REGIONAL EMS CONSORTIUM

Marion County Emergency Medical Services

PATIENT TREATMENT PROTOCOLS

2025

These patient care protocols will go into effect 04/01/2025 for EMRs, EMTs AEMTs, Intermediates and Paramedics of Santiam Ambulance, Stayton Fire District, Turner Fire District, Gates Fire District, Lyons Fire District, Mill City Fire District, Aumsville Fire District, Sublimity Fire District, and Idanha/Detroit Fire District.

These protocols, we believe, are the best of their type. Where evidence has been available, the Marion County EMS Protocol Development Committee has diligently evaluated the material and drafted protocols that will assist us in providing excellent patient care. Where evidence is lacking, we have relied on best practices, expert advice and consensus to guide the development of the protocol or procedure. These protocols are reviewed on a regular basis and updated when necessary to reflect advances in the art and science pertaining to the care of the acutely ill and injured.

Remember that these protocols are guidelines. EMS is performed in a stressful environment with time-critical decisions and no specific patient care matrix can be developed that will cover every type of injury, illness, and complicating circumstance that EMT providers will encounter while providing on-scene care. It is our expectation that providers will use these protocols in conjunction with their training and experience to do what is best for each patient. From time to time, it is expected that circumstances will arise that are not covered within these protocols. In such instances, providers should function within their scope of practice and use all available resources (including On-Line Medical Control) to provide the best possible patient care.

Thanks to everyone who has aided in protocol development and review. Anything that is complex and includes detail is prone to errors. Please review these protocols carefully and route any potential errors, unclear directions, or suggestions for improvement to your agency's EMS Office.

Finally, we thank every one of you for your dedication and commitment every day to providing the best possible pre-hospital medical care to the citizens of our respective communities.

Steve Vets DO Medical Director

Date: 03/31/2025

HOSPITALS	ADDRESS		ED PHONE	FAX NUMBER
Salem Hospital	890 Oak St SE Salem OR 97301	503-561-5200	503-814-1300	503-814-1055
Santiam Hospital	1401 N 10th St Stayton OR 97383	503-769-2175	503-769-9256	503-769-4023
Legacy Silverton Hospital	342 Fairview St, Silverton, OR 97381	(503) 873-1500		
Good Sam Albany Hospital	1046 6th Ave SW Albany, OR 97321	541-812-4000		
Good Sam Regional Medical Center Corvalllis	3600 NW Samaritan Drive Corvallis, OR 97330	541-768-5111		
Samaritan Lebanon Community Hospital	525 Santiam Hwy SE, Lebanon, OR 97355	541-258-2101		
West Valley Hospital	525 SE Washington Street Dallas OR 97355	503-623-8301		
Providence Newberg Medical Center	1001 Providence Dr. Newberg. OR 97132	503-537-1555	503-537-1785	
Shriners Hospital for Children -Portland	3101 SW Sam Jackson Park Rd Portland OR 97239	am Jackson Park Rd 503-241-5090		
Legacy Emanuel	2801 N. Gantenbein Ave. Portland OR 97227	503-413-2200	503-413-4121	
Randall Children's Hospital at Legacy Emanuel	2801 N. Gantenbein Ave. Portland OR 97227	503-276-6500		
OHSU ED	3181 SW Sam Jackson Park Rd Portland OR 97239	503-494-4036	503-494-7551	
VA Portland ED	tland ED 3710 SW US Veterans Hospital Rd, Portland, OR 97239		503-721-7803	
Willamette Valley Medical Center	2700 SE Stratus Ave. McMinnville OR 97128	503-472-6131		
Kaiser Sunnyside	10180 Sunnyside Rd Clackamas OR 97015	503-813-2000	503-571-9516	
Legacy Meridian Park	19300 S.W. 65th Ave. Tualatin Oregon 97062	503-692-1212	503-692-7467	
Portland Adventist Medical Center ED	10300 SE Main St Portland OR 97216	503-251-6168	503-257-2500	
Providence St Vincent ED	9205 SW Barnes Rd. Portland OR 97225	503-216-1234	503-216-2444	
Providence Willamette Falls ED	1500 Division St Oregon City 97045	503-656-1631	503-657-6702	
Mckenzie Willamette Medical Center	1460 G Street Springfield OR 97477	541-726-4400		
Sacred Heart University District	1255 Hilyard Street Eugene OR 97401	541-686-7300		
Saint Charles Bend	1253 NE Neff Rd Bend OR 97701	541-382-4321		

Table of Contents

Scope of Practice	00.010
Medical Control of Medications and Procedures	00.040
Treatment – 10.000	
Universal Patient Care	
Abdominal Pain	
Agitated Patient	
Altered Mental Status & Coma	
Anaphylaxis & Allergic Reactions	
Brief Resolved Unexplained Event (BRUE)	
Burns	
Cardiac Arrest – AED / CPR	
Cardiac Arrest – <i>General Guidelines</i>	
Cardiac Arrest – <i>Pulseless Arrest</i>	
Cardiac Arrest – <i>Trauma</i>	
Cardiac Arrest – <i>Pregnancy</i>	
Cardiac Arrest – Post Resuscitation	
Cardiac Dysrhythmias – <i>Bradycardia</i>	
Cardiac Dysrhythmias – <i>Tachycardia - Stable</i>	
Cardiac Dysrhythmias – <i>Tachycardia - Unstable</i>	
Chest Pain / Acute Coronary Syndrome	
Crush Injury / Entrapment	
Diabetic Emergencies-Hyperglycemia	
Diabetic Emergencies-Hypoglycemia	
Epistaxis	
Eye Emergencies	
Fever Management	
Hyperkalemia	
Hyperthermia/Heat Related Emergencies	
Hypothermia	
Musculoskeletal Trauma – Extremity / Hemorrhage	
Musculoskeletal Trauma – Spinal Injury	
Nausea & Vomiting	
Neonatal Resuscitation	

Treatment – 10.000 (Continued)

Obstetrical Childbirth	10.130
Obstetrical Complications	
Pain Management	10.135
Poisoning & Overdose	10.140
Respiratory Distress- CHF/Pulmonary Edema	10.160
Respiratory Distress- COPD/Asthma	
Respiratory Distress- Pediatrics	10.160
Seizures	10.170
Sepsis	10.175
Shock	10.180
Stroke / CVA	
Submerged Patient/Drowning	
Traumatic Brain Injury	

Medications – 20.000

Acetaminophen	
Activated Charcoal	
Adenosine (Adenocard [®])	
Albuterol (Ventolin [®])	
Amiodarone (Cordarone®)	
Aspirin	
Atropine Sulfate	
Calcium Gluconate	
Dexamethasone (Decadron®)	
Dextrose	
Diltiazem	
Diphenhydramine (Benadryl [®])	
Droperidol (Inapsine®)	
Epinephrine	
Etomidate (Amidate [®])	

Medications – 20.000 (Continued)

Fentanyl (Sublimaze [®])	20.117
Glucagon	
Glucose – Oral	20.140
Haloperidol (Haldol [®])	20.142
Hydromorphone (Dilaudid [®])	20.144
Hydroxocobalamin (CYANOKIT®)	20.145
lbuprofen	20.147
Ipratropium Bromide (Atrovent [®])	20.150
Ketamine Hydrochloride	20.155
Ketorolac Tromethamine (Toradol [®])	20.157
Labetalol	20.158
Lidocaine	
Magnesium Sulfate	20.170
Midazolam (Versed®)	
Morphine Sulfate	
Naloxone (Narcan®)	
Nitroglycerin	
Norepinephrine (Levophed®)	
Ondansetron (Zofran [®])	
Oxygen	
Oxymetazoline Hydrochloride (Afrin®)	20.245
Oxytocin (Pitocin [®])	
Pralidoxime (Protopam® / 2-PAM [®])	
Rocuronium (Zemuron [®])	20.257
Sodium Bicarbonate	20.260
Sodium Thiosulfate 25%	20.265
Succinylcholine (Anectine®)	
Tranexamic Acid (TXA)	20.277
Vecuronium (Norcuron®)	

Table of Contents

Procedures – 30.000

	AICD Deactivation	
	Behavioral Health Emergencies	
	Breath Actuated Nebulizer	
	Continuous Positive Airway Pressure (CPAP)	
	Emergency Cricothyrotomy	
	Endotracheal Intubation	
	End Tidal CO ₂ Monitoring	
	i-gel [®] Supraglottic Airway Device	
	Intraosseous Access & Infusion	
	Left Ventricular Assist Devices (LVAD)	
	Orogastric Tube Insertion and Maintenance	
	Patellar Dislocation Reduction	
	PICC Line Access	
	Positive End-Expiratory Pressure	
	Sports Equipment Removal	
	Suctioning	
	TASER [®] Barb Removal	
	Tension Pneumothorax Decompression	
	Tourniquet Placement	
	Transcutaneous Pacing	
Оре	rations – 50.000	
	Ambulance Diversion Guidelines	
	Crime Scene Response	
	Death and Dying	
	Death Notifications	

Table of Contents

Operations – 50.000 (Continued)

Documentation	
Hand-Off Reports	
Hazardous Materials Response	
Hospice and Palliative Care	
Lift Assists	
Multiple Toxic Exposure	
Medical Control of Scene	
On-Line Medical Consult	
Refusal and Informed Consent	
Rehabilitation	
Trauma System – 60.000	
Trauma System Guidelines	
Multi-Casualty Incidents – 65.000	
MCI General Guidelines	
Hazardous Materials – 70.000	
Decontamination	
Hydrogen Cyanide	
Hydrogen Fluoride	
Organophosphates	70.040

Scope of Practice

Medical Control for Medications and Procedures

EMERGENCY MEDICAL RESPONDER SCOPE OF PRACTICE

An Emergency Medical Responder may:

- A. Conduct primary and secondary patient examinations;
- B. Take and record vital signs;
- C. Utilize noninvasive diagnostic devices in accordance with manufacturer's recommendation;
- D. Open and maintain an airway by positioning the patient's head;
- E. Provide external cardiopulmonary resuscitation and obstructed airway care for infants, children, and adults;
- F. Provide care for musculoskeletal injuries;
- G. Provide hemorrhage control;
- H. Provide emergency moves for endangered patients;
- I. Assist with prehospital childbirth;
- J. Complete a clear and accurate prehospital emergency care report form on all patient contacts and provide a copy of that report to the senior emergency medical services provider with the transporting ambulance;
- K. Administer medical oxygen;
- L. Maintain an open airway through the use of:
 - 1. A nasopharyngeal airway device;
 - 2. An oropharyngeal airway device;
 - 3. A pharyngeal suctioning device;
- M. Operate a bag mask ventilation device with reservoir;
- N. Provide care for suspected medical emergencies, including administering liquid oral glucose for hypoglycemia;
- O. Prepare and administer aspirin by mouth for suspected myocardial infarction (MI) in patients with no known history of allergy to aspirin or recent gastrointestinal bleed;
- P. Prepare and administer epinephrine by automatic injection device for anaphylaxis;
- Q. Administer and distribute short-acting opioid antagonist kit and distribute the necessary medical supplies to administer the short-acting opioid antagonist as provided in ORS 689.800;
- R. Perform cardiac defibrillation with an automated external defibrillator; and
- S. Perform other emergency tasks as requested if under the direct visual supervision of a physician and then only under the order of that physician.

EMERGENCY MEDICAL TECHNICIAN SCOPE OF PRACTICE

An EMT may:

- A. Perform all procedures that an Emergency Medical Responder may perform;
- B. Ventilate with a non-invasive manual or continuous positive pressure delivery device;
- C. Insert a supraglottic airway device to facilitate ventilation through the glottic opening by displacing tissue and sealing of the laryngeal area;
- D. Perform tracheobronchial tube suctioning;
- E. Provide care for suspected shock;
- F. Provide care for suspected medical emergencies, including:
 - 1. Obtain a capillary blood specimen for blood glucose monitoring;
 - 2. Prepare and administer epinephrine for anaphylaxis;
 - 3. Administer activated charcoal for poisonings; and
 - 4. Prepare and administer nebulized and metered dose albuterol or levalbuterol with or without ipratropium for known asthmatic and chronic obstructive pulmonary disease (COPD) patients suffering from suspected bronchospasm.
- G. Transport stable patients with saline locks, heparin locks, foley catheters, or in-dwelling vascular devices;
- H. Assist the on-scene Advanced EMT, EMT-Intermediate, or Paramedic by:
 - 1. Assembling and priming IV fluid administration sets; and
 - 2. Opening, assembling and uncapping preloaded medication syringes and vials;
- I. Complete a clear and accurate prehospital emergency care report form on all patient contacts;
- J. Assist a patient with administration of sublingual nitroglycerin tablets or spray and with metered dose inhalers that have been previously prescribed by that patient's personal physician and that are in the possession of the patient at the time the EMT is summoned to assist that patient;
- K. In the event of a release of organophosphate agents, the EMT who has completed Authorityapproved training may prepare and administer atropine sulfate and pralidoxime chloride by autoinjector, using protocols approved by the Authority and adopted by the supervising physician; and
- L. In the event of a declared Mass Casualty Incident (MCI) as defined in the local Mass Casualty Incident plan, monitor patients who have isotonic intravenous fluids flowing
- M. Administer over-the-counter medications in unit dose packaging for immediate use under specific written protocols authorized by the supervising physician or direct orders from a licensed physician.
- N. Acquire and transmit cardiac monitoring and electrocardiogram (ECG).
- O. Prepare and administer immunizations in the event of an outbreak or epidemic as declared by the Governor of the state of Oregon, the State Public Health Officer, or a county health officer, as part of an emergency immunization program, under the agency's supervising physician's standing order. Prior to vaccine administration, the EMT must be trained by the supervising physician or their designee. The EMT and the EMS agency or employer much maintain records or training;
- P. Prepare and administer immunizations for seasonal and pandemic influenza vaccinations according to the CDC Advisory Committee on Immunization Practices (ACIP), and/ or the Oregon State Public Health Officer's recommended immunization guidelines as directed by the agency's supervising physician's standing order. Prior to vaccine administration, the EMT must be trained by the supervising physician or their designee. The EMT and the EMS agency or employer much maintain records or training.

ADVANCED EMERGENCY MEDICAL TECHNICIAN SCOPE OF PRACTICE

Advanced Emergency Medical Technician (AEMT) may:

- A. Perform all procedures that an EMT may perform;
- B. Initiate and maintain peripheral intravenous (I.V.) lines;
- C. Initiate saline or similar locks;
- D. Obtain peripheral venous blood specimens;
- E. Initiate and maintain an intraosseous infusion; and
- F. Prepare and administer the following medications under specific written protocols authorized by the supervising physician or direct orders from a licensed physician:
 - 1. Analgesics for acute pain: nitrous oxide.
 - 2. Anaphylaxis: epinephrine;
 - 3. Hypoglycemia reversal agents:
 - a. Hypertonic dextrose;b. Glucagon;
 - 4. Intraosseous infusion anesthetic: Lidocaine;
 - 5. Bronchodilators:
 - a. Albuterol;
 - b. Ipratropium bromide;
 - 6. Vasodilators: nitroglycerin;
 - 7. Isotonic crystalloid solutions.
- G. Distribute medications at the direction of the Oregon State Public Health Officer as a component of a mass distribution effort. The AEMT must be trained by the supervising physician or their designee. The AEMT and EMS agency or employer must maintain records of the training; and
- H. Prepare and administer routine or emergency immunization and tuberculosis skin testing, as part of an EMS Agency's occupational health program, to the AEMT's EMS agency personnel, under the supervising physician's standing order. Prior to administration, the AEMT must be trained by the supervising physician or their designee. The AEMT and the EMS agency or employer must maintain records of training.

EMERGENCY MEDICAL TECHNICIAN – INTERMEDIATE SCOPE OF PRACTICE

An EMT-Intermediate may:

- A. Perform all procedures that an Advanced EMT may perform;
- B. Prepare and administer the following medications under specific written protocols authorized by the supervising physician, or direct orders from a licensed physician:
 - 1. Vasoactive medications:
 - a. Epinephrine;
 - 2. Antiarrhythmics:
 - a. Atropine sulfate;
 - b. Lidocaine;
 - c. Amiodarone;
 - 3. Analgesics for acute pain:
 - a. Morphine;
 - b. Ketorolac tromethamine;
 - c. Fentanyl;
 - 4. Antihistamine: Diphenhydramine;
 - 5. Anti-Emetic: Ondansetron;
- C. Insert an orogastric tube;
- D. Maintain during transport any intravenous medication infusions or other procedures which were initiated in a medical facility, if clear and understandable written and verbal instructions for such maintenance have been provided by the physician, nurse practitioner or physician assistant at the sending medical facility;
- E. Perform electrocardiographic rhythm interpretation; and
- F. Perform cardiac defibrillation with a manual defibrillator.
- G. Administer benzodiazepines for seizures or agitation. Prior to administration of benzodiazepines, the EMT-I must be trained by the supervising physician or their designee. The EMT-I and the EMS agency or employer must maintain records of training.

PARAMEDIC SCOPE OF PRACTICE

A Paramedic may:

- A. Perform all procedures that an EMT-Intermediate may perform;
- B. Initiate and maintain mechanical ventilation during transport if formally trained on the particular equipment and if acting under written protocols specific to the particular equipment.
- C. Initiate the following airway management techniques:
 - 1. Endotracheal intubation;
 - 2. Cricothyrotomy; and
 - 3. Transtracheal jet insufflation which may be used when no other mechanism is available for establishing an airway;
- D. Initiate a nasogastric tube;
- E. Provide advanced life support in the resuscitation of patients in cardiac arrest;
- F. Perform emergency cardioversion in the compromised patient;
- G. Transcutaneous pacing of bradycardia that is causing hemodynamic compromise;
- H. Initiate needle thoracostomy for tension pneumothorax;
- I. Obtain peripheral arterial blood specimens under specific written protocols authorized by the supervising physician;
- J. Access indwelling catheters and implanted central IV ports for fluid and medication administration;
- K. Initiate and maintain urinary catheters under specific written protocols authorized by the supervising physician or under direct orders from a licensed physician; and
- L. Prepare and initiate or administer any medications or blood products under specific written protocols authorized by the supervising physician or under direct orders from a licensed physician
- M. Interpret electrocardiogram (ECG).

Medical Control for Medications & Procedures – 00.040

The following drugs and procedures are considered **CATEGORY A** and will be used at the EMT clinician's discretion in accordance with these EMS Treatment Protocols.

