delaware.gov/dhsoph/emergency/forms.htm](http://dhs.delaware.gov/dhsoph/emergency/forms.htm) or email us at OEMS@state.de.us

**EMERGENCY MEDICAL CARE PROVIDER:**

- Report Exposure to Agency's Designated Officer and (if needed) the medical facility as soon as possible (within 24 hours)
- Follow your agency's exposure control plan
- Complete Section (A): Report of Potential Exposure Form
- Follow Agency's Designated Officer and/or Medical Evaluator's Recommendations

**AGENCY'S DESIGNATED OFFICER (DO):**

- Help complete Section (A): Report of Potential Exposure Form
- Consult with medical evaluator for source blood testing; send form with Emergency Medical Care Provider to receiving Medical Facility
- NO KNOWN Exposure: Maintain record keeping
- KNOWN Exposure: Maintain ongoing monitoring of exposed Provider through course of employment.
- Provide copy of Report of Potential Exposure Form to OEMS. Black out provider/source information.
- Submit Designated Officer's Confirmation of Exposures to OEMS

**MEDICAL EVALUATOR:**

- Complete Section (B): Report of Potential Exposure Form
- Counsel and treat prehospital responder as needed
- Make three copies of the completed Report of Potential Exposure Form
- Keep a completed copy of the Report of Potential Exposure Form as a confidential medical record for the hospital
- Send the completed original Report of Potential Exposure Form to the submitting agency for their records
- Notify Agency's Designated Officer of exposure results (and Public Health if required) within 48 hours of confirmed exposure
- Provide copy of Report of Potential Exposure Form to OEMS. Black out provider/source information.

Additional Information may be found in Delaware Law under Title 16 Chapter 12A, Notification of Emergency Medical Providers of Persons with Communicable Disease.
Appendix E

NON-INVASIVE GAS MONITORING
PULSE OXIMETRY and CO-OXIMETRY

INDICATIONS: Pulse oximetry and CO-oximetry is an adjunctive technique that can help to detect hypoxia and to assess the impact of oxygen therapy. The EMT assessment and treatment of the patient is much more important than the pulse oximeter or CO-oximeter reading. The pulse oximeter and CO-oximeter supplies one additional small piece of information.

"Room" Carbon monoxide monitoring is an adjunctive technique that can help to detect the presence of carbon monoxide in the air that would pose a threat to the patient and EMS crews.

- "Room" Carbon monoxide monitoring: *
  - If audible alert sounds suspect the presence of carbon monoxide.
  - Consider scene safety and additional resources
  - As soon as practical, remove patient to non-contaminated area

- Pulse oximetry and CO-oximetry:
  - Provide appropriate supplemental oxygen.
  - Obtain a pulse oximeter reading (SpO2). **
    - The pulse oximeter reading can be assessed prior to giving oxygen if this does not significantly delay oxygen therapy. A reading taken after oxygen has been administered can be compared to the first reading for signs of improvement or deterioration of oxygenation.
    - Make sure the pulse-ox reading correlates with the patient's palpated pulse rate
  - Obtain CO-oximeter reading, if available ***
    - If carboxyhemoglobin is less than 5%, consider other possible causes of symptoms.
    - If carboxyhemoglobin is greater than 5%, and patient has suffered a loss of consciousness or altered mental status, suspect CO poisoning

- Continue oxygen therapy utilizing non-rebreather mask or BVM and transport
  - Regardless of pulse-oximetry reading

- Always treat the patient, not the pulse oximeter or CO-oximeter reading. Do not let the pulse oximeter or CO-oximeter delay other assessment or treatment.

*Certain medical conditions will give a falsely high pulse oximeter reading. The most common condition is carbon monoxide poisoning. Do not rely on a pulse oximeter reading if carbon monoxide toxicity is a consideration.

**CO oximetry may be performed as an option by agencies carrying CO monitoring equipment
Appendix F

For updated Hospital Contacts, click link below:
https://www.dhss.delaware.gov/dph/ems/ems.html

Hospital Contacts

Christiana E.D. 302-733-1638 or -1621
Wilmington E.D. 302-428-4182 or -4886
Saint Francis E.D. 302-421-4333
A.I. DuPont E.D. 302-651-4183
Crozier E.D. 610-447-2186 or -2188
Kent General E.D. 302-744-7122
Milford Memorial E.D. 302-430-5720 or -5721
Nanticoke E.D. 302-629-6611 x2555
Beebe E.D. 302-645-3554
P.R.M.C. E.D. 410-543-7100
South Coastal E. D. 302-291-6900

Christiana Medic Room 302-733-1617
Kent General Medic Room 302-744-6953
Milford Medic Room 302-430-5654

Adult Protective Services 800-223-9074
Child Abuse Reporting Hotline 800-292-9582
Domestic Violence 302-678-3886
Division of Long-Term Care hotline 877-453-0012
Delaware Victim Services 800-842-8461

DE Office of EMS 302-223-1350
Poison Control 800-222-1222
NCC Fireboard 302-571-7331
Kent Center 302-734-6040
Sussex Fireboard 302-855-2970
Appendix G

Suspected Stroke Assessment Tool
VA N Stroke Assessment

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arm and/or leg drift, unilateral weakness, or paralysis – <strong>NO</strong> weakness noted</td>
<td><strong>VAN Negative</strong></td>
</tr>
<tr>
<td>Arm and/or leg drift, unilateral weakness, or paralysis – <strong>Weakness noted</strong></td>
<td>Continue with <strong>VAN</strong> assessment</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>V</th>
<th>Visual Disturbance</th>
<th>Double-vision, visual field cut, or new loss of vision?</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>A</th>
<th>Aphasia</th>
<th>Difficulty forming words, or understanding/following commands? Difficulty recognizing objects correctly?</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>N</th>
<th>Neglect</th>
<th>Present with a gaze deviation or inability to cross midline? Unable to feel both sides at the same time when touched? Unable to recognize their own arm, or ignoring one side?</th>
</tr>
</thead>
</table>

| Arm and/or leg weakness **AND** visual/aphasia/neglect symptoms | **VAN Positive** |
| Arm and/or leg weakness and **NO** visual/aphasia/neglect symptoms | **VAN Negative** |

Transport to the nearest appropriate State of Delaware or Joint Commission certified Stroke Center without delays and request prehospital stroke alert for the following categories:

- VAN negative and LKW less than 4.5 hours → go to nearest certified Stroke Center.
- VAN positive and LKW less than 4.5 hours → contact local medical control to discuss destination.
- VAN positive and LKW greater than 4.5 hours (including wake up stroke or unknown LKW) or considering hemorrhagic stroke → consider transport to certified Thrombectomy Capable or Comprehensive Stroke Center.
**DELWARE MEDICAL ORDERS FOR SCOPE OF TREATMENT (DMOST)**

- **FIRST,** follow the orders below. **THEN** contact physician or other health care practitioner for further orders, if indicated.
- The DMOST form is voluntary and is to be used by a patient with serious illness or frailty whose health care practitioner would not be surprised if the patient died within next year.
- Any section not completed requires providing the patient with the full treatment described in that section.
- Always provide comfort measures, regardless of the level of treatment chosen.
- The Patient or the Authorized Representative has been given a plain-language explanation of the DMOST form.
- The DMOST form must accompany the patient at all times. It is valid in every health care setting in Delaware.