Drugs – Category A:

- Acetaminophen
- Activated Charcoal (aspirin or acetaminophen < 2 hrs post ingestion)
- Adenosine (Adenocard[®])
- Albuterol (Ventolin®)
- Amiodarone (Cordarone[®])
- Aspirin
- Atropine Sulfate
- Calcium Gluconate
- Dexamethasone (Decadron[®])
- Dextrose
- Diltiazem
- Diphenhydramine (Benadryl[®])
- Dopamine (Intropin[®])
- Droperidol (Inapsine[®])
- DuoNeb (albuterol and ipratropium)
- Epinephrine
- Etomidate (Amidate[®])
- Fentanyl (Sublimaze[®])
- Glucagon
- Glucose, Oral
- Haloperidol (Haldol[®])
- Hydromorphone (Dilaudid®)
- Hydroxocobalamin (Cyanokit®)
- IV solutions
- Ibuprofen
- Ipratropium Bromide (Atrovent[®])
- Ketamine Hydrochloride
- Ketorolac Tromethamine (Toradol®)
- Labetalol
- Lidocaine
- Magnesium Sulfate (wide complex irregular tachycardia/torsades and
- adult asthma)
- Midazolam (Versed[®])
- Morphine Sulfate
- Naloxone (Narcan®)
- Nitroglycerin
- Norepinephrine (Levophed[®])
- Olanzapine (Zyprexa[®])

Medical Control for Medications & Procedures – 00.040

Drugs – Category A (continued):

- Ondansetron (Zofran®)
- Oxygen
- Oxymetazoline Hydrochloride (Afrin®)
- Oxytocin (Pitocin[®])
- Pralidoxime (Protopam[®] / 2-PAM[®]) •
- Rocuronium (Zemuron[®])
- Sodium Bicarbonate
- Succinylcholine
- Tranexamic Acid (TXA)
- Vecuronium (Norcuron[®])

Procedures – Category A:

- Defibrillation in cardiac arrest (to include DSED)
- Drug Assisted Airway Management (DAAM)
- End-tidal CO₂ monitoring
- Endotracheal intubation
- Emergency cricothyrotomy
- i-gel[®] Supraglottic Airway Device
- Intranasal medication administration
- Intraosseous access & infusion
- Intravenous access & infusion
- Left Ventricular Assist Device (LVAD) management
- Modified Valsalva Maneuver
- Non-invasive positive pressure ventilation
- Orogastric tube insertion and maintenance
- Patellar dislocation reduction
- Physical patient restraint
- PICC line access
- Pelvic immobilization with sling/wrap
- Pharmacological sedation of the agitated patient
- Positive end-expiratory pressure (PEEP)
- Sports equipment removal
- Suctioning
- Synchronized cardioversion
 - o Unstable V-Tach, OR
 - o SVT, unstable patient
- Taser barb removal
- Tension pneumothorax decompression

Medical Control for Medications & Procedures – 00.040

Procedures – Category A (continued):

- Tourniquet placement
- Transcutaneous pacing
- Ventilator management
- XSTAT

The following drugs and procedures are considered **CATEGORY B** and require On-line Medical Consult authorization. Confirmation of dosage or procedure will be obtained directly from a physician on duty at OLMC.

Drugs – Category B:

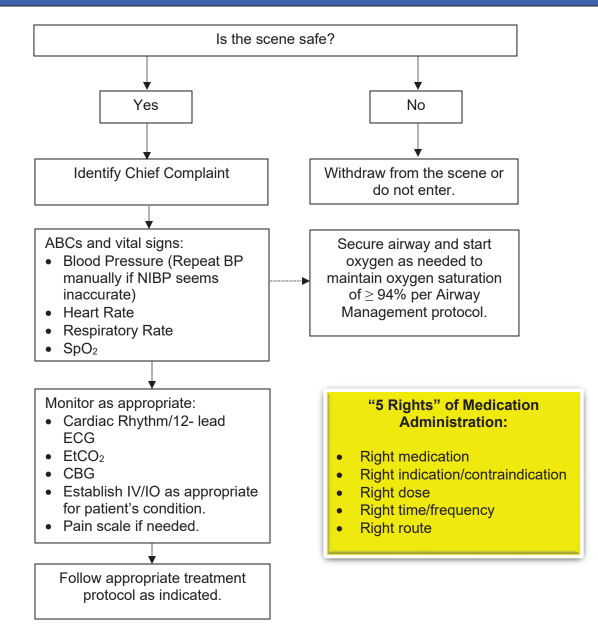
- Activated Charcoal (aspirin or acetaminophen > 2 hours post ingestion and all other poisons)
- Droperidol in patients ≤ 12
- Hydroxocobalamin (CYANOKIT®), repeat doses in pediatric patients
- Magnesium Sulfate (pediatric asthma OR seizures in eclampsia/pre-eclampsia)
- Ondansetron in patients < 6 months, except for children in spinal motion restriction or children receiving chemotherapy.
- Pralidoxime (2-Pam[®]), for IV use
- Sodium Bicarbonate for pediatric hyperkalemia and crush injuries
- Sodium Thiosulfate 25%

Procedures – Category B:

• Automatic Implantable Cardio-Defibrillator (AICD) deactivation with magnet.

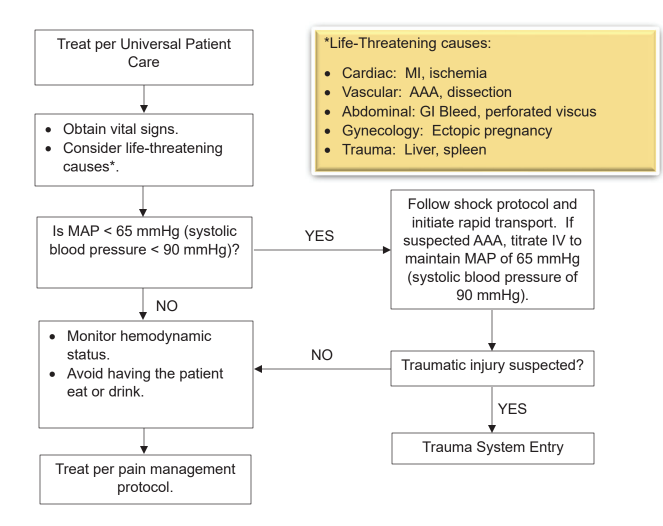
Treatment

Universal Patient Care – 10.005



NOTES & PRECAUTIONS

- If patient is unable to provide medical history, check for medical bracelets and necklaces, which can provide critical medical information and treatment.
- If any uncertainty exists about the gender of a patient, ask for and use preferred pronouns. In certain conditions such as abdominal pain, you may also need to ask about the menstrual history (e.g., female to male transgender). When obtaining a 12-lead ECG, use the sex assigned at birth for computerized interpretations.



NOTES & PRECAUTIONS:

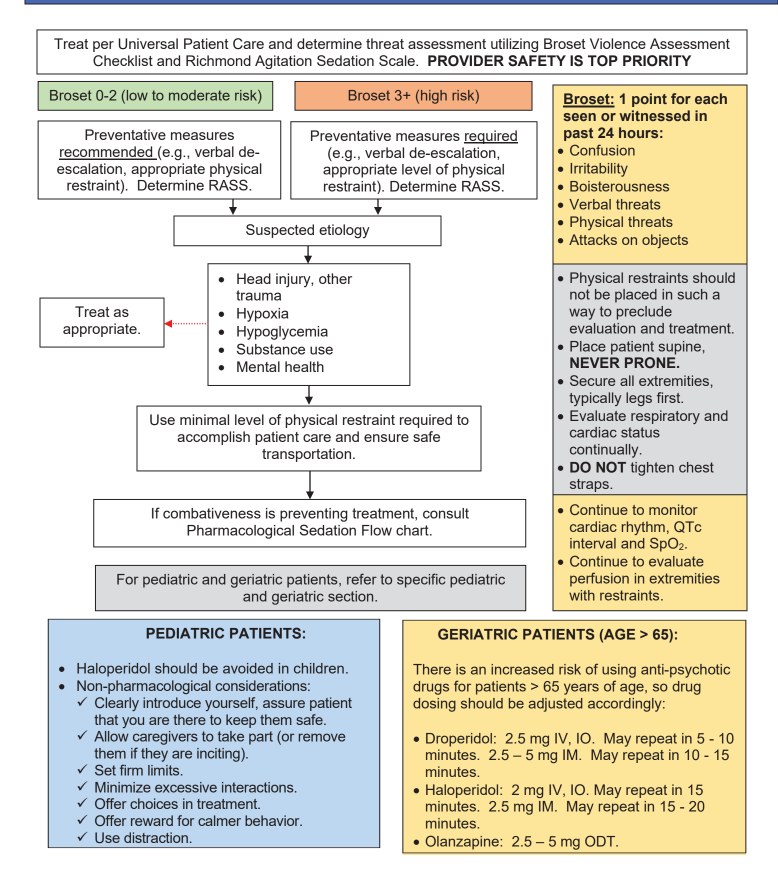
- Abdominal pain may be the first sign of catastrophic internal bleeding (ruptured aneurysm, liver, spleen, ectopic pregnancy, perforated viscus, etc.).
- Monitor the patient closely for signs of shock.
- For transgender and non-binary patients, ask about the presence of intact reproductive organs and consider gynecological (i.e., pregnancy issues) or urological (i.e., testicular torsion) related complications in your differential diagnosis.

PEDIATRIC PATIENTS:

- Consider non-accidental trauma.
- Closely monitor vital signs; blood pressure may drop quickly.
- If systolic BP is inappropriate for age, treat per Shock protocol.

Lowest normal pediatric systolic blood pressure by age:

- Less than one month: > 60 mmHg.
- One month to 1 year: > 70 mmHg.
- Greater than 1 year: 70 + 2 x age in years.

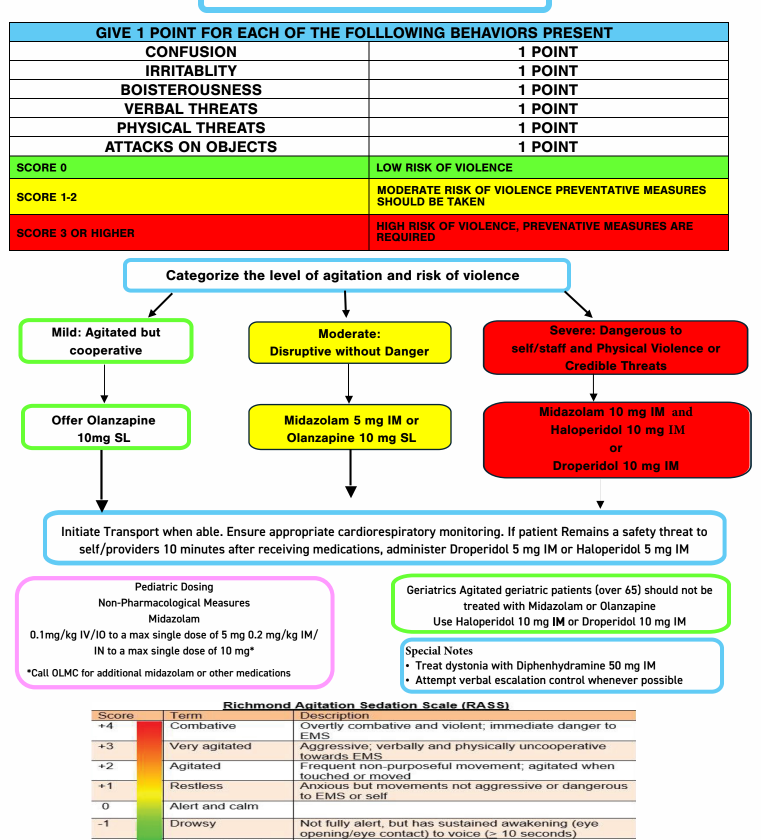


NOTES & PRECAUTIONS:

- All patients who receive IV, IO, or IM pharmacological sedation must be fully monitored, when possible, with cardiac monitor, SpO₂, and EtCO₂.
- Droperidol, haloperidol, and ziprasidone may induce Torsades de Pointes in patients with history of prolonged QTc or patients taking QTc-prolonging drugs. Monitor patient's ECG, if possible. If prolonged QTc is present (> 500 msec.), administer 2 grams magnesium sulfate IV/IO.
- Droperidol, haloperidol, or ziprasidone are preferred for patients with known psychiatric disorders.
 Midazolam is preferred for patients who are known or suspected to be under the influence of stimulants or other intoxicants, who are in withdrawal, or who are postictal.
- If patient has Parkinson's Disease or takes dopamine agonist medications such as carbidopa-levodopa (Sinemet), pramipexole (Mirapex), or ropinirole (Requip), <u>do not use</u> droperidol or haloperidol. In these patients, use olanzapine first (2.5 - 5.0 mg ODT), then midazolam (5 mg IM or 2.5 mg IV/IO) if needed.

Pharmacological Sedation Flow Chart

Assess Risk using Broset Check List



Briefly awakens with eye contact to voice (< 10

Movement or eye opening to voice (but no eye

 -4
 Deep sedation
 No response to voice but movement or eye opening to physical stimulation

 -5
 Unarousable
 No response to voice or physical stimulation

 Treatment - Revised 4/25/2025
 Vertical stimulation

Moderate sedation

seconds)

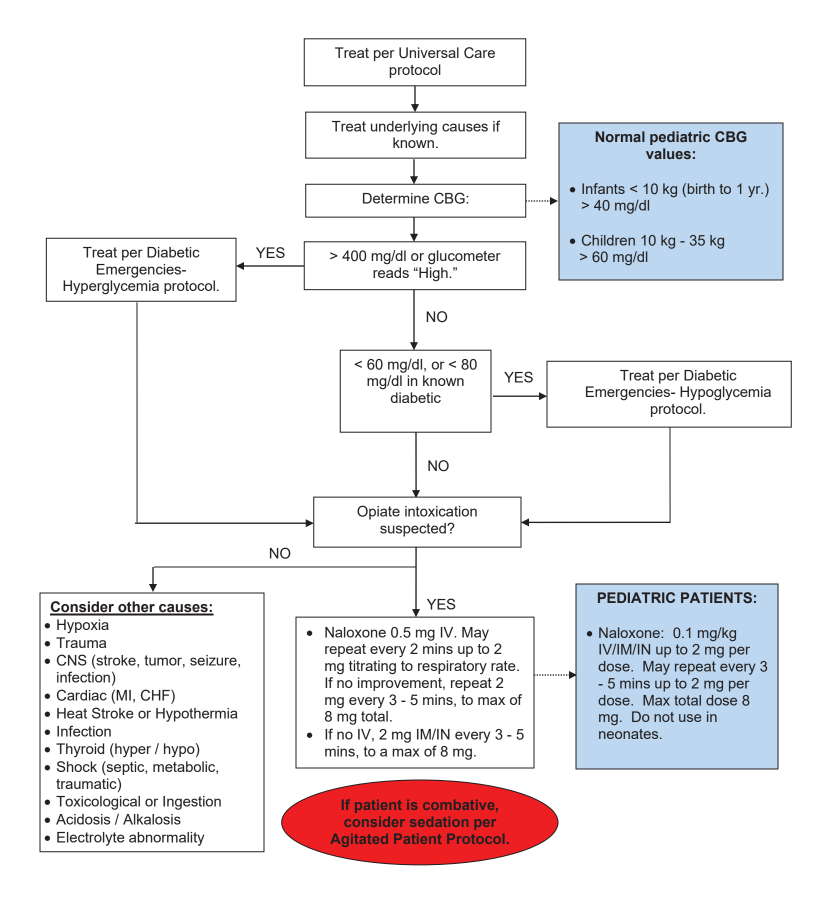
contact)

Light Sedation

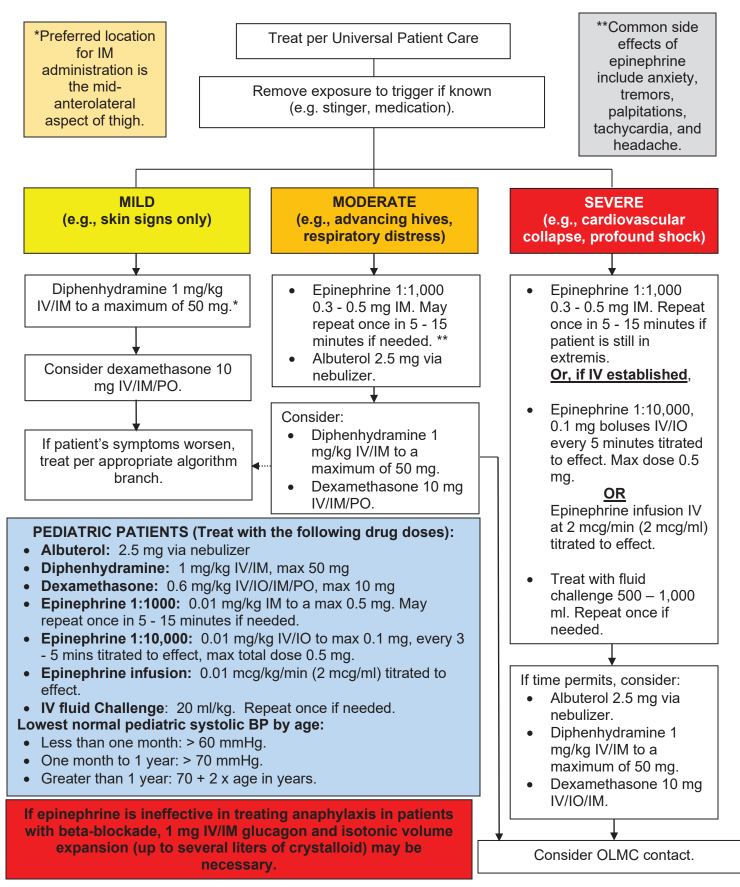
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Altered Mental Status & Coma – 10.020



Anaphylaxis and Allergic Reaction – 10.030



Treatment - Revised 4/5/24

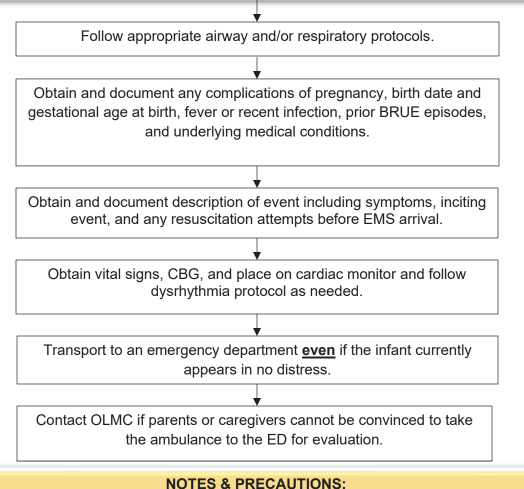
Brief Resolved Unexplained Event (BRUE) – 10.035

DEFINITION:

Event lasting <1 minute in an infant <1 year of age associated with at least one of the following:

- Cyanosis or pallor
- Absent, decreased, or irregular breathing
- Marked change in muscle tone (hypertonia or hypotonia)
- Altered level of responsiveness

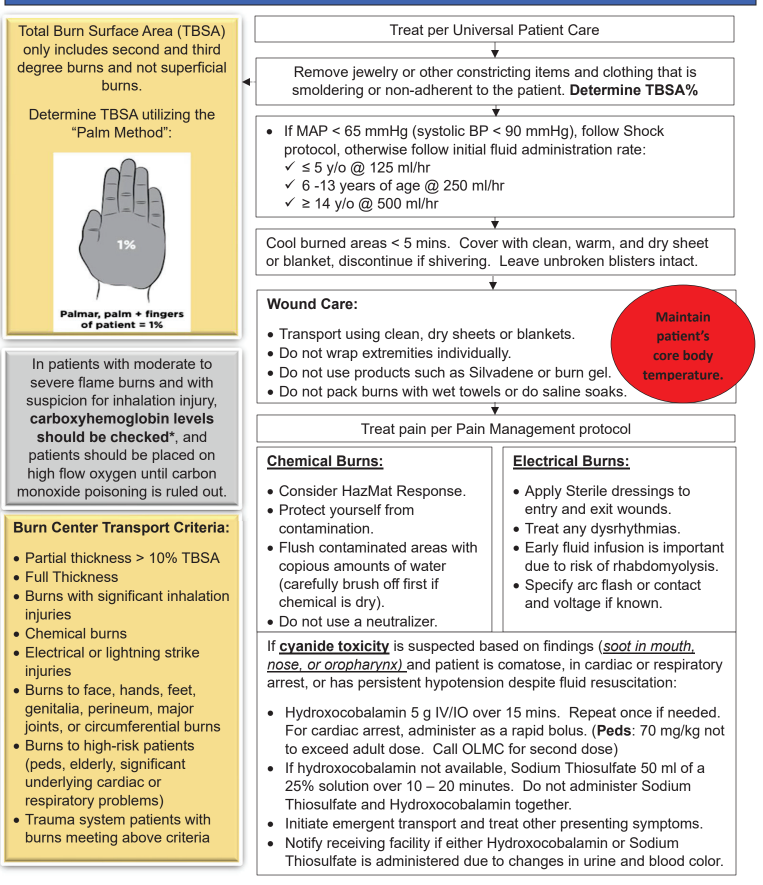
Patient must appear well and be at baseline health.



BRUE is a group of symptoms, not a specific disease. BRUEs are most common in

- infants under one year of age but may occur up to two years of age.
- Many infants appear normal by the time EMS arrives.
- Consider non-accidental trauma.
- Serious underlying causes of BRUE can include pneumonia, bronchiolitis, seizures, sepsis, intracranial hemorrhage, and meningitis.
- BRUEs are more frequent in premature infants and infants with other health conditions such as cystic fibrosis, bronchiolitis, and congenital heart disease.

Burns – 10.040



AIRWAY CONSIDERATIONS:

- Singed nasal hairs and facial burns alone are not indications for intubation.
- Mild inhalation injuries with normal SpO₂ and no signs of respiratory distress can be safely observed.
- Indications for early intubation:
 - ✓ Respiratory distress, stridor, accessor muscle use
 - ✓ New onset hoarseness
 - ✓ Blisters or edema of oropharynx
 - ✓ Deep burns to lower face or neck

PEDIATRICS:

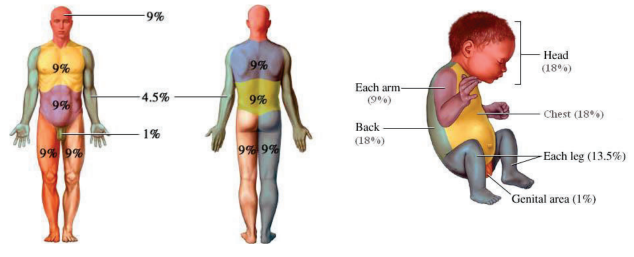
- Consider non-accidental causes of trauma in children.
- If systolic BP is inappropriate for age, treat per Shock protocol.

Lowest normal pediatric systolic blood pressure by age:

- Less than one month: > 60 mmHg.
- One month to 1 year: > 70 mmHg.
- Greater than 1 year: 70 + 2 x age in years.

NOTES & PRECAUTIONS:

- *Apply carbon monoxide monitor (e.g., Rad-57) if available.
- Remove rings or other constricting items immediately.
- Be prepared to use RSI/DSI early to control the airway.
- For firefighters, consider potential for traumatic injury or MI.



RULE OF NINES

Treatment - Revised 9/18/2024

Cardiac Arrest (AED/CPR) – 10.050				
CPR GUIDELINES				
Component	Adults and Adolescents	Child 1 year to puberty	Infant under 1 year, excluding neonates	
Airway	Head tilt-chi	n lift. Jaw thrust if suspected		
Breathing: Without CPR	10 to 12 breaths/min (approximate)	1 breath every 2 - 3 seconds (20 -30 breaths/min) (approximate)		
Breathing: CPR with advanced airway	1 breath every 6 secs. (10 breaths/min) asynchronous with compressions. About 1 second per breath. Visible chest rise. Optional method 30:2 comp./vent. ratio with advanced airway until ROSC.	1 breath every 2 - 3 seconds (approximately 20 - 30 breaths/min) asynchronous with compressions. About 1 second per breath. Visible chest rise. Optional method, 15:2 compression/ventilation ratio with advanced airway until ROSC.		
Foreign Body – Conscious patient	and obese patients or if	<i>chest thrusts in pregnant</i> <i>abdominal thrusts are not</i> <i>ctive</i>)		
Compression landmarks	Lower half of sternum between nipples		Just below nipple line (lower half of sternum)	
Hand Placement	Heel of one hand, other hand on top	As for adults (may use both hands or the heel of one hand depending on the size of patient and rescuer)	2 thumb-encircling hands preferred for two rescuers	
Compression depth	At least 2 inches	Approximately one-third anterior/posterior depth of chest. (Approximately 2" in child and 1 ½" in infant)		
Compression rate		100 - 120 per minute		
Compression/Ventilation ratio w/o advanced airway	30:2 or 10:1 with continuous compressions	15:2		
AED GUIDELINES				
AED Defibrillation	Use Adult pads	Use pediatric dose-attenuator system for children and infants if available. Use pediatric pads. If unavailable, use adult pads		
NEONATAL GUIDELINES (LESS THAN 1 DAY OLD)				
Assisted ventilations should be delivered at a rate of 40 - 60 breaths/minute to achieve or maintain a heart rate > 100 bpm. The ratio of compressions to ventilations should 3:1, with 90 compressions and 30 breaths to achieve approximately 120 events per minute.				

Cardiac Arrest- Guidelines – 10.050

COMPRESSIONS AND VENTILATIONS

- Use a pit crew approach to assign responders to positions.
- Initiate and maintain high quality chest compressions with limited interruptions (< 10 secs).
- CPR should be provided at a rate of 30:2 or continuous compressions with interposed ventilations every 6 seconds throughout resuscitation until ROSC is achieved or termination of resuscitation.
- There should be no interruptions to CPR when securing an airway. Consider early use of a supraglottic airway to minimize CPR interruptions or when ALS resources are limited.
- If mechanical CPR device is available, avoid extra or prolonged pauses in CPR when applying.

VASCULAR ACCESS

- Preferred order of vascular access in adults is:
 - 1. Upper extremity IV (or external jugular)
 - 2. Upper extremity IO
 - 3. Lower extremity IO
- Preferred access site for pediatrics is the proximal tibia or the distal femur. Humeral IO <u>not</u> recommended for infants and toddlers.
- Medications should be administered IV if multiple means of vascular access are established.

ROSC

If patient has return of spontaneous circulation, reassess vital signs to ensure stability before packaging for transport. Follow Cardiac Arrest Post-Resuscitation protocol to include targeted temperature management, obtaining a 12-lead ECG (ideally > 8 mins post ROSC), and managing blood pressure.

TRANSPORT

- In general, continue resuscitation for a minimum of 30 minutes.
- If persistent/refractory VF/pVT, consider early transport, especially if mechanical CPR is available.

EPINEPHRINE ADMINISTRATION

- For patients in a non-shockable rhythm, epinephrine should be administered as soon as feasible, ideally within 5 minutes of EMS arrival to patient side.
- For shockable rhythms, administer epinephrine as soon as feasible after the second defibrillation attempt has failed.

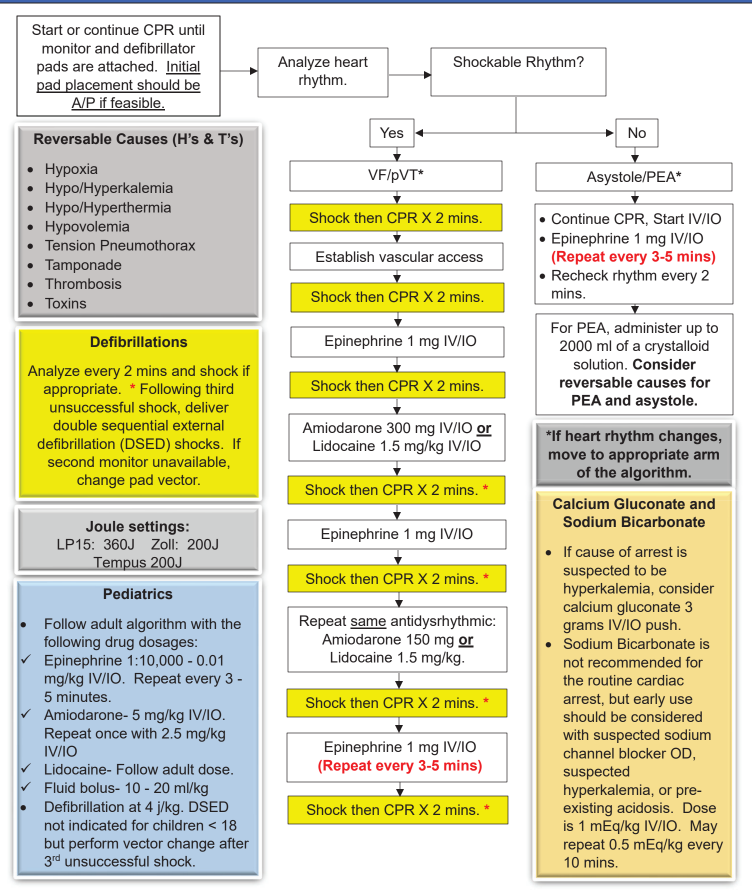
CPR INDUCED CONSCIOUSNESS

- With high quality CPR and the addition of mechanical CPR devices, a growing number of patients have been reported to experience "CPR Induced Consciousness". Assess for signs of consciousness by checking for spontaneous eye opening, purposeful movement, or verbal response including moaning.
- If signs of CPR Induced Consciousness are present, treat as follows (repeat vital signs between medications):
 - 1. 50 mcg of fentanyl IV/IO, then
 - 2. 2.5 mg of midazolam IV/IO _
 - 3. May repeat as needed every 5 10 minutes.

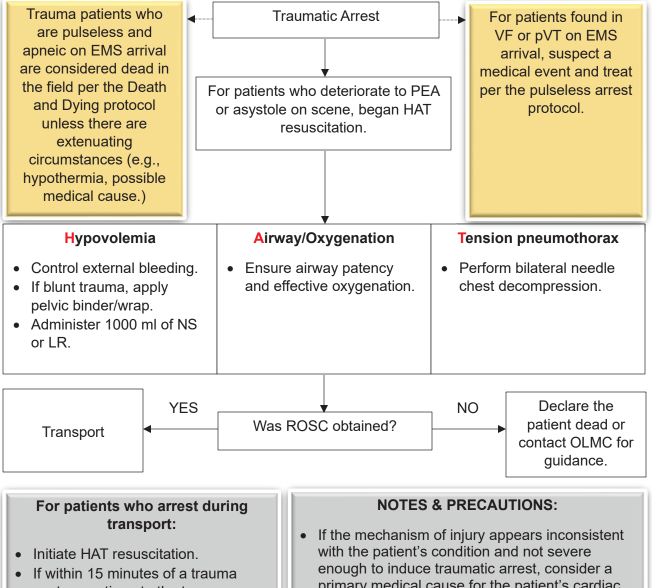
TERMINATION OF RESUSCITATION

- For patients in whom the asystole protocol has been used throughout the resuscitation, refer to Death and Dying protocol for guidelines regarding termination of resuscitation prior to 30 minutes without OLMC contact.
- Survival from PEA is based on identifying and correcting the responsible factors; consider a broad differential diagnosis, with early and aggressive treatment of possible reversible causes.
- Death in the field for PEA may be determined with EtCO₂ ≤ 10 after 30 minutes of attempted ACLS resuscitation. For patients with EtCO₂ > 10 continue resuscitation and contact OLMC to stop resuscitation.

Cardiac Arrest- Pulseless Arrest – 10.050

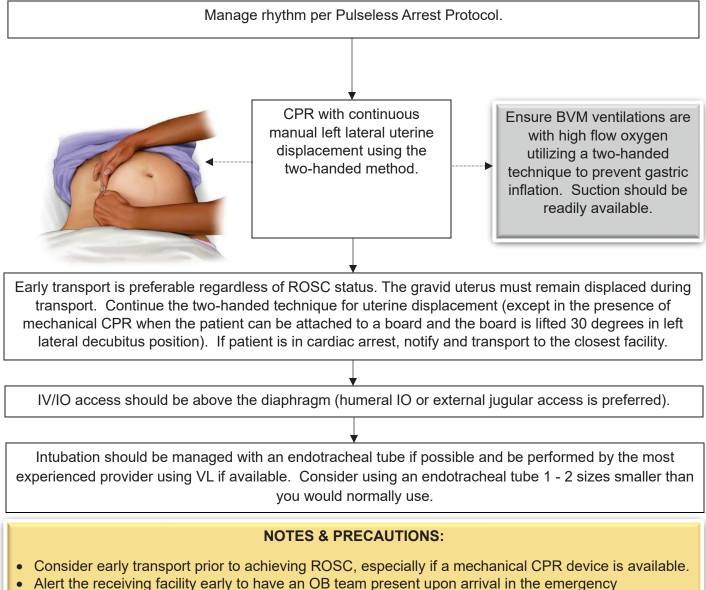


Cardiac Arrest (Trauma) – 10.050



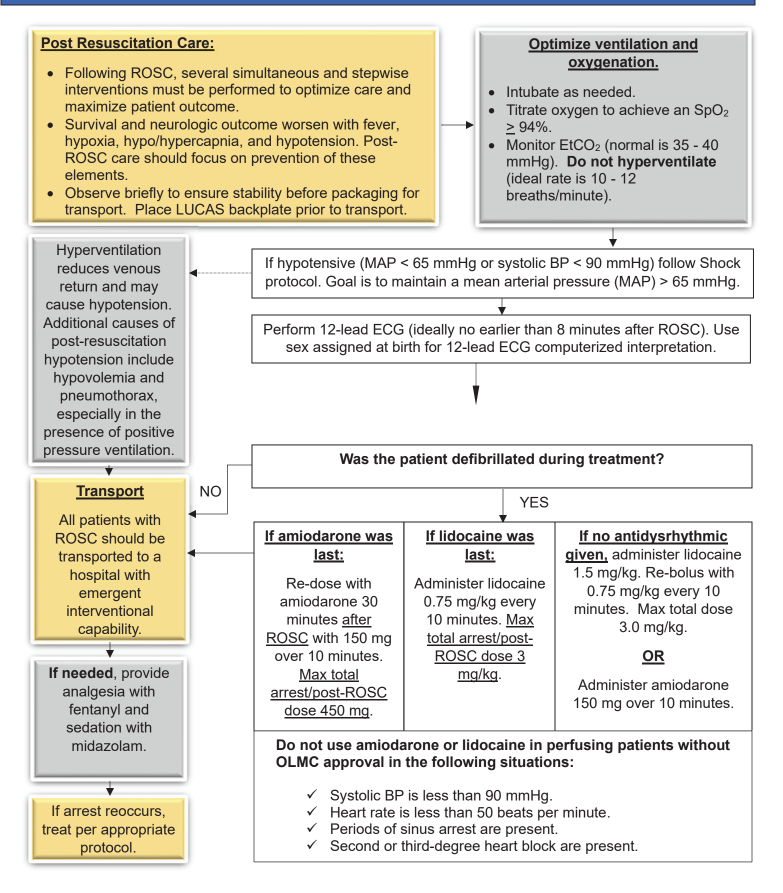
- center, continue to the trauma center.
- If farther than 15 minutes to the trauma center, consider pulling over for crew safety and personnel resource reasons. If ROSC is not achieved, you may declare the patient dead or contact OLMC for guidance.
- primary medical cause for the patient's cardiac arrest.
- If there is concern for a medical cause of the arrest, transport to the nearest cath lab capable facility if ROSC is achieved. If the patient is still in presumed medical cardiac arrest, then transport to the closest facility.
- · Perform chest compressions in traumatic arrest, but DO NOT allow compressions to interfere with addressing the reversible causes of a traumatic arrest in the HAT resuscitation.
- Post-ROSC cooling in the traumatic arrest patient should be deferred to the hospital.

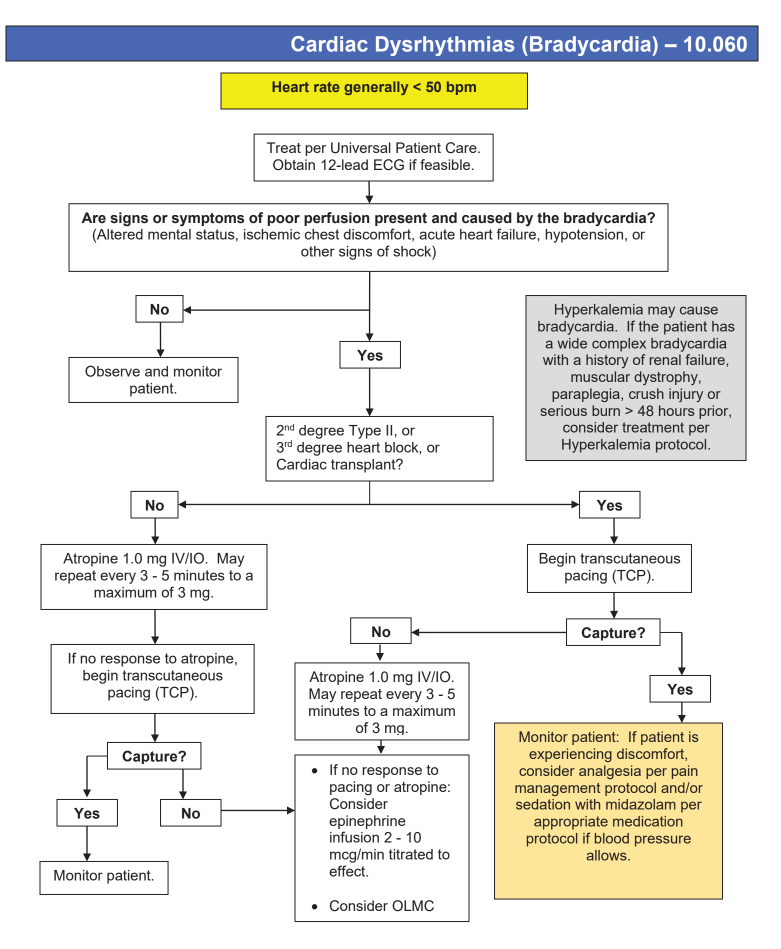
Cardiac Arrest with Pregnancy (> 22 weeks) – 10.050



- Alert the receiving facility early to have an OD team present upon anival in the emergency department. If you have not achieved ROSC, go to the closest facility regardless of OB capabilities.
 If ROSC has been achieved and maintained prior to, or during transport, bypass to an OB and NICU
- If ROSC has been achieved and maintained prior to, or during transport, bypass to an OB and NICU capable facility.
- Lidocaine is preferable (Class B in Pregnancy) to amiodarone (Class C in Pregnancy) in the setting of ventricular fibrillation or pulseless ventricular tachycardia.
- In the setting of ventricular fibrillation or pulseless ventricular tachycardia, no adjustments need to be made to defibrillation energy settings. Immediately following defibrillation, resume the left lateral uterine displacement.
- If mechanical CPR is in place, continue the left lateral uterine displacement by tilting the backboard 30° to the left or by continuing manual displacement.
- If ROSC is achieved, continue left lateral uterine displacement by placing the patient in the left lateral decubitus position or by manually displacing the gravid uterus.
- High flow oxygen needs to be maintained in all peri-arrest patients.
- Consider OG placement when possible.