<table>
<thead>
<tr>
<th>Print Patient's Name (last, first, middle)</th>
<th>Date of Birth last four digits of SSN</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Goals of Care</strong></td>
<td>(see reverse for instructions. This section does not constitute a medical order.)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>B</strong></th>
<th><strong>C</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient has no pulse and/or is not breathing</td>
<td>Medical Interventions: Patient is breathing and/or has a pulse.</td>
</tr>
<tr>
<td>□ Attempt resuscitation/CPR.</td>
<td>□ Full Treatment: Use all appropriate medical and surgical interventions, including intubation and mechanical ventilation in an intensive care setting, if indicated to support life. Transfer to a hospital, if necessary.</td>
</tr>
<tr>
<td>□ Do not attempt resuscitation/DNAR.</td>
<td>□ Limited Treatment: Use appropriate medical treatment, such as antibiotics and IV fluids, as indicated. May use oxygen and noninvasive positive airway pressure. Generally avoid invasive care.</td>
</tr>
<tr>
<td></td>
<td>□ Transfer to hospital for medical interventions.</td>
</tr>
<tr>
<td></td>
<td>□ Transfer to hospital only if comfort needs cannot be met in current setting.</td>
</tr>
<tr>
<td></td>
<td>□ Treatment of Symptoms Only/Comfort Measures: Use any medications, including pain medication, by any route, positioning, wound care, and other measures to keep clean, warm, dry, and comfortable. Use oxygen, suctioning, and manual treatment of airway obstruction as needed for comfort.</td>
</tr>
<tr>
<td></td>
<td>Use antibiotics only to promote comfort. Transfer only if comfort needs cannot be met in current location.</td>
</tr>
<tr>
<td></td>
<td>□ Other Orders:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>D</strong></th>
<th><strong>E</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Artifically Administered Fluids and Nutrition: Always offer fluids by mouth if feasible and desired.</td>
<td>Print Name of Authorized Representative Relation to Patient Address Phone #</td>
</tr>
<tr>
<td>□ Long-term artificial nutrition</td>
<td>If I lose capacity, my Authorized Representative may not change or void this DMOST</td>
</tr>
<tr>
<td>□ Defined trial period of artificial nutrition: Length of trial: ___________ Goal: ___________</td>
<td>Patient Signature</td>
</tr>
<tr>
<td>□ No artificial nutrition □ hydration only □ none (check one box)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Orders Discussed With:</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>□ Patient ph.#</td>
<td>□ Guardian Surrogate (per DE Surrogacy Statute) Printed Name &amp; phone number</td>
</tr>
<tr>
<td>□ Other Agent under healthcare POA/AHCD</td>
<td></td>
</tr>
<tr>
<td>□ Parent of a minor</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>F</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>SIGNATURES: Patient/Authorized Representative/Parent (mandatory) I have discussed this information with my Physician/APRN/PA</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Signature</th>
<th>Date</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Print Name</td>
<td>Print Address</td>
<td>License Number</td>
</tr>
</tbody>
</table>
DIRECTIONS FOR HEALTH CARE PROFESSIONALS

COMPLETING A DMOST FORM
- Must be signed by a Licensed Physician, Advance Practice Registered Nurse, or Physician Assistant.
- Use of original form is highly encouraged. Photocopies and faxes of signed DMOST forms are legal and valid.
- Any incomplete section of a DMOST form indicates the patient should get the full treatment described in that section.

REVIEWING A DMOST FORM – It is recommended that a DMOST form be reviewed periodically, especially when:
- The patient is transferred from one care setting or care level to another,
- There is a substantial change in the patient’s health status, or
- The patient’s treatment preferences change.

MODIFYING AND VOIDING INFORMATION ON A COMPLETED DMOST FORM
A patient with decision-making capacity can void a DMOST form at any time in any manner that indicates an intent to void. Any modification to the form voids the DMOST form. A new DMOST form may be completed with a health care practitioner.
Forms are available online at www.delaware.gov.

SECTION A This section outlines the specific goals that the patient is trying to achieve by this treatment plan. Health care professionals shall share information regarding prognosis with the patient in order to assist the patient in setting achievable goals. Examples may include:
- Longevity, cure, remission or better quality of life
- To live long enough to attend an important event (wedding, birthday, graduation)
- To live without pain, nausea, shortness of breath or other symptoms
- Eating, driving, gardening, enjoying time with family, or other activities

SECTION B This is a medical order. Mark a selection for the patient’s preferences regarding CPR.

SECTION C This is a medical order. When “limited treatment” is selected, also indicate whether the patient prefers or does not prefer transfer to a hospital for additional care:
- IV medication to enhance comfort may be appropriate treatment for a patient who has indicated “symptom treatment only.”
- Non-invasive positive airway pressure includes continuous positive airway pressure (CPAP) and bi-level positive airway pressure (Bi-PAP).
- The patient will always be provided with comfort measures.
- Patients who are already receiving long-term mechanical ventilation may indicate treatment limitations on the “Other Orders” line.

SECTION D This is medical order. Mark a selection for the patient’s preferences regarding nutrition and hydration. Check one box:
- Oral fluids and nutrition should always be offered if feasible and consistent with the goals of care.

SECTION E This section documents with whom the medical orders were discussed, the name of any health care professional who assisted in the completion of the Form, the name of any authorized representative and if the authorized representative may not modify/void the form.

SECTION F To be valid, all information in this section must be completed.