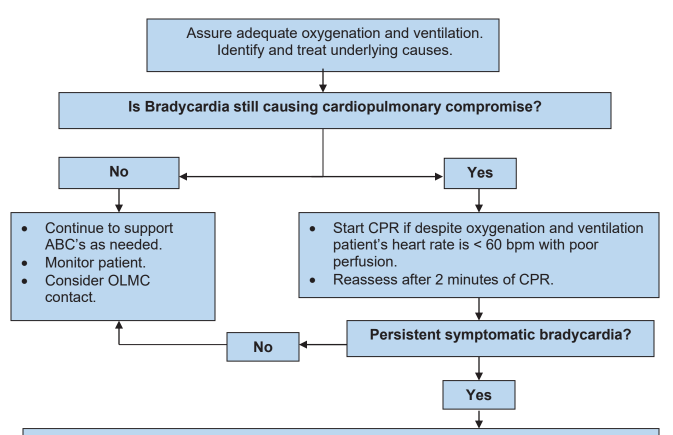
Cardiac Arrest Post Resuscitation – 10.050





PEDIATRIC PATIENTS:

BRADYCARDIA WITH A PULSE AND POOR PERFUSION



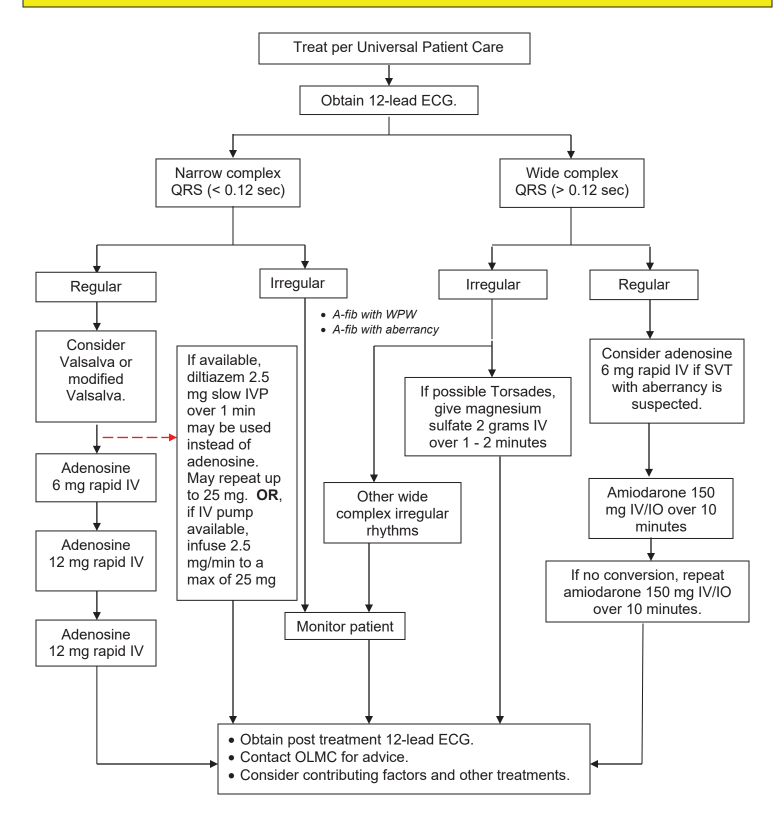
- Give 1:10,000 epinephrine 0.01 mg/kg IV/IO. Repeat every 3 5 minutes.
- Consider pacing per Transcutaneous Pacing procedure. If patient is experiencing discomfort, consider analgesia per pain management protocol and/or sedation with midazolam per appropriate medication protocol if blood pressure allows.
- If capture is not achieved, try repositioning pads.
- Goal of therapy is to improve perfusion.

NOTES & PRECAUTIONS:

- Hypoxia is a common cause of bradycardia.
- Bradycardia may be protective in the setting of cardiac ischemia and should only be treated if associated with serious signs and symptoms of hypoperfusion. Increasing heart rate may worsen ischemia or increase infarct size.
- Immediate TCP can be considered in unstable patients when vascular access is not available.
- TCP is at best a temporizing measure and is not useful in asystole.
- If TCP capture is not achieved, try repositioning pads.
- Atropine will likely be ineffective in heart transplant recipients because they lack vagal innervation.
- 3rd degree heart blocks with a wide complex QRS (>0.12 sec) are less likely to respond to atropine than those with a narrow complex.

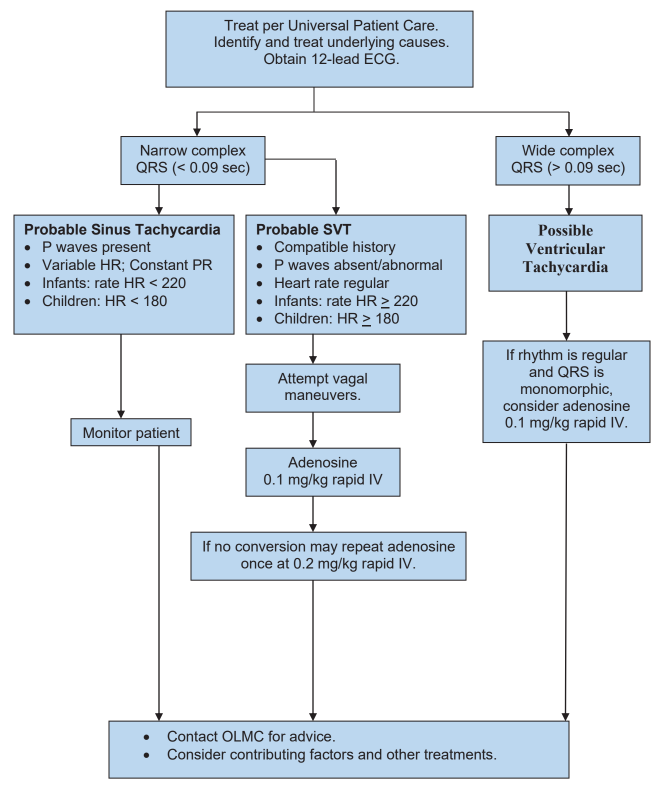
Cardiac Dysrhythmias (Tachycardia Stable) – 10.060

Patient <u>does not</u> have signs or symptoms of poor perfusion caused by the dysrhythmia. (e.g., Altered mental status, ischemic chest discomfort, acute heart failure, hypotension, or other signs of shock) *Rate related symptoms uncommon if HR <150 bpm. Consider other causes.*



Cardiac Dysrhythmias (Tachycardia Stable) – 10.060

PEDIATRIC PATIENTS:

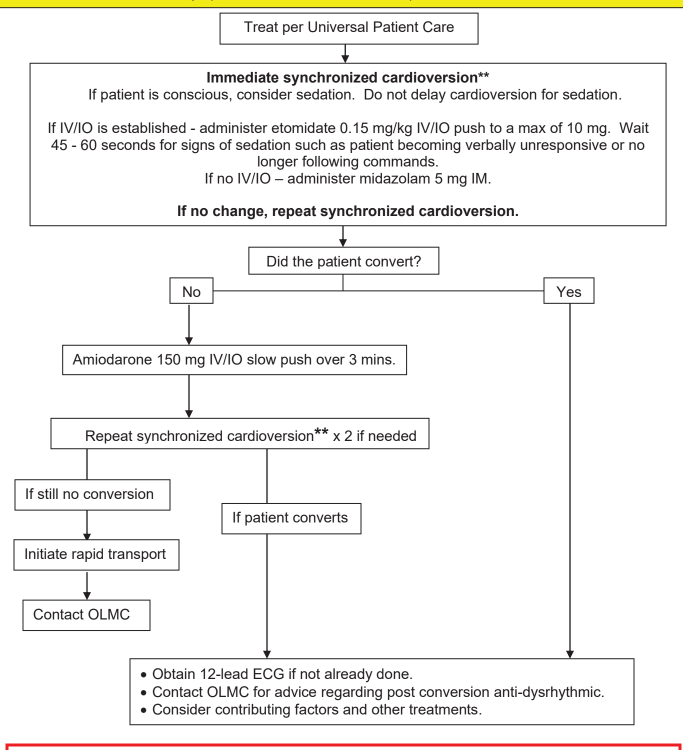


Cardiac Dysrhythmias (Tachycardia Stable) – 10.060

- In stable wide complex tachycardia, which is monomorphic, consider adenosine if SVT with aberrancy is suspected.
- If the patient is asymptomatic, tachycardia may not require treatment in the field. Continue to monitor the patient for changes during transport. The acceptable upper limit for heart rate for sinus tachycardia is 220 minus the patient's age.
- Other possible causes of tachycardia include:
 - ✓ Acidosis
 - ✓ Hypovolemia
 - ✓ Hyperthermia/fever
 - ✓ Hypoxia
 - ✓ Hypo/Hyperkalemia
 - ✓ Hypoglycemia
 - ✓ Infection
 - ✓ Pulmonary embolus
 - ✓ Tamponade
 - ✓ Toxic exposure
 - ✓ Tension pneumothorax
- If pulseless arrest develops, follow appropriate Cardiac Arrest protocol.
- All doses of adenosine should be reduced to one-half (50%) in the following clinical settings:
 - ✓ History of cardiac transplantation.
 - ✓ Patients who are on carbamazepine (Tegretol) and dipyridamole (Persantine, Aggrenox).
 - ✓ Administration through any central line.
- Adenosine may initiate atrial fibrillation with rapid ventricular response in patients with Wolff-Parkinson-White syndrome.
- Adenosine should be used with caution in patients with asthma as it may cause a reactive airway response in some cases.
- The Modified Valsalva Maneuver may increase the likelihood of converting SVT to sinus rhythm. Have the patient sit in an upright position. With the assistance of a 10 ml syringe, encourage the patient to strain for a full 15 seconds, trying to push out the plunger by forced expiration. Lay the patient flat and elevate their legs to 45-90 degrees for 15 seconds. Lay the patient's legs flat for 60 seconds. May repeat x1 if patient has not converted to sinus rhythm.
- Consider the following Valsalva techniques for pediatric patients:
 - ✓ For infants and toddlers, apply ice or chilled IV fluid to the patient's face.
 - \checkmark For preschool age and up, have the patient blow on a syringe.

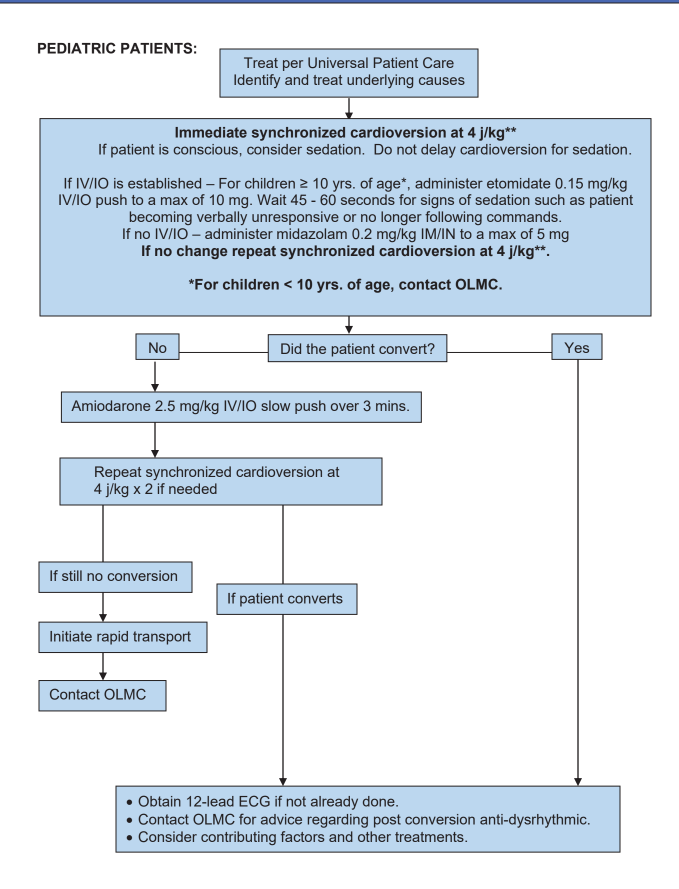
Cardiac Dysrhythmias (Tachycardia Unstable) - 10.060

Patient <u>has</u> signs or symptoms of poor perfusion caused by the dysrhythmia (e.g., Altered mental status, ischemic chest discomfort, acute heart failure, hypotension or other signs of shock). *Rate related symptoms uncommon if HR<150 bpm. Consider other causes.*



**If patient is in a wide complex irregular tachycardia use defibrillation (un-synchronized).

Cardiac Dysrhythmias (Tachycardia Unstable) – 10.060



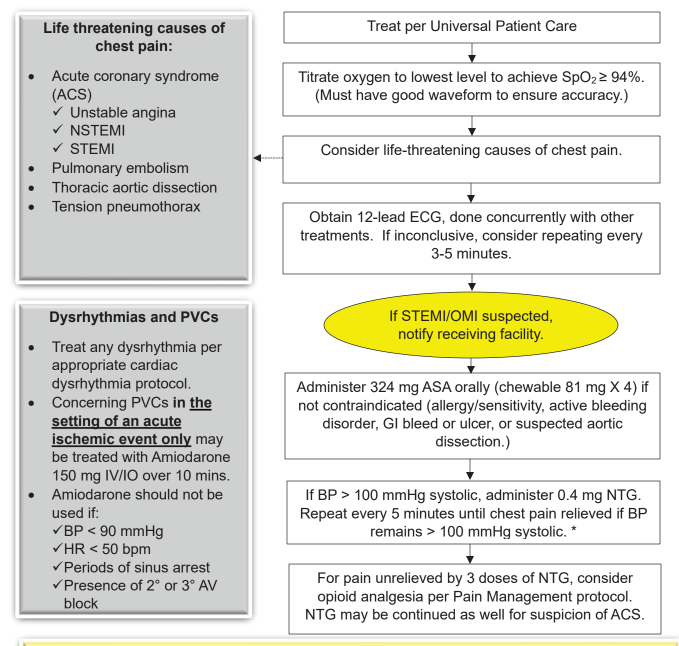
Cardiac Dysrhythmias (Tachycardia Unstable) – 10.060

- Possible causes of tachycardia include: •
 - ✓ Acidosis
 - ✓ Hypovolemia
 - ✓ Hyperthermia/fever
 - ✓ Hypoxia
 - ✓ Hypo/Hyperkalemia
 - ✓ Hypoglycemia
 - ✓ Infection
 - ✓ Pulmonary embolus✓ Tamponade

 - ✓ Toxic exposure
 - ✓ Tension pneumothorax
- If pulseless arrest develops, follow Cardiac Arrest protocol.
- Defibrillation is recommended for wide complex irregular tachycardia.
- Etomidate may result in myotonic jerking, apnea and/or pain at the injection site. •

Heart Monitor Adult Synchronous Cardioversion Settings (Joules)		
Physio LifePak [®]	360 j	
Zoll E/M Series®	200 j	
TEMPUS®	200 j	

Chest Pain/Acute Coronary Syndromes – 10.065



*Nitroglycerin Precautions

- Establish vascular access prior to administration for patients having not previously taken NTG or who are at risk of hemodynamic instability.
- NTG can cause hypotension in 10% of patients.
- Use with caution in patients with an inferior MI as profound hypotension can occur due to an associated right ventricular infarction (RVI can occur in up to 50% of inferior MIs).
- 12-lead clues to RVI include STE in III > II or STE ≥ 1 mm in V₄R. Current guidelines recommend avoidance of NTG in RVI.
- Do not administer NTG without OLMC if patient has taken sildenafil (Viagra[®]), vardenafil (Levitra[®]) in last 24 hours or tadalafil (Cialis[®]) in last 48 hours, given risk of profound hypotension with concomitant administration.

ST Elevation MIs (STEMIs)

- STEMI is defined by:
 - ✓ At least 1 mm ST elevation in two contiguous leads (except V2 and V3) in the absence of a LBBB or paced rhythm.
 - ✓ For leads V2 and V3, ≥ 2.5 mm STE for men < 40, ≥ 2 mm in men ≥ 40, and ≥ 1.5 mm in woman of all ages.</p>
- Field identified STEMI is a 12-lead ECG with:
 - ✓ Automatic ECG interpretation of "Acute MI", or
 - ✓ Paramedic concern for STEMI or OMI based on provider ECG review and clinical presentation.

Occlusive MIs (OMIs)

ECG findings concerning for an ongoing coronary occlusion also warrant cath lab activation. Findings consistent with OMI include:

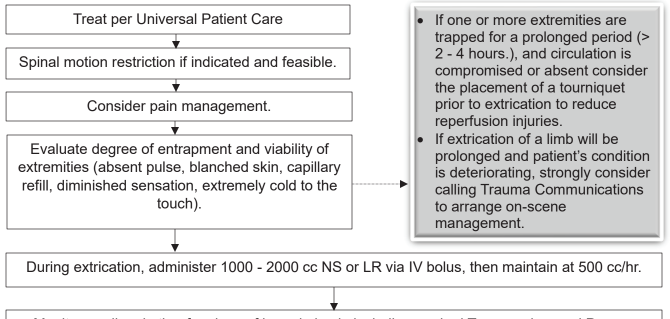
- Hyperacute T waves
- DeWinter T waves
- Mild inferior STE with reciprocal depression in aVL
- Anterior ST depression in the absence of posterior STE
- LBBB or paced rhythm with Smith-Modified Sgarbossa Criteria
- Wellens syndrome: Deep inverted T waves in V2/V3
- Aslanger's Pattern: Inferior STE in Lead III only, ST depression in any V4-V6 with positive T-Wave, ST segment in V1 > V2.

STEMI/OMI Actions

- If possible, transmit 12-lead ECG to destination hospital.
- Early notification to receiving hospital of "STEMI activation" ideally within 5 mins of identification.
- Apply defibrillation pads.
- Rapid transport to destination with cardiac interventional capability.

For pediatrics, consider pleuritic causes or trauma. Contact OLMC for advice

Crush Injury / Entrapment – 10.070



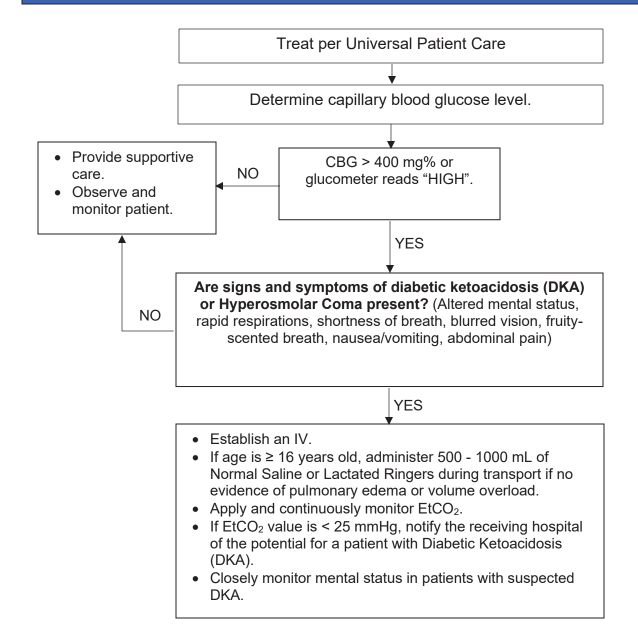
Monitor cardiac rhythm for signs of hyperkalemia including peaked T-waves, lowered P-wave amplitude or the loss of the P-wave, prolonged PR interval, second-degree AV block, and a widened QRS. If present, treat per Hyperkalemia protocol.

Wound Care

- Remove all restrictive dressings (clothing, jewelry, etc.).
- Monitor distal pulse, motor, and sensation in involved extremity.
- Bandage all open wounds (irrigate if needed).
- Stabilize all protruding foreign bodies (impaled objects).
- Splint/immobilize injured areas.
- For suspected pelvic crushing injuries, apply Pelvic Wrap if indicated.

- Crush injuries may elevate blood potassium levels (hyperkalemia) causing bradycardia, hypotension, weakness, weak pulse, and shallow respirations.
- Plan extrication activities to allow for periodic patient assessment. Plan for occasional extrication equipment "shut down" to assess vital signs.
- Carefully track vital signs, IV fluids, cardiac rhythm, and medications during extrication.
- Protect patient from environment (rain, snow, direct sun, etc.). If needed, begin warming methods (warm blankets, heated air with blower, warm IV fluids) to prevent hypothermia.
- Carefully assess collateral injuries that may have occurred during event.
- If patient is trapped in a heavy dust environment, consider methods to provide filtered oxygen to the patient. If patient is in respiratory distress, consider dust impaction injuries and prepare to administer nebulized albuterol per OLMC direction.
- Do not allow any personnel into extrication area (inner circle) without proper protective equipment and thorough briefing including review of the evacuation signal.
- Notify the receiving Trauma Center through Trauma Communications early in the extrication process for additional recommendations if needed.

Diabetic Emergencies- Hyperglycemia – 10.072



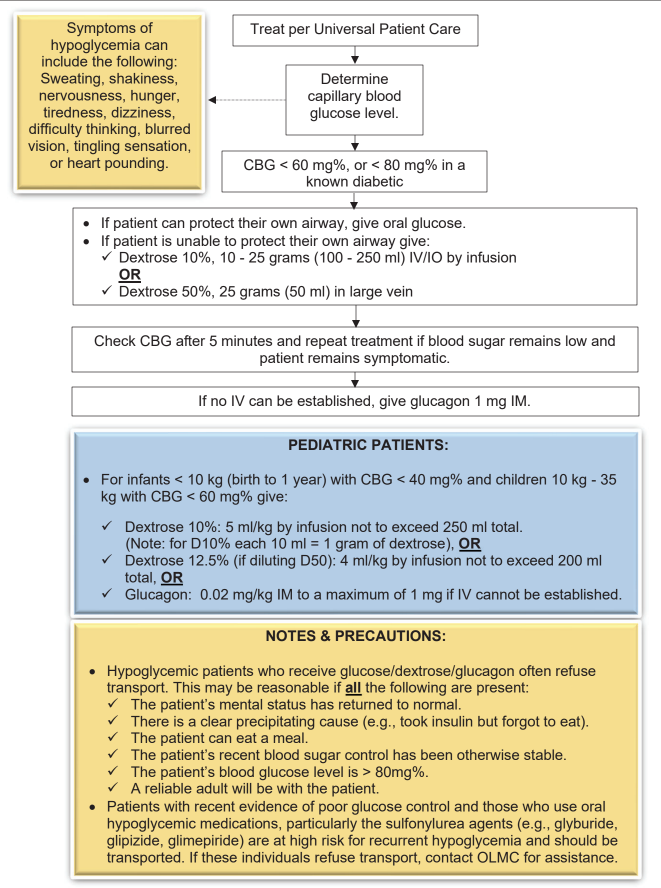
PEDIATRIC PATIENTS:

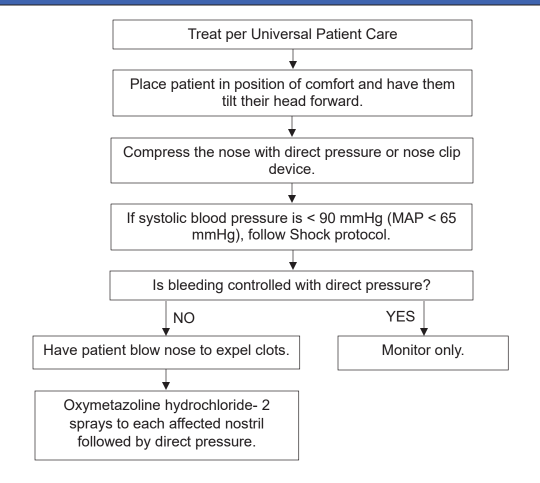
- Follow adult algorithm.
- If age is < 16 years old, consider administration of 10 mL/kg of Normal Saline or Lactated Ringers during transport if no evidence of pulmonary edema or volume overload.

NOTES & PRECAUTIONS:

If concern for DKA, avoid intubation unless the patient cannot protect their airway or there is evidence of extreme fatigue with an inability to ventilate or oxygenate. If intubation becomes necessary, the ventilation goal should be to maintain pre-intubation EtCO₂ levels.

Diabetic Emergencies- Hypoglycemia – 10.072



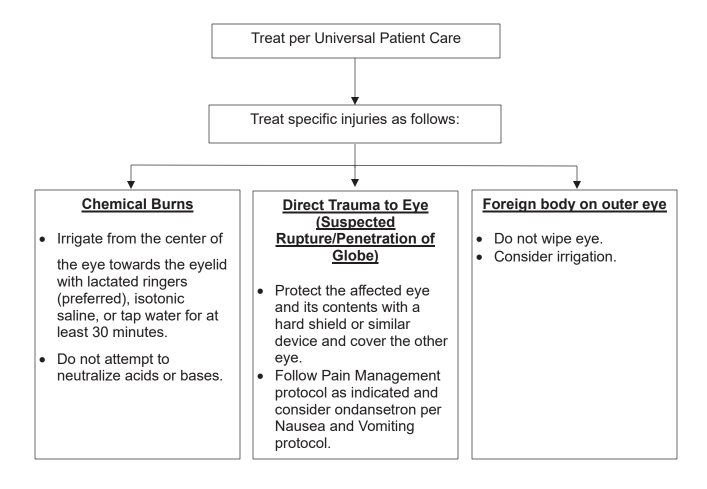


PEDIATRIC PATIENTS:

- Follow adult algorithm.
- Oxymetazoline Hydrochloride should be avoided if child cannot follow instructions to blow their nose or are unable to tolerate the administration of a nasal medication.

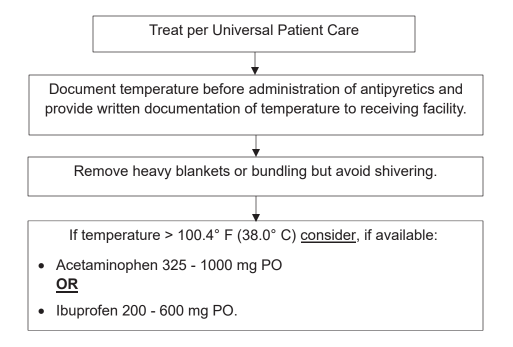
- Blood loss in epistaxis can be hard to quantify.
- Bleeding may be also occurring posteriorly. Evaluate for posterior blood loss by examining the back of the throat.
- Posterior epistaxis may be an emergency and may require advanced ED techniques such as balloon tamponade or interventional radiology. Do not delay transport. Be prepared for potential airway issues.
- Detailed medication history should be obtained to assess for the use of agents such as NSAIDs, antiplatelet agents, or anticoagulant medications that may contribute to bleeding.
- For patients on home oxygen via nasal cannula, place the cannula in the patient's mouth while the nares are compressed for active bleeding.

Eye Emergencies – 10.076



- Unless contraindicated, patients should be transported in a seated position of at least 30 degrees in order to decrease intraocular pressure.
- Document new onset of blurring, double vision, perceived flashes of light, or other visual changes.
- Contact lenses should be removed, if possible.

Fever Management – 10.077



PEDIATRIC DOSING

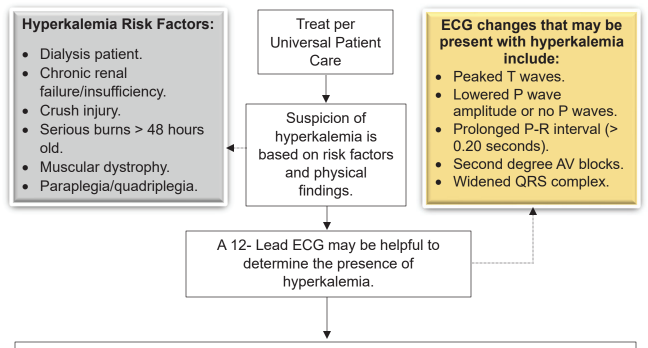
• Acetaminophen-

15 mg/kg PO liquid only to a maximum of 1000 mg

• Ibuprofen-

10 mg/kg PO <u>liquid only</u> to a maximum of 600 mg. **Do not give ibuprofen to children** less than 6 months old or with signs of dehydration.

- There is no evidence that treating fever decreases the likelihood of febrile seizure or has other therapeutic benefit. Treatment of fever is to improve patient comfort and is optional.
- Do not give acetaminophen if known liver disease, alcohol abuse, acute intoxication, or has taken acetaminophen in last 4 hours.
- Do not give ibuprofen to infants under 6 months, or patients with known renal disease, dehydration, ulcer, GI bleeding, gastric reflux disease (heartburn), pregnancy, or if a NSAID has been taken within the last 6 hours.
- Antipyretics are not indicated for environmental hyperthermia.



If hyperkalemia is suspected based on history and physical findings:

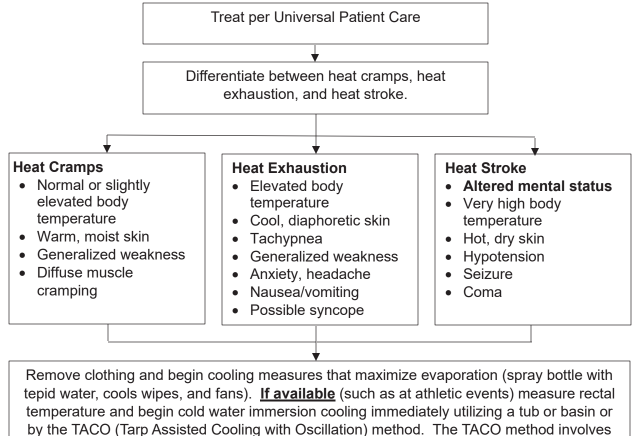
- Administer 10% calcium gluconate 1 3 grams IV/IO slowly over 5 10 minutes in a proximal port.
- If no change in rhythm following calcium administration and transport time is prolonged, consider alternate therapy:
 - ✓ High dose albuterol (10 mg by nebulizer)
 - ✓ Sodium bicarbonate 50 mEq IV/IO

<u>DO NOT</u> mix sodium bicarbonate solutions with calcium preparations. Slowly flush remaining calcium gluconate from the catheter prior to administering sodium bicarbonate.

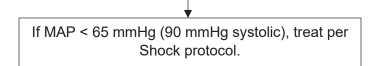
PEDIATRIC PATIENTS:

- Calcium gluconate- 0.6 ml/kg IV/IO slowly over 5-10 minutes. Max dose 10 ml.
- Albuterol-
 - ✓ < 25 kg, 2.5 mg via nebulizer
 - ✓ 25-50 kg, 5.0 mg via nebulizer
 - \checkmark > 50 kg, 10 mg via nebulizer
- Call OLMC regarding the use of sodium bicarbonate.

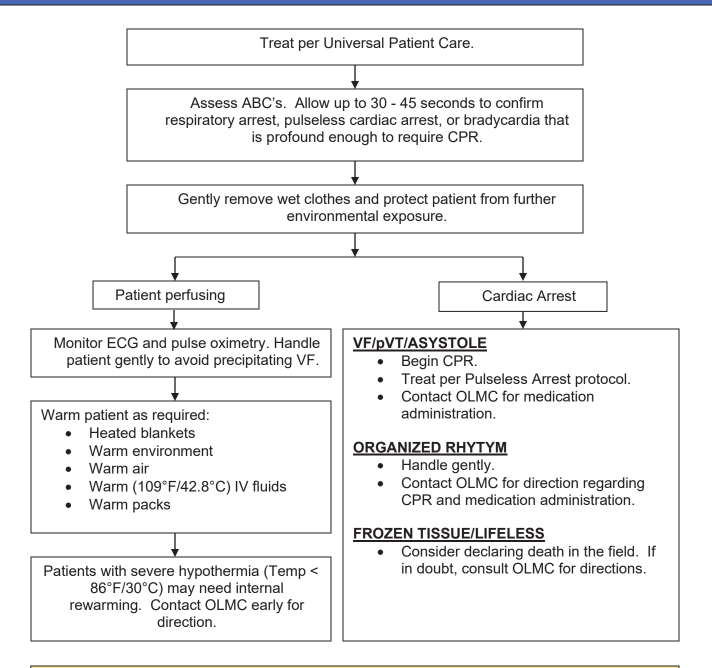
Hyperthermia/Heat-Related Emergencies – 10.080



placing a patient onto a tarp and pouring ice and cold water over the patient while simultaneously lifting the tarp and moving it back and forth. Athletic trainers are familiar with rectal temperature monitoring/TACO and can provide assistance.

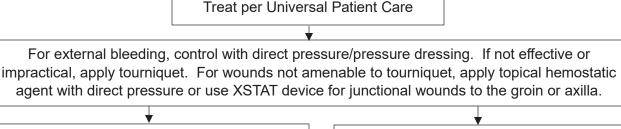


- Heat stroke is a medical emergency. Be aware that heat exhaustion can progress to heat stroke.
- Wet sheets over a patient without good airflow will increase temperature and should be avoided.
- Suspect hyperthermia in patients with altered mental status or seizure on a hot, humid day.
- Consider sepsis and/or contagious disease. Examine patient closely for rashes and nuchal rigidity.



- At-risks groups for hypothermia include trauma victims, alcohol and drug abuse patients, houseless persons, elderly, low-income families, infants and small children, and entrapped patients.
- Hypothermia may be preceded by other disorders (alcohol, trauma, OD, hypoglycemia) so look for and treat any underlying conditions while managing the hypothermia.
- The hypothermic heart may be unresponsive to cardiovascular medications, external cardiac pacing, or defibrillation.

Musculoskeletal Trauma – Extremity / Hemorrhage – 10.100



FRACTURES/SPRAINS/DISLOCATIONS

- Check for pulses, movement, and sensation (PMS), distal to the injury site before and after immobilization.
- Splint fractures/dislocations in the position found. If PMS is compromised distal to <u>fracture</u>, consider applying axial traction to bring extremity into normal anatomical position. If patient complains of increase in pain or resistance is felt, stop, and immobilize. If PMS is compromised distal to dislocation, contact OLMC.
- If fracture/dislocation is open, place a moist sterile dressing over wound and cover with a dry dressing.
- Elevate and/or place cold packs over fracture site if time/injuries allow.
- Apply traction splint to femur shaft fractures.
- For suspected pelvic fractures, utilize pelvic sling and secure to backboard to minimize blood loss.
- Treat per Pain Management protocol.

AMPUTATIONS

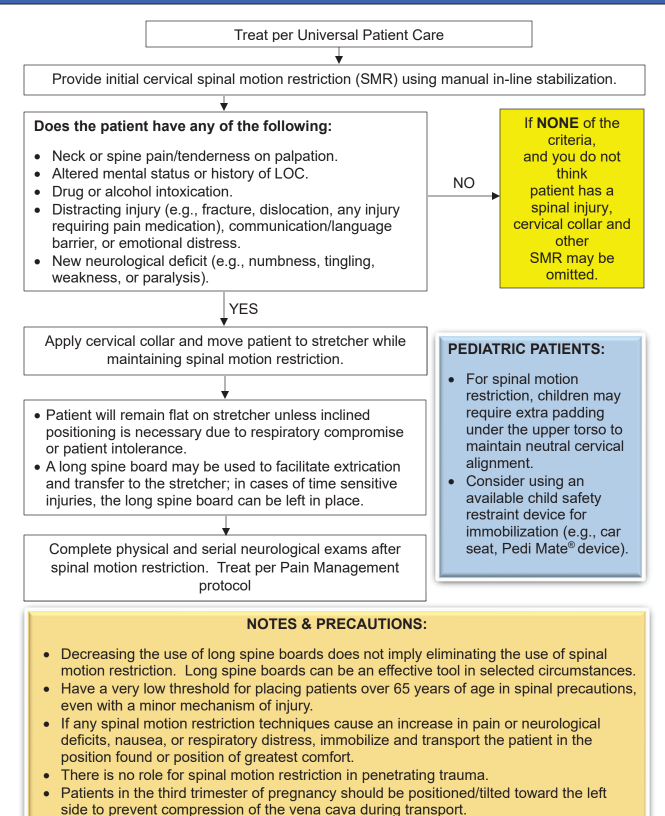
- Cover stump or partial amputation with moist sterile dressing.
- Splint partial amputations in anatomical position to avoid torsion and angulation.
- Wrap amputated part in a sterile dressing, and place in a plastic bag to keep dry. Place bag in ice water if available.
- If transport time is prolonged (extended extrication, etc.) consider sending the amputated part ahead to be prepared for reimplantation.
- Treat per Pain Management protocol.

PEDIATRIC PATIENTS:

- Treat pain per Pain Management protocol.
- Consider non-accidental trauma.

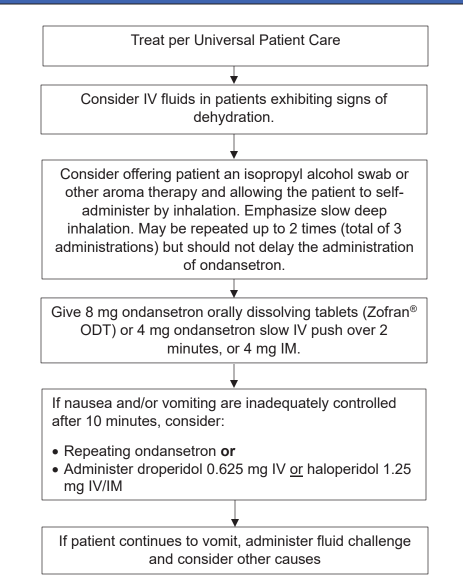
- Use of tourniquet for extremity hemorrhage is strongly recommended if sustained direct pressure is ineffective or impractical; use a commercially produced, windlass, pneumatic, or ratcheting device, which has been demonstrated to occlude arterial flow and avoid narrow, elastic, or bungee-type devices. Utilize improvised tourniquets only if no commercial device is available. If an improvised tourniquet is present before medical provider arrival, place a commercial tourniquet per protocol and remove the improvised tourniquet if operationally feasible. Time tourniquet was placed must be recorded.
- Apply a topical hemostatic agent, in combination with direct pressure, for wounds in anatomical areas where tourniquets cannot be applied, and sustained pressure alone is ineffective or impractical. Only apply topical hemostatic agents in a gauze format that supports wound packing.
- XSTAT is for the control of severe, life-threatening bleeding from junctional wounds in the groin or axilla that are not amenable to tourniquet applications in adults and adolescents. It should only be used for patients at high risk for immediate life-threatening bleeding from hemodynamically significant, non-compressible junctional wounds.

Musculoskeletal Trauma - Spinal Injury – 10.100



- If feasible, especially in prolonged scene transports, pad backboards.
- If sports injury, follow Sports Equipment Removal protocol.

Nausea & Vomiting – 10.110

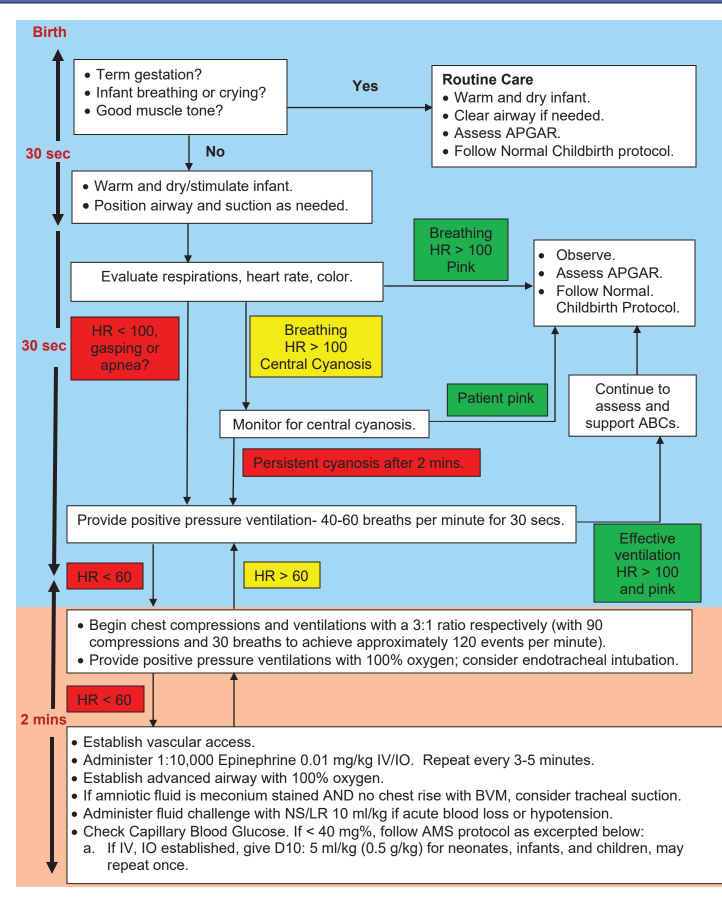


PEDIATRIC PATIENTS:

- Ondansetron use in patients under 6 months of age requires OLMC consultation except for children in spinal immobilization or children receiving chemotherapy.
- For children 6 months 2 years of age, administer 2 mg ondansetron orally dissolving tablet (Zofran[®] ODT). For children 2 12 years of age, administer 4 mg ondansetron orally dissolving tablet (Zofran[®] ODT) or administer ondansetron 0.1mg/kg via slow IV push over 2 minutes up to a total maximum single IV dose of 4mg. Consider IM at same dose if unable to start IV and ODT tablet is contraindicated.