HIPAA PERMITS DISCLOSURE OF DMOST TO OTHER HEALTH CARE PROFESSIONALS AS NECESSARY FOR TREATMENT.

SEND FORM WITH PATIENT WHENEVER MOVED TO A NEW SETTING
- Faxed, Copied, or Electronic Versions of the Form are legal and valid.
Delaware Medical Orders for Scope of Treatment (DMOST)

DMOST is a process for documenting treatment choices. The DMOST form is voluntary. It is a portable, standardized Medical Order that will be recognized and followed by Delaware health care providers.

The DMOST conversation is an opportunity to understand the likely course of your health and medical condition, so that you may make informed choices that are appropriate and reflect what you want. If you choose, you may invite loved ones to join this conversation.

Q. What is DMOST?
A. The Delaware Medical Orders for Scope of Treatment (DMOST) form is a portable medical order form. It allows you to make choices about life-sustaining treatments, including among other treatments, CPR (resuscitation) and artificial nutrition. You may request full treatment, limited treatment, or comfort care only.

Q. Who is it for?
A. A DMOST form can be used by a person with a serious illness or frailty, whose health-care practitioner would not be surprised if they died within the next year.

Q. When should it be discussed and signed? Who signs it?
A. A DMOST form is completed after a conversation you have with a health care practitioner. It is signed by you and a physician (MD or DO), an advanced practice registered nurse (APRN), or a physician assistant (PA). The physician/APRN/PA signature makes the choices into portable medical orders.

Q. Who is required to follow the wishes documented on the DMOST form?
A. These orders will be followed by health care providers in any setting (ambulance, long-term care facility, emergency room, hospital, hospice, home, assisted living facility, etc.). It travels with you and is honored when you move to a new setting.

Q. Can someone else make DMOST decisions for me?
A. You make health-care decisions for yourself as long as you have decision-making capacity. You have the right to change your authorized representative at any time while you have decision-making capacity.

If a physician determines that a person lacks decision-making capacity, an authorized representative can sign a DMOST form on behalf of that person. DMOST form does not change the decision-maker designated by an Advance Health Care Directive, a Health Care Power of Attorney document, a guardian of person appointed by a Court, or Delaware law on health care surrogates.

If you have capacity and complete a DMOST form, you can sign on the form saying that if you lose capacity, your authorized representative cannot void the form you signed.

Q. What if I change my mind?
A. If your condition or your choices change, you or your authorized representative should void (cancel) your DMOST form and request a new DMOST be completed with your new choices. You cannot void a DMOST form if you change your mind but do not want to create a new one. You may not make any changes to the content of the DMOST form. If you want to change your DMOST form you must void your previous form and complete a new one with your health-care practitioner. If your DMOST form does not agree with your advance directive, the most recent document will be followed.

Q. Must I do this?
A. The DMOST form is always voluntary and can be voided at any time. A Health care organization is prohibited from requiring you to complete a DMOST form for any reason, including as part of a person’s admission to a health care facility.

It is important to understand that this form contains medical orders. It will be followed by health care providers. For example, if you choose “Do Not Attempt Resuscitation”, and your heart stops, no attempt will be made to restart your heart. If you choose “Intubate/Use Artificial Ventilation”, then you may be placed on a breathing machine with a tube in your throat and transferred to an intensive care setting in a hospital.

Q. What will happen to my choices if I travel out of state?
A. Many states, including all the states in our region, currently use a form similar to the DMOST form. Forms from those states which are valid under the Delaware Law will be honored in Delaware. DMOST forms will be honored in other states which have reciprocity.
## Appendix I

# Mass Casualty Triage Charts

### JumpSTART Mass Casualty Triage (pediatric)

<table>
<thead>
<tr>
<th>Move the walking wounded</th>
<th>➔</th>
<th>MINOR</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RESPIRATIONS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No respirations</td>
<td>➔</td>
<td>DEAD</td>
</tr>
<tr>
<td>No peripheral pulse</td>
<td>➔</td>
<td>DEAD</td>
</tr>
<tr>
<td>No respirations</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td>With peripheral pulse</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td>Give 5 breaths with barrier</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiration resume</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td>No spontaneous respirations</td>
<td>➔</td>
<td>DEAD</td>
</tr>
<tr>
<td>Respirations labored</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td>Respirations &gt; 45 or &lt;15 per minute</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td><strong>PERFUSION</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No peripheral pulse</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td>Cap refill &gt;2 sec</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
<tr>
<td><strong>MENTAL STATUS (AVPU)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AV</td>
<td>➔</td>
<td>DELAYED</td>
</tr>
<tr>
<td>PU</td>
<td>➔</td>
<td>IMMEDIATE</td>
</tr>
</tbody>
</table>
Appendix J