- Do not administer ondansetron (Zofran[®]) to patients with a hypersensitivity to the drug or other 5-HT₃ type serotonin receptor agonists (e.g., dolasetron, palonosetron, and granisetron.)
- Do not administer alkaline medications or preparations in the same IV as ondansetron as it may cause precipitation.

Neonatal Resuscitation – 10.120



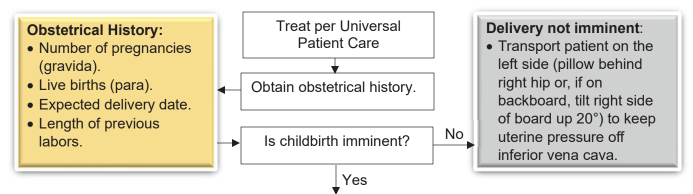
POST RESUCITATION CARE:

- Continue to provide assisted ventilations as needed.
- Closely monitor respiratory effort, heart rate, blood glucose, and pulse oximetry.
- Keep newborn normothermic. Hypothermia significantly increases risk of morbidity.
- Babies who required prolonged PPV, intubation and/or chest compressions are likely to have been severely stressed and are at risk for multi-organ dysfunction that may not be immediately apparent.

- Tracheal suctioning <u>is not</u> indicated in the vigorous infant born with meconium-stained fluid, whatever the consistency. Simply use a bulb syringe or large bore catheter to clear secretions from the mouth and nose as needed. However, if the newborn is having respiratory distress, then meconium aspiration should be performed per suctioning protocol.
- Volume expanders should not be given during resuscitation in the absence of a history or indirect evidence of acute blood loss. Giving a large volume load to a baby whose myocardial function is already compromised by hypoxia can decrease cardiac output. If fluid resuscitation is needed, administer 10 ml/kg NS over 5 - 10 minutes. Contact OLMC for repeat dosing.
- An electronic cardiac monitor is the preferred method for assessing heart rate.
- The ratio of compressions to ventilations should be 3:1, with 90 compressions and 30 breaths to achieve approximately 120 events per minute.
- Pulse oximeter should be applied to the right hand preferentially.
- 100% oxygen should not be used to initiate resuscitation. Begin resuscitation with room air and add supplemental oxygen if infant remains cyanotic or oxygen saturation < 70% after 2 minutes.
- Expected oxygen saturation of full-term newborn:

1 min	60% - 65%
2 min	65% - 70%
3 min	70% - 75%
4 min	75% - 80%
5 min	80% - 85%
10 min	85% - 95%

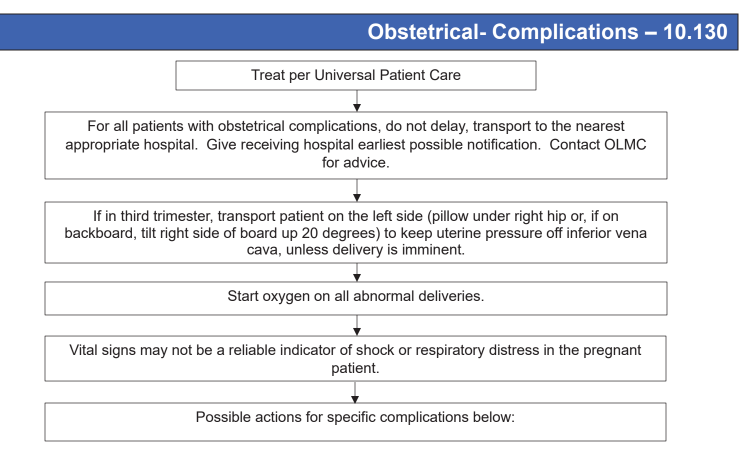
Obstetrical- Childbirth – 10.130

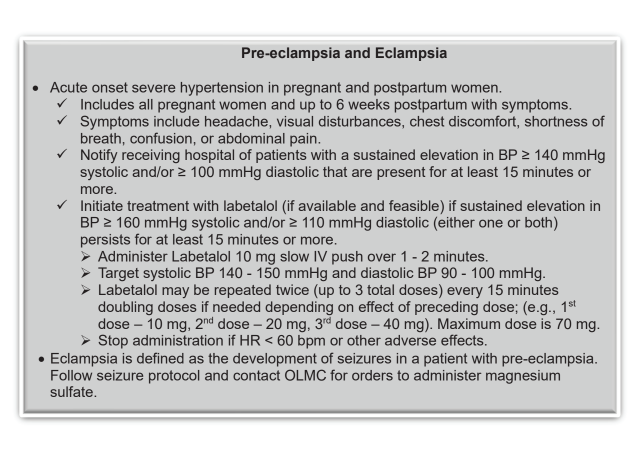


Normal Childbirth Procedure:

- Ask if the patient has had prenatal ultrasound and the possibility of multiple births. If multiple, or abnormal birth, consider second transport unit.
- Use sterile or clean technique.
- Guide/control but do not retard or hurry delivery.
- Check for cord around neck and gently remove if found.
- After delivery, assess infant per Neonatal Resuscitation protocol. If no resuscitation is needed (term infant, breathing or crying, good muscle tone), proceed as below.
 - ✓ Administer 10 IU oxytocin IV/IM within one minute of delivery when feasible if the neonate is a singleton. For multiple births, administer oxytocin only after last neonate has delivered.
 - ✓ Do not suction infant's nose and mouth unless there is meconium present, <u>and</u> the infant is depressed; or there is a need to clear the airway.
- Briefly dry infant and place on mother's chest, in skin-to-skin contact. Cover both with a clean, dry blanket.
- Assess infant using APGAR at time of birth and five minutes later. (Documentation should describe the infant using criteria rather than giving a numerical score).
- At 30 60 seconds after delivery, clamp and cut the umbilical cord about 6 inches from infant after cord pulsations have ceased. If resuscitation is needed, cord may be clamped and cut as soon as necessary.
- Do not delay transport to deliver the placenta. After the placenta has delivered, gently externally massage uterus to encourage contractions and prevent bleeding.
- If mother has significant postpartum hemorrhage (> 500 ml), administer tranexamic acid 2 g slow IV push.
- Unless infant needs treatment, keep on mother's chest for transport.
- Monitor vital signs of mother and infant during transport.

APGAR SCORE	0	1	2
Appearance	Blue/Pale	Body pink, extremities blue	Completely pink
Pulse	Absent	Slow (<100 bpm)	> 100 bpm
Grimace	No response	Grimace	Cough or sneeze
Activity	Limp	Some flexion	Active motion
Respirations	Absent	Slow, irregular	Good, crying





Breech Delivery (buttocks first):

- If delivery is imminent, prepare the mother as usual and allow the buttocks and trunk to deliver spontaneously then support the body while the head is delivered.
- If the head does not deliver within three minutes, suffocation can occur.
 - \checkmark Place a gloved hand into the vagina, with your palm toward the baby's face.
 - ✓ Form a "V" with your fingers on either side of the baby's nose and push the vaginal wall away from the baby's face to create airspace for breathing.
 - ✓ Assess for the presence of pulse in umbilical cord if able.

Limb Presentation

- The presentation of an arm or leg through the vagina is an indication for immediate transport to the hospital.
- Assess for presence of pulse in umbilical cord if presenting.

Prolapsed Cord

- Place the mother in left lateral Trendelenburg position.
- If the cord is visible, gently displace presenting part of baby off cord and maintain displacement. DO NOT pull or over-handle cord to prevent cord compression and spasm.

Abruptio Placenta

- Occurs in the third trimester of pregnancy when the placenta prematurely separates from the uterine wall leading to intrauterine bleeding.
- The patient experiences lower abdominal pain and the uterus often becomes rigid.
- Shock may develop without significant vaginal bleeding (concealed abruption).

Placenta Previa

• Occurs when the placenta covers the cervical opening, which can result in vaginal bleeding and prevents delivery of the infant through the vagina. The infant needs to be delivered via caesarian section.

Pain Management – 10.135

Treat per Universal Patient Care

- Consider and treat underlying causes of pain.
- Use non-pharmacological pain management (i.e., position of comfort, hot/cold pack, elevation, splinting, padding, wound care, and therapeutic calming and communication).

Determine location of pain and severity using numeric scale (1 - 10) or faces scale.

↓ For mild pain, consider:

- Acetaminophen 325 1000 mg PO, or
- Ibuprofen 200 600 mg PO

Controlled medications (opioids and ketamine) are to be avoided in the following patients: Active labor, headache, non-traumatic neck or back pain, any chronic pain (head, neck, back, fibromyalgia, abdominal/pelvic pain), or dental pain. <u>Contact OLMC</u>

- Monitor SpO₂ and EtCO₂.
- Document vital signs, response to treatment and pain scale rating prior to and after each administration of pain medication.

Opioids and dissociative medications (ketamine) can be used in the same patient to achieve pain relieve if necessary. For moderate to severe pain, consider:

Non-Opioid medications

• Ketorolac (patients aged 2 - 80): 30 mg IM or 15 mg IV. Do not repeat. Use for musculoskeletal pain or flank pain with suspected kidney stones.

Opioid medications

- Fentanyl: 25 100 mcg IV/IN/IM. May repeat with 25 100 mcg for IV/IN or 50 100 mcg for IM q 10 15 mins to max of 500 mcg. If BP < 100 mmHg or minor AMS or resp. depression, the first dose is 25 mcg all routes, repeating with 25 50 mcg q 10 -15 mins, to max of 500 mcg. Monitor patient closely.
- Hydromorphone: 0.25 0.5 mg IV or 0.5 1.0 mg IM q 15 20 mins., to max of 2 mg. Do not administer if systolic BP < 100 mmHg.
- **Morphine:** 2 10 mg IV q 15 20 mins, to max of 20 mg. Or, 5 10 mg IM, repeating with 5 mg q 15 20 mins, to a max of 20 mg. Do not administer if systolic BP < 100 mmHg.

Dissociative medications

Ketamine: 12.5 - 25 mg IV/IO slowly over 5 mins, or by IV infusion over 15 mins., or 25 - 50 mg IM. May repeat once in 30 mins., unless patient develops nystagmus, hallucinations, or other psychiatric symptoms. Must be diluted prior to IV or IO administration to a min. of 10 ml for slow IV push or 100 ml for IV infusion. Aternatively,1 mg/kg <u>VIA BREATH ACTUATED</u> <u>NEBULIZER (BAN)</u> MAY be used. Add saline for total volume of 5 ml.

PEDIATRIC PATIENTS:

- Acetaminophen: 15 mg/kg PO liquid only to a maximum of 1000 mg.
- **Ibuprofen:** 10 mg/kg PO <u>liquid only</u> to a maximum of 600 mg.
- Ketorolac (age 2 16 years): 1 mg/kg IM to a max of 30 mg or 0.5 mg/kg IV to a max of 15 mg. Do not repeat.
- Fentanyl (not to exceed adult dose):
 - ✓ 1 mcg/kg IV. May repeat with 0.5 1 mcg/kg every 10 15 minutes as needed to a maximum of 4 mcg/kg IV.
 - ✓ 2 mcg/kg IN. May repeat with 1 mcg/kg every 10 15 minutes as needed to a maximum of 4 mcg/kg IN.
 - ✓ If no IV/IN, may give fentanyl 1 2 mcg/kg IM. May repeat every 10 15 minutes to a max of 4 mcg/kg IM.
 - ✓ IN is preferred if no IV.
- Hydromorphone: For patients ≥ 12 months: 0.01 mg/kg IV/IM not to exceed the adult dose. May repeat every 15 - 20 minutes to a maximum of 2 mg. Hydromorphone is not preferred in young infants and toddlers if fentanyl or morphine is available.
- **Morphine:** 0.1 mg/kg IV or IM. (IM may repeat after 15 20 minutes). Do not exceed adult dosing.
- Ketamine: For children ≥ 15, dose is 0.3 mg/kg IV slow push over 5 minutes, up to a max of 25 mg. Dose must be diluted in normal saline prior to administration. Alternatively, for children ≥ 7, 1 mg/kg VIA BREATH ACTUATED NEBULIZER (BAN) MAY be used. Add saline for total volume of 5 ml.
- Do not administer fentanyl, morphine or hydromorphone if patient's systolic blood pressure is lower than what is normal for child's age.

Lowest normal pediatric systolic blood pressure by age:

- Less than one month: > 60 mmHg.
- One month to 1 year: > 70 mmHg.
- Greater than 1 year: 70 + 2 x age in years

- Acetaminophen potentiates the analgesic effect of opioids, and they can be given together.
- Benzodiazepines do not have an analgesic effect. Their anxiolytic effects may potentiate the analgesic effect of opioids but also increase the likelihood of respiratory depression. OLMC consult is required for use of midazolam or along with opioids for pain management.
- Do not give oral medication to patients with abdominal pain or open or obviously angulated fractures.
- Ketorolac should not be used in patients less than 2 or over 80.
- Do not administer ketamine to patients who are pregnant or have non-traumatic chest pain.
- Ketamine should not be given to patients with schizophrenia or history of psychosis due to the potential for exacerbating the mental health condition.

Poisoning & Overdose – 10.140

Treat per Universal Patient Care

*

If systolic BP < 90 mmHg, follow Shock Protocol. Goal is to maintain a mean arterial pressure (MAP) <u>></u> 65 mmHg.

▼

If unknown poison or overdose and the patient has a decreased LOC, treat per Altered Mental Status protocol. Manage airway per the Airway Management protocol. Contact OLMC and/or Poison Center (**1-800-222-1222**) for advice.

↓

Treat specific <u>symptomatic</u> poisoning/overdose patient as outlined below. Strongly consider Haz-Mat Team activation when appropriate.		
OVERDOSE/POISONING	TREATMENT	
Aspirin and/or Acetaminophen	 Activated Charcoal 1 g/kg if < 2 hours since ingestion. Max dose 50 g. If ingestion involves other substances, contact OLMC. Avoid intubation for ASA ODs unless necessary. If intubation becomes necessary, the ventilation goal is to maintain pre-intubation EtCO₂ levels. 	
Beta Blockers	Treat bradycardia/hypotension with push dose epinephrine as bridge until an epinephrine drip at 2 - 10 mcg/min can be started. Titrate to effect.	
Calcium Channel Blocker	Calcium gluconate, 1 - 3 g slow IV/IO over 5 - 10 minutes.	
Carbon Monoxide	 Place all suspected CO poisoning patients on CPAP/BiPAP with high flow O₂. Recommend NRB with nasal cannula if contraindications to or if patient does not tolerate CPAP/BiPAP. Measure CO level with SpCO monitor when possible. All symptomatic patients (e.g., headache, dizziness, nausea) or patients with an SpCO monitor reading ≥ 15% should be transported. Transport patients with severe symptoms (e.g., cardiac ischemia, coma, syncope, seizures, loss of consciousness) to a hyperbaric facility if available, or nearest facility if unavailable. Treat symptoms per appropriate protocol (e.g., 12-lead ECG for suspected cardiac ischemia.) If cyanide poisoning is also suspected, consider obtaining SpCO, if possible, before administration of CYANOKIT[®] since the latter will interfere with the carboxyhemoglobin monitor. SpCO levels may be elevated in smokers. Levels can range from 3 - 10% depending on the number of packs smoked. Pulse oximeter may provide a false reading in patients with elevated SpCO levels. 	

Chlorine Inhalation	 Treat symptomatic patients with: Albuterol- 2.5 mg nebulized. Dexamethasone- 10 mg IV/IO/IM/PO. Sodium bicarbonate 8.4%- 2.5 ml via nebulizer. 	
Cyanide	Hydroxocobalamin (CYANOKIT [®]) 5 g IV/IO over 15 minutes. Repeat once if needed. For cardiac arrest, hydroxocobalamin should be administered as a rapid bolus.	
Hydrofluoric Acid	Dermal: Calcium gluconate 3 g mixed with 5 oz water soluble lubricant and applied to burn.	
Organophosphate	 Prepare to handle copious secretions. In mild to moderate poisonings (e.g., headache, mild bronchorrhea, nausea, vomiting, diarrhea but normal mentation), administer atropine 1 - 2 mg IV/IO/IM every 3 - 5 minutes until symptoms improve. For severe poisoning (e.g., altered mental status, unconsciousness, seizures), administer atropine 3 - 5 mg IV/IO/IM every 3 - 5 minutes until symptoms begin to improve. Treat seizures per seizure protocol. See Haz-Mat Protocol for more specifics of treatment. For large organophosphate poisonings, refer to HazMat protocol. 	
Sodium Channel Blocker (TCA, diphenhydramine, Type 1a and 1c antiarrhythmics)	 If patient exhibits arrhythmias or a widening QRS complex, administer sodium bicarbonate 1 mEq/kg IV/IO. Treat hypotension per Shock protocol. 	
Do not neutralize acids or alkalis.		
If the patient exhibits extrapyramidal symptoms/Dystonia with a history of phenothiazine use, consider diphenhydramine.		

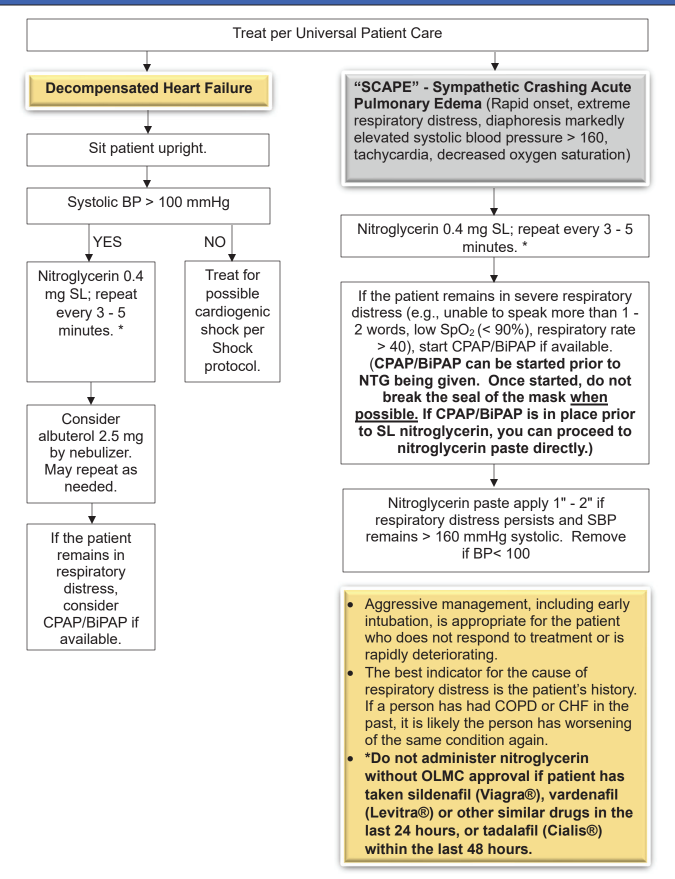
PEDIATRIC PATIENTS:

- Consider possibility of neglect or abuse.
- For organophosphate poisoning, atropine dose is 0.05 mg/kg IV/IO. Contact OLMC for frequency of dosing.
- Activated charcoal dose is 1 g/kg, max of 50 g.
- For children < 1-year, dilute sodium bicarbonate by one-half with normal saline prior to administration.
- Hydroxocobalamin for cyanide poisoning- 70 mg/kg IV/IO to a max of 5 g over 15 minutes. For cardiac arrest, hydroxocobalamin should be administered as a rapid bolus. Contact OLMC for advice regarding second dose.

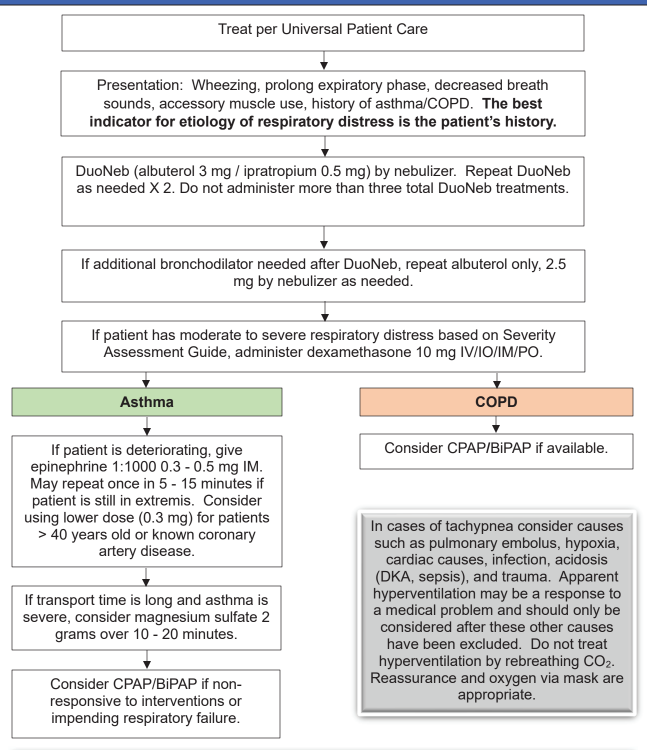
Poisoning & Overdose – 10.140

TOXIDROME TABLE			
Toxidrome	Examples	Clinical Features	Antidotes/Treatment
Sympathomimetic	Cocaine Methamphetamine Ecstasy/MDMA	Agitation Hyperthermia Diaphoresis Dilated pupils HTN Tachycardia	Midazolam or lorazepam (OLMC)
Opioid	Heroin/Fentanyl Hydromorphone Methadone Oxycodone	Depressed mental status Hypoventilation Constricted pupils	Naloxone
Cholinergic (Anti- cholinesterase)	Pesticides • Carbamates • Organophosphates Nerve agents	Muscarinic* Nicotinic** Central***	Atropine Pralidoxime (2-Pam) Midazolam (Hazmat, OLMC)
Sedative-Hypnotic	Barbiturates Benzodiazepines GHB	Depressed mental status Hypotension Hypothermia	Supportive care
Cardiotoxic drugs	Beta-blockers Calcium channel blockers	Bradycardia Conduction issues Hypotension	Epinephrine Calcium (OLMC)
Anticholinergic	Atropine Jimson Weed Scopolamine Diphenhydramine	Delirium Hyperthermia Tachycardia Warm, dry skin	Supportive treatment Physostigmine (ED)
Sodium channel blockade	 Tricyclic antidepressants Antiarrhythmics Type 1A – quinidine, procainamide Type 1C – flecainide, propafenone 	Altered mental status Hypotension Seizures Wide complex tachycardia	Sodium Bicarbonate (OLMC)
Methemoglobinemia (nitrate/nitrite poisoning)	Contaminated well water (nitrates) Inhalation injuries Topical anesthetics (benzocaine, lidocaine) Amyl Nitrites (poppers)	Cyanosis SpO ₂ 75-85% despite supp. O ₂ Headache Weakness Seizures/Coma Dysrhythmias Chocolate brown blood	Supportive Care O₂ administration Methylene blue (ED)
salivation, sweating.	tachycardia, weakness, hy	rdia, bronchospasm, bronchorrhea pertension, hyperglycemia, fascici	

Respiratory Distress- CHF/Pulmonary Edema – 10.160



Respiratory Distress- COPD/Asthma – 10.160

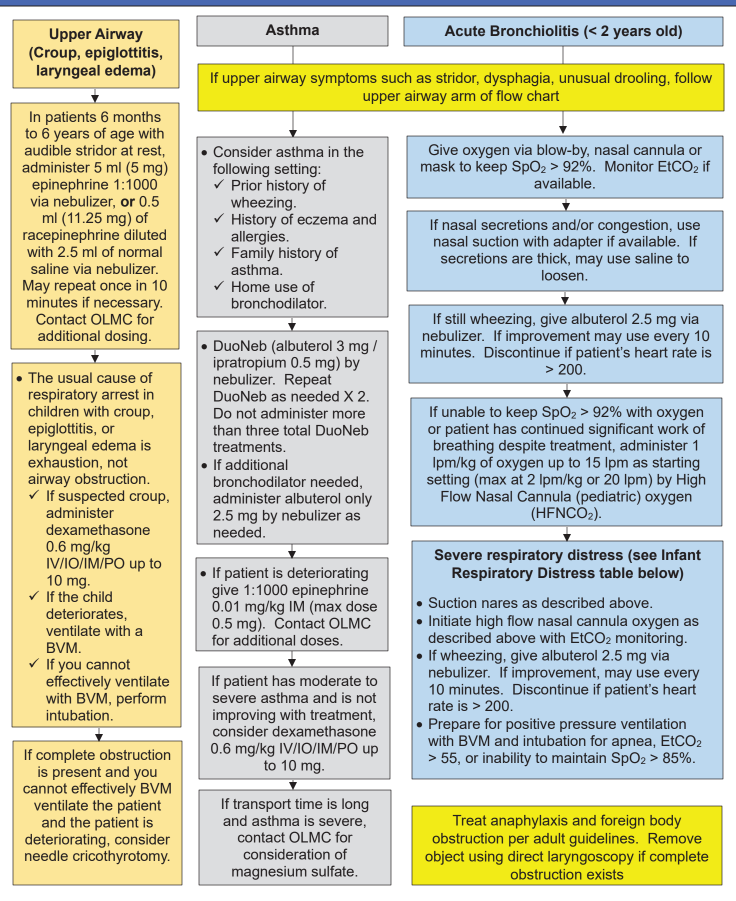


- Aggressive management, including early intubation, is appropriate for the patient who does not respond to treatment or is rapidly deteriorating.
- COPD and asthma patients receiving CPAP/BiPAP need to be monitored closely due to the higher risk of secondary pneumothorax from positive pressure ventilation.

Respiratory Distress- COPD/Asthma – 10.160

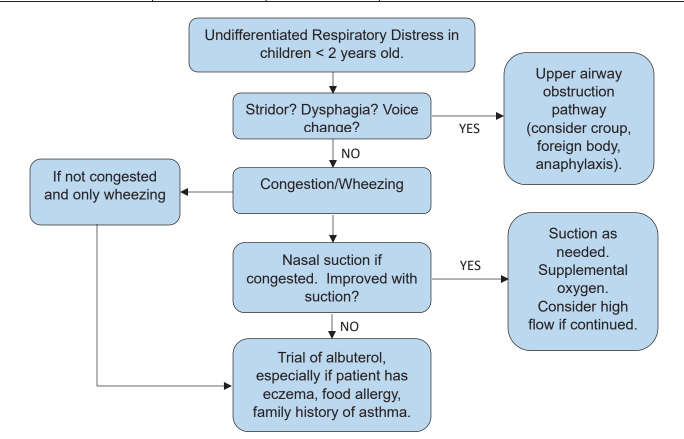
ASTHMA SEVERITY ASSESSMENT GUIDE			
	MILD	MILD MODERATE	
Short of breath	Walking	Talking	At rest
Able to speak	In sentences	In phrases In words	
Heart rate	< 100	100 - 120	> 120
Respiratory rate	Elevated	Elevated	> 30
Lung sounds	End expiratory wheezes	Full expiratory wheezes	Wheezes both phases or absent
Accessory muscle use	Not usually	Common	Usually
Alertness	Possibly agitated	Usually agitated	Usually agitated
EtCO ₂	20 - 30	30 - 40 > 50	

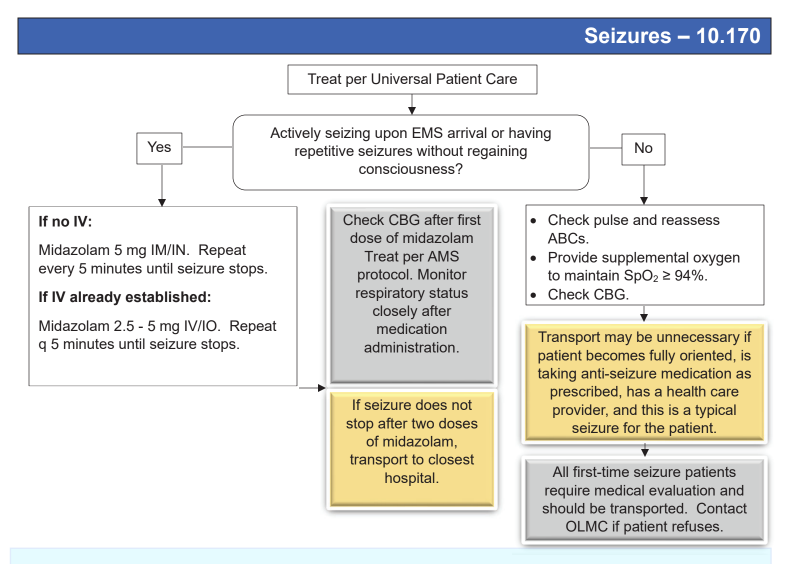
Respiratory Distress- Pediatrics – 10.160



Respiratory Distress- Pediatrics – 10.160

INFANT RESPIRATORY DISTRESS ASSESSMENT GUIDE			
	MILD	MODERATE	SEVERE
Respiratory Rate			
≤ 2 months	≤ 60	61 - 69	≥ 70
2 - 12 months	≤50	51 - 59	≥ 60
1 - 2 years	≤ 40	41 - 44	≥ 45
Retractions	Subcostal or intercostal	2 of: subcostal, intercostal, substernal retractions, OR nasal flaring	3 of: subcostal, intercostal, substernal, suprasternal, supraclavicular retractions, OR nasal flaring, OR head bobbing
Dyspnea	1 of: difficulty feeding, decreased vocalization or agitation	2 of: difficulty feeding, decreased vocalization or agitation	Stops feeding, no vocalization OR drowsy and confused
Auscultation	End-expiratory wheeze only	Expiratory wheeze only	Inspiratory and expiratory wheezing OR diminished breath sounds OR both





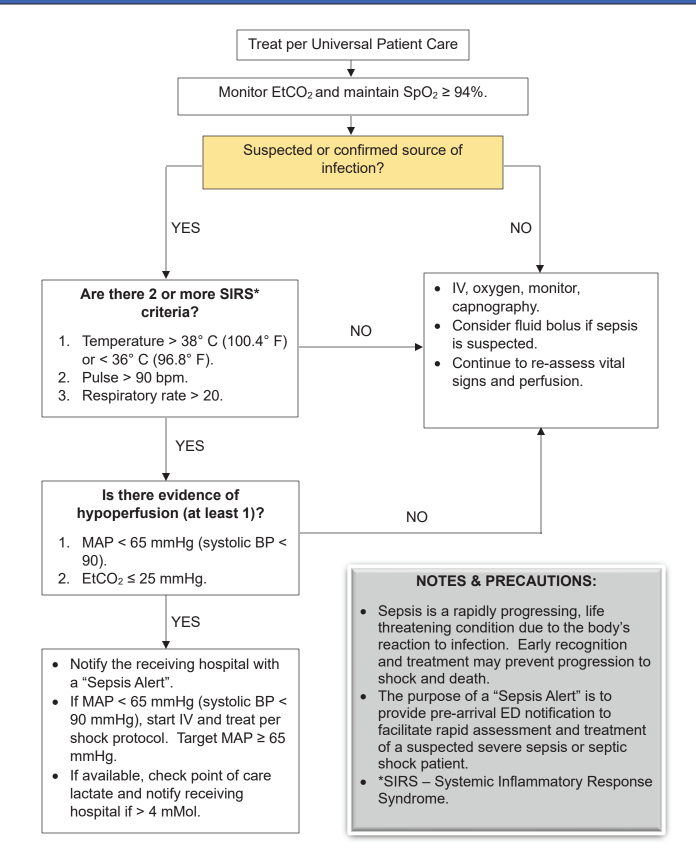
PEDIATRIC PATIENTS:Refer to Pediatric Guide

- If patient is actively seizing upon EMS arrival or having repetitive seizures without regaining consciousness:
 ✓ 0 11 months (16 29", 0 8 kg) or patient is extremely small for age: Follow pediatric guide and administer midazolam 0.3 mg/kg IM/IN to a max of 10 mg. Repeat every 5 minutes until seizure stops.
 ✓ 12 months 13 years old (use reported age; if unknown, measure patient and use corresponding length in inches to determine dose:
- If an IV/IO is already available, follow pediatric guide and administer midazolam 0.1 mg/kg IV/IO to a max of 10 mg. Repeat every 5 minutes until seizure stops.
- If no IV/IO access is available 0.3 mg/kg IM/IN to a max of 10 mg.
- If seizure does not stop after two doses of midazolam transport to closest hospital.
- If on arrival, the patient is not actively seizing (post-ictal), an IV is not required.
- All hypoglycemic or first-time pediatric seizures patients should be transported.
- Febrile seizures are typically found between the ages of 6 months 6 yrs. and are usually brief.

NOTES & PRECAUTIONS:

- Seizures in patients > 50 years of age can be caused by dysrhythmias. Monitor rhythm and treat per appropriate protocol. Remember to check a pulse once a seizure stops.
- The longer status seizure lasts, the more difficult it is to control. Seizures that aren't responsive to midazolam may require alternative antiepileptic agents in a timely manner.
- New onset of seizures in a pregnant patient, especially in the third trimester, may indicate eclampsia. Contact OLMC for consideration of magnesium sulfate. Normal dose is 4 grams IV over 15 20 minutes.

Sepsis – 10.175



Shock - 10.180

Treat per Universal Patient Care and prepare for rapid transport			
Determine type of shock and treat as follows:			
Hypovolemic or Hemorrhagic Shock	Obstructive Shock (Tamponade, Pneumothorax, PE)		
 Control external bleeding with direct pressure, elevation, tourniquet, and/or hemostatic dressing. Administer 500 - 1000 ml fluid challenge to maintain MAP > 65 mmHg (SBP > 90 mmHg). Repeat fluid boluses if continued signs of shock and no pulmonary edema. For shock secondary to trauma or suspect AAA do not over resuscitate. MAP 55 - 65 mmHg (Goal is SBP 70 - 90 mmHg). If <u>hemorrhagic shock</u> with blunt or penetrating trauma and MAP < 50 mmHg (SBP < 70 mmHg), administer 2 grams TXA slow IV/IO push. Contact OLMC for advice. 	 If tension pneumothorax is suspected, decompress per the Tension Pneumothorax Decompression procedure protocol. Administer 500 - 1000 ml fluid challenge to maintain MAP > 65 mmHg (SBP > 90 mmHg). Repeat fluid boluses if continued signs of shock and no pulmonary edema. If <u>not</u> responding to fluid administration begin norepinephrine infusion at 4 mcg/min. If no response, increase every 5 minutes in 4 mcg/min increments to max of 24 mcg/min. Goal is MAP > 65 mmHg (SBP > 90 mmHg). While drip is being set up, consider push dose epinephrine, per epinephrine protocol, for temporary hemodynamic support. Contact OLMC for advice. 		
Cardiogenic Shock (STEMI, cardiomyopathy)	Distributive Shock (septic, neurogenic, anaphylactic) or unknown type of shock		
 If suspected cardiac event, follow Chest Pain protocol. Monitor cardiac rhythm and follow Cardiac Dysrhythmia protocol. Administer 250 - 500 ml fluid challenge to maintain MAP > 65 mmHg (SBP > 90 mmHg). May repeat once if continued signs of shock and no pulmonary edema/volume overload. Max of 1000 ml. If <u>not</u> responding to fluid administration, begin norepinephrine infusion at 4 mcg/min. If no response, increase every 5 minutes in 4 mcg/min increments to max of 24 mcg/min. Goal is MAP > 65 mmHg (SBP > 90 mmHg). While drip is being set up, consider push dose epinephrine, per epinephrine protocol, for temporary hemodynamic support. Contact OLMC for advice. 	 If anaphylaxis is suspected, follow Anaphylaxis and Allergic Reaction protocol. Administer 500 - 1000 ml fluid challenge to maintain MAP > 65 mmHg (SBP > 90 mmHg). Repeat once if continued signs of shock and no pulmonary edema. If not responding to fluid administration, begin norepinephrine infusion at 4 mcg/min. If no response, increase every 5 minutes in 4 mcg/min increments to max of 24 mcg/min. Goal is MAP > 65 mmHg (SBP > 90 mmHg). While drip is being set up, consider push dose epinephrine, per epinephrine protocol, for temporary hemodynamic support. Contact OLMC for advice. 		

Shock – 10.180

PEDIATRIC PATIENTS:

FEDIATRIC FATILITS.	PEDIATRIC PATIENTS:				
Treat per Universal Patient Care and prepare for rapid transport					
Determine type of shock and treat as follows:					
Hypovolemic or Hemorrhagic Sh	ock	Obstructive Shock (Tamponade, Pneumothorax, PE)			
 Control external bleeding with direct p elevation, tourniquet, and/or hemostat Administer 20 ml/kg fluid challenge (1) neonates) to maintain age appropriate Repeat twice if continued signs of sho pulmonary edema to a max of 60 ml/k neonates) Contact OLMC for advice. 	ic dressing. 0 ml/kg in e SBP. ck and no	 If tension pneumothorax is suspected, decompress per the Tension Pneumothorax Decompression procedure protocol. Administer 20 ml/kg fluid challenge (10 ml/kg in neonates) to maintain age appropriate SBP. Repeat twice if continued signs of shock and no pulmonary edema to a max of 60 ml/kg (30 ml/kg in neonates) If <u>not</u> responding to fluid administration begin norepinephrine infusion at 0.1 mcg/kg/min. If no response, in 5 minutes, increase to 0.2 mcg/kg/min. 			
Lowest normal pediatric systolic blood age: Less than one month: > 60 One month to 1 year: > 70 r Greater than 1 year: 70 + 2 years. 	mmHg. nmHg.	 If still no response after 5 more minutes, may increase to 0.4 mcg/kg/min. Goal is age appropriate SBP. While drip is being set up, consider push dose epinephrine, per epinephrine protocol, for temporary hemodynamic support. Contact OLMC for advice. 			
Cardiogenic Shock (STEMI, cardiom	yopathy)	Distributive Shock (septic, neurogenic, anaphylactic) or unknown type of shock			
 If suspected cardiac event, follow Cheprotocol. Monitor cardiac rhythm and follow CarDysrhythmia protocol. Administer 20 ml/kg fluid challenge (1 neonates) to maintain age appropriate Repeat twice if continued signs of shopulmonary edema to a max of 60 ml/k neonates). If blood pressure remains low, begin norepinephrine infusion at 0.1 mcg/kg response in 5 minutes, increase to 0.2 If still no response after 5 more minute increase to 0.4 mcg/kg/min. Goal is a appropriate SBP. While drip is being set up, consider puepinephrine, per epinephrine protocol, temporary hemodynamic support. 	rdiac 0 ml/kg in e SBP. ck and no g (30 ml/kg in /min. If no ex, may ge ush dose	 If anaphylaxis is suspected, follow Anaphylaxis and Allergic Reaction protocol. Administer 20 ml/kg fluid challenge (10 ml/kg in neonates) to maintain age appropriate SBP. Repeat twice if continued signs of shock and no pulmonary edema. If blood pressure remains low, begin norepinephrine infusion at 0.1 mcg/kg/min. If no response in 5 minutes, increase to 0.2 mcg/kg/min. If still no response after 5 more minutes, may increase to 0.4 mcg/kg/min. Goal is age appropriate SBP. While drip is being set up, consider push dose epinephrine, per epinephrine protocol, for temporary hemodynamic support. Contact OLMC for advice. 			