Best Practices for Pediatric Transportation

All children should be properly restrained when riding in a vehicle. An ill or injured child must be restrained in a manner that minimizes injury in an ambulance crash. The best location for transporting a pediatric patient is on the ambulance cot. The method of restraint should be determined by various circumstances including the child’s medical condition and weight.

1. Convertible car seat with two belt paths (front and back) with four points for belt attachment to the cot is considered best practice for pediatric patients who can tolerate a semi-upright position.
   - Position safety seat on cot facing foot-end with backrest fully elevated.
   - Secure safety seat with 2 pairs of belts at both forward and rear points of seat.
   - Place shoulder straps of the harness through slots just below child’s shoulders and fasten snugly to child.
   - Follow manufacturer’s guidelines regarding child’s weight.
   **Note:** Non-convertible safety seats cannot be secured safely to cot. If child’s personal safety seat is not a convertible seat, it cannot be used on the cot.

2. Car bed with both a front and rear belt path (example: Cosco Dream Ride SE)
   - For infants who cannot tolerate a semi-upright position or who must lie flat.
   - Position car bed so infant lies perpendicular to cot, keeping infant’s head toward center of patient compartment.
   - Fully raise backrest and anchor car bed to cot with 2 belts, utilizing 4 loop straps supplied with car bed.
   - Only appropriate for infants from 5 – 20 lbs.

3. Restraint device (marketed to EMS) with 5-point harness (examples: Ferno Pedi-Mate, SafeGuard Transport)
   - Attach securely to cot utilizing upper back strap behind cot and lower straps around cot’s frame.
   - 5-point harness must rest snugly against child.
   - Adjust head portion of cot according to manufacturer’s recommendation.
   - Pedi-mate fits children weighing 10 – 40 lbs. SafeGuard Transport fits children weighing 22 – 100 lbs.
4. Child belted directly to backboard and/or cot in manner to prevent ramping or sliding in a front or rear end crash
   - Loop narrow belt under each arm and extend over child’s shoulder securing belt at shoulder level so no gap exists above shoulder.
   - Use soft, sliding, or breakaway connector to hold shoulder straps together on chest.
   - Anchor 2 belts to non-sliding cot member and route over thighs and hips, not around waist.

5. Isolette restraint device with 3-point harness (example: International Biomed papoose)
   - Rest harness securely on child with no blanket or sheet between harness and child.
   - Attach to isolette tray at four points.
   - Additional soft Velcro straps may be added for lateral security.
   - Blanket or towels may be used to provide stabilization of the head.

NON-PATIENT TRANSPORT

Best practice is to transport well children in a vehicle other than the ambulance, whenever possible, for safety.

If no other vehicle is available and circumstances dictate that the ambulance must transport a well child, he/she may be transported in the following locations:
   - Captain’s chair in patient compartment using a size appropriate integrated seat or a convertible safety seat that is secured safely in relationship to the orientation of the captain’s chair.
   - Passenger seat of the driver’s compartment if child is large enough (according to manufacturer’s guidelines) to ride forward-facing in a child safety seat or booster seat. Airbag should be turned off. If the air bag can be deactivated, an infant, restrained in a rear-facing infant seat, may be placed in the passenger seat of the driver’s compartment.

USE OF PATIENT’S CHILD PASSENGER SAFETY SEAT AFTER INVOLVEMENT IN MOTOR VEHICLE CRASH

The patient’s safety seat may be used to transport the child to the hospital after involvement in a minor crash if ALL of the following apply:
   - It is a convertible seat with both front and rear belt paths.
   - Visual inspection, including under movable seat padding, does not reveal cracks or deformation.
   - Vehicle in which safety seat was installed was capable of being driven from the scene of the crash.
   - Vehicle door nearest the child safety seat was undamaged.
   - The air bags (if any) did not deploy.