NOTES & PRECAUTIONS:

- Closely monitor patient's respiratory status and vital signs. Avoid fluid overload.
- Mean Arterial Pressure targets:
 - ✓ Uncontrolled traumatic hemorrhagic shock without TBI or suspected AAA, target MAP is 55 65 mmHg (SBP 70 - 90).
 - ✓ Uncontrolled traumatic hemorrhagic shock with TBI or shock from all other causes, target MAP is ≥ 65 mmHg (SBP ≥ 90).
- For patients in shock with known or suspected adrenal insufficiency (AI) consider administration of dexamethasone 10 mg (0.6 mg/kg for pediatric patients) in addition to fluids and/or norepinephrine.
- If an improvised tourniquet is present before medical provider arrival, place a commercial tourniquet per protocol and remove the improvised tourniquet if operationally feasible.

Treat per Universal Patient Care

- Apply cardiac monitor as soon as possible and continuously assess rhythm.
- Place 18g IV or larger in AC when possible.
- Check CBG: If low, treat per Diabetic Emergencies-Hypoglycemia protocol.
- No oxygen if $SpO_2 \ge 94\%$ with good waveform.

Complete the **BEFAST stroke assessment** if last known well time is \leq 24 hours ago.

BEFAST STROKE SCREEN			
Neurological examination	Normal	Abnormal (any positive)	
Balance			
Symptoms:			
Acute loss of balance, coordination, trouble walking			
Test:			
Perform bilateral index finger-to-nose test (FTN test)			
Have the patient walk if normally able to (walk next to them in			
case of gait instability)If patient unable to walk, have the patient sit up (truncal stability)	Normal	Abnormal	
test)			
Positive findings:			
Patient overshoots or undershoots intended target (FTN test)			
Patient falls over to one side (truncal instability)			
Unsteady gait (shuffling, wide based gait, falling to one side) that the patient reports is acutely abnormal			
Eyes			
Symptoms:			
Acute onset of vision loss, double vision, or part of vision loss (visual field cut)			
Test:			
• Ask the patient if they have double vision or loss of vision in one			
or both eyesMake sure the patient can move their eyes all the way from left	Normal	Abnormal	
to right up and down (extraocular movements)			
Positive findings:			
• Trouble seeing out of one or both eyes or acute onset of double vision or visual field cut			
Eyes are deviated together to the left or to the right or are unable to perform full movement			

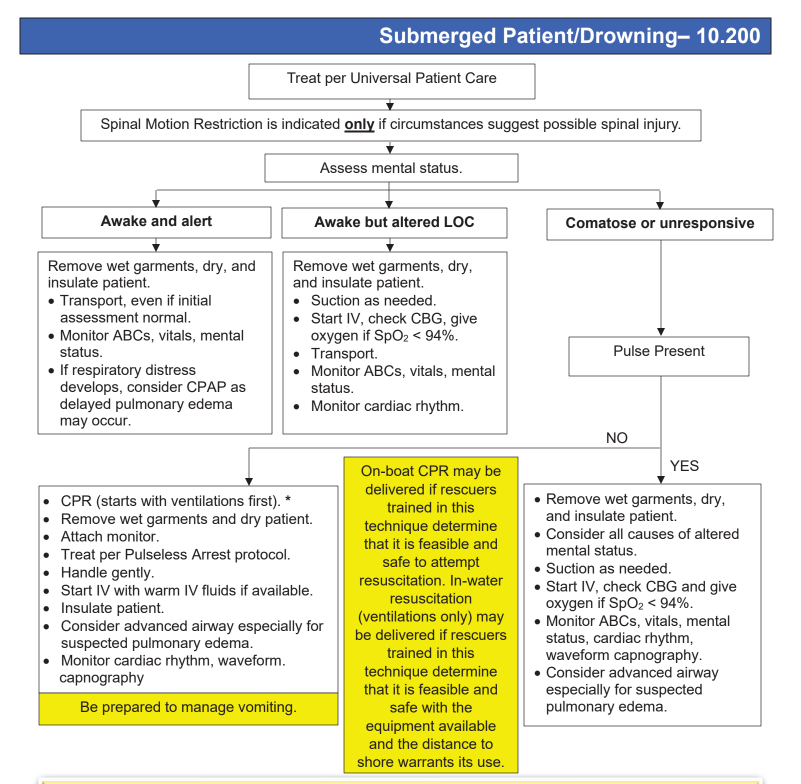
Stroke / CVA – 10.190

Neurological examination	Normal	Abnormal (any	positive)
 Face <u>Symptoms:</u> Acute onset facial droop <u>Test:</u> Ask the patient to smile or show their teeth <u>Positive findings:</u> The patient's face looks uneven, is drooping, or has numbness on one side 	Normal	Right	Left
 Arms/Legs <u>Symptoms:</u> Acute onset numbness or weakness of the <u>arm/leg on one side of the body</u> <u>Test:</u> Ask the patient to raise and extend both arms with their palms up for 10 seconds, then close their eyes Ask the patient to raise one leg at a time for 5 seconds Touch each side of the patient's extremities and ask if they feel each equally <u>Positive finding:</u> One arm or leg drifts downward Unequal extremity sensation 	Normal	Right	Left
 Speech Symptoms: Acute onset slurred speech, trouble speaking, or understanding Test: Ask the patient to repeat the phrase, "The sky is blue." Ask the patient to (1) squeeze AND let go of your hand (2) open AND close their eyes Ask the patient to name common objects (e.g., glove, pen, watch) Positive findings: Slurred speech, trouble finding words, unintelligible words Patient is unable to follow simple commands Patient is unable to recognize common objects 	Normal Abnormal		al
 Time What time was the patient last known well (i.e., last appear normal)? 	L	ast Known Well Tin	ne:

Stroke / CVA – 10.190

If BEFAST is positive (at least 1 of the neurological examination findings are **ABNORMAL**), the patient is considered to have a **POSITIVE** stroke screen. Continue to **Cincinnati – Stroke Triage Assessment Tool (C-STAT)** to screen for a large vessel occlusion (LVO) stroke.

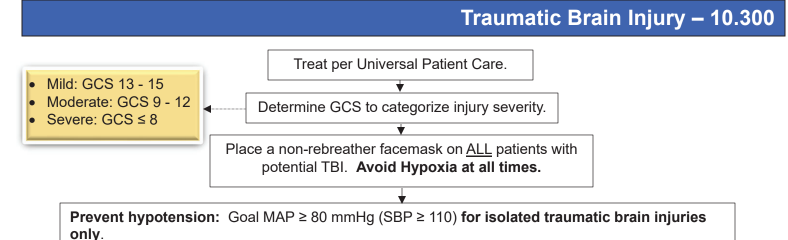
	Ļ	
2 C-STAT – CINCINNATI STROKE TRIAGE ASSESSMENT TOOL		
	Points	
Conjugate Gaze Deviation - Eyes are de	eviated togethe	r to the left or to the right or are unable to
perform full movement.		1
Absent	0	_
Present	2	
Arm Weakness - Cannot hold up one arm		
Absent Present	0	-
	I Swers at least (one of two LOC questions (1) what is your
		least one of two commands (1) squeeze AND
let go of your hand (2) open AND close th		
Absent	0	
Present	1	-
	•	
 If BEFAST and C-STAT positive (≥ 2), t 		
 notify the receiving facility of acute stro If BEFAST positive and C-STAT negative 		
receiving facility of acute stroke alert a		
 If BEFAST is negative, transport to any 		
 Notify the receiving facility if the patient 		
, , , , , ,		
▼ Transport patient with head elevated at least 30 degrees.		
i ransport patien	t with head ele	valed at least 30 degrees.
Document serial neurologic examinations.		
		NOTES & PRECAUTIONS:
Do not treat hypertension or give a		d to the nearest CT conclusion with
All potential stroke patients should	be transported	d to the nearest CT capable hospital.
reatment - Revised 10/28/2024		Stroke / CVA - 10.190



*Do not attempt resuscitation if patient has been submerged for more than 30 minutes, with the following exceptions:

Resuscitation may be initiated if the patient is recovered within 60 minutes if:

- Children < 6 years of age and water temperature at recovery depth of < 40° F.
- Patients who may have been trapped in an underwater air pocket.
- Water temperature at recovery depth is < 40° F and information suggests that patient may have been swimming on the surface for at least 15 minutes prior to becoming submerged.
- Paramedic discretion (contact OLMC).



Initiate a bolus of normal saline or lactated ringers.

• Continue fluid boluses to maintain the MAP ≥ 80 mmHg (SBP ≥ 110 mmHg).

AIRWAY CONSIDERATIONS:

- If patient is unable to maintain airway, consider oral airway (nasal airways should not be used in the presence of significant facial injury or possible basal skull fracture).
- Place an advanced airway (oral endotracheal intubation, supraglottic device, surgical airway) if BVM ventilation ineffective in maintaining oxygenation or if airway is continually compromised. Nasal intubation should not be attempted.
- If the patient has an airway placed (oral or advanced), carefully manage ventilations to minimize hyperventilation.
 - ✓ Monitor EtCO₂ with goal of EtCO₂ of 40 mmHg.
 - ✓ If available, use a pressure-controlled bag (PCB) and ventilation rate timer (VRT).
 - \checkmark If a transport ventilator is available, begin with the following settings:
 - Tidal volume of 7ml/kg,
 - > Rate of 10 BPM. Adjust rate to keep EtCO₂ within target range.

If there are signs of herniation, then $\underline{\text{MILD}}$ hyperventilation to an EtCO₂ of 35 mmHg may be performed. Signs of herniation include:

- Blown pupil
- Posturing

For moderate to severe blunt or penetrating head trauma: **<u>If available</u>**, administer 2 grams Tranexamic acid (TXA) slow IV/IO push if **<u>both</u>** of the following indications are met:

• Age \geq 15 (or \geq 50 kg if age unknown).

• GCS between 3 and 12, with a reactive pupil.

Contraindications to TXA:

- Time of head injury > 2 hours.
- GCS of 3 with no reactive pupil.
- EMS chest compressions at any time (manual or mechanical).
- Patients with a clinical concern for: Epilepsy, seizures, MI, stroke, PE, DVT, renal failure dialysis.
- Known or suspected pregnancy.
- Drowning, hanging, or burns > 20% TBSA.

Consider and treat reversible causes of AMS including hypoxia, hypoglycemia, and overdose.

PEDIATRIC PATIENTS (follow adult flow chart with the following considerations):

- Manage blood pressure. Avoid hypotension.
 - ✓ Initiate a 20 ml/kg bolus of normal saline or lactated ringers.
 - ✓ Continue fluid boluses to maintain SBP goals:
 - Infants/children age < 10: 70 mmHg + (age X 2).</p>
 - > Children age \geq 10: 110 mmHg (same as adults).
- Pediatric ventilatory rates:
 - ✓ Infants: (age 0 24 months): 25 breaths per minute (bpm);
 - ✓ Children: (age 2 14): 20 bpm;
 - ✓ 15 years: 10 bpm (same as adults).

NOTES & PRECAUTIONS:

- The main goal is to address the three H's that increase mortality with isolated TBI:
 ✓ Avoid Hypoxia.
 - ✓ Avoid **H**yperventilation.
 - ✓ Avoid **H**ypotension.
- A single episode of hypoxia is independently associated with DOUBLING of the mortality rate.
- Hyperventilation is independently associated with a mortality rate that is between TWO and SIX times higher. Inadvertent hyperventilation happens reliably if not meticulously prevented by proper external means.
- A single episode of hypotension is independently associated with DOUBLING of the mortality rate and persistent hypotension is independently associated with a mortality rate that is eight times higher.

Medications

SUPPLIED:

Acetaminophen liquid 160 mg/5 mL Acetaminophen 325 mg and 500 mg tablets, capsules, gel, suppositories

PHARMACOLCOGY AND ACTIONS:

Acetaminophen (paracetamol) targets the cyclooxygenase enzymes that produce prostaglandins responsible for pain and fever. It has little anti-inflammatory effect. It is metabolized into toxic and non-toxic products in the liver by:

- <u>Glucuronidation</u> (45-55%)
- Sulfate conjugation (20–30%)
- N-hydroxylation and dehydration, then GSH conjugation (less than 15%)

All three pathways yield final products that are non-toxic. In the third pathway, however, the intermediate product NAPQI is toxic. At usual doses, NAPQI is quickly detoxified by conjugation with glutathione. In overdose, glutathione is used up and the toxic metabolite can cause potentially fatal <u>liver damage</u>. It is metabolized by the liver and is <u>hepatotoxic</u>. Toxicity is multiplied when combined with alcoholic drinks, and very likely in <u>chronic</u> <u>alcoholics</u> or patients with liver damage.

INIDCATIONS:

- A. Mild to moderate pain.
- B. Fever.

CONTRAINDICATIONS

- A. Known liver disease.
- B. Current alcohol abuse.
- C. Acute intoxication.
- D. Has taken acetaminophen in last 4 hours.

ADULT & PEDIATRIC DOSING:

Acetaminophen 15 mg/kg PO to maximum of 1000 mg

Approximate dosing using 160 mg/5 mL liquid

Weight	Dose	Volume
11 lbs/5kg	80 mg	2.5 mL
22 lbs/10 kg	160 mg	5 mL
45 lbs/20 kg	320 mg	10 mL
66 lbs/30 kg	480 mg	15 mL
88 lbs/40 kg	640 mg	20 mL
110 lbs/50 kg	800 mg	25 mL
130 lbs/60 kg	960 mg	30 mL

- A. Aspirin and acetaminophen with time of ingestion > two hours.
- B. All other poisons or ingestions regardless of time from ingestion.

SUPPLIED: 25 grams / 120 ml bottle.

PHARMACOLOGY AND ACTIONS:

Activated charcoal adsorbs toxic substances ingested and inhibits GI adsorption by forming an effective barrier between the particulate material and the gastrointestinal mucosa. The effect is greatest if used within one hour of ingestion.

INDICATIONS:

Management of poisoning or overdose of many substances.

CONTRAINDICATIONS:

- A. Patients with altered mental status or the inability to maintain their own airway.
- B. Patients who have aspirated or with a potential for aspiration.

PRECAUTIONS:

- A. Activated charcoal may be ineffective in some ingestions.
- B. Milk, ice cream and other dairy products will decrease the adsorption capacity substantially.

SIDE EFFECTS AND NOTES:

May cause nausea, vomiting, and constipation.

ADULT DOSING:

Poisoning & overdose -1 gram / kg PO or OG to a max of 50 grams.

PEDIATRIC DOSING:

Same as adult.

SUPPLIED: 6 mg / 2 ml and 12 mg / 4 ml pre-filled syringes

PHARMACOLOGY AND ACTIONS:

Adenosine is a naturally occurring nucleoside that has the ability to slow conduction through the AV node. Since most cases of PSVT involve AV nodal re-entry, adenosine is capable of interrupting the AV nodal circuit and stopping the tachycardia, restoring normal sinus rhythm. It is eliminated from the circulation rapidly and has a half-life in the blood of less than ten seconds.

INDICATIONS:

To convert PSVT to a normal sinus rhythm, including PSVT that is associated with accessory bypass tracts (e.g. Wolff-Parkinson-White Syndrome).

CONTRAINDICATIONS:

- A. Second- or third-degree heart block.
- B. Sick Sinus Syndrome.
- C. Known hypersensitivity.
- D. Atrial fibrillation.

PRECAUTIONS:

- A. When doses larger than 12 mg are given by injection, there may be a decrease in blood pressure secondary to a decrease in vascular resistance.
- B. The effects of adenosine are antagonized by methylxanthines such as theophylline and caffeine. Larger doses of adenosine may be required.
- C. Adenosine effects are potentiated by dipyridamole (Persantine) resulting in prolonged asystole.
- D. In the presence of carbamazepine (Tegretol), high degree heart block may occur.
- E. Adenosine is not effective in converting atrial fibrillation, atrial flutter or ventricular tachycardia.
- F. All doses of adenosine should be reduced to one-half (50%) in the following clinical settings:
 - 1. History of cardiac transplantation.
 - 2. Patients who are on carbamazepine (Tegretol) and dipyridamole (Persantine).
 - 3. Administration through any central line.
- G. Adenosine should be used with caution in patients with asthma as it may cause a reactive airway response in some cases.

SIDE EFFECTS AND NOTES:

May cause facial flushing, shortness of breath, chest pressure, nausea, headache, and lightheadedness.

ADULT DOSING:

6 mg rapid IV. May repeat with 12 mg IV x 2 if patient fails to convert after 6 mg dose. Use a large proximal IV site with fluid bolus flush.

PEDIATRIC DOSING:

0.1 mg/kg rapid IV. May repeat with 0.2 mg/kg once if patient fails to convert after first dose. Use a large proximal IV site with fluid bolus flush.

SUPPLIED: 2.5 mg / 3 ml vial individually or 3 mg packaged with 0.5 mg ipratropium (Duo-Neb).

PHARMACOLOGY AND ACTIONS:

Albuterol is a potent, relatively selective beta-2 adrenergic bronchodilator and is associated with relaxation of bronchial smooth muscle and inhibition of release of mediators of immediate sensitivity from cells, especially MAST cells. The onset of improvement in pulmonary function is within 2 - 15 minutes after the initiation of treatment and the duration of action is from 4 - 6 hours. Albuterol has occasional beta-1 overlap with clinically significant cardiac effects.

INDICATIONS:

- A. To treat bronchial asthma and reversible bronchial spasm that occurs with chronic obstructive pulmonary disease.
- B. To treat hyperkalemia.
- C. Chlorine Inhalation.

CONTRAINDICATIONS: None in the prehospital setting.

PRECAUTIONS:

- A. The patient's rhythm should be observed for arrhythmias. Stop treatment if frequent PVC's develop or any tachyarrhythmias other than sinus tachycardia appears or if heart rate increases by more than 20 beats/minute.
- B. Paradoxical bronchospasm may occur with excessive administration.

SIDE EFFECTS AND NOTES:

Clinically significant arrhythmias may occur, especially in patients with underlying cardiovascular disorders such as coronary insufficiency and hypertension.

ADULT DOSING:

Respiratory distress -

3.0 ml DuoNeb (3.0 mg albuterol/0.5 mg ipratropium) via nebulizer. Repeat DuoNeb as needed X 2. Do not administer more than three total treatments. If no response to DuoNeb, continue with Albuterol only at 2.5 mg via nebulizer. May repeat as needed.

Hyperkalemia (including secondary to crush injury) -

10 mg via nebulizer.

Chlorine Inhalation-

2.5 mg via nebulizer.

PEDIATRIC DOSING:

Same as adult except in hyperkalemia.

Hyperkalemia-

< 25 kg, 2.5 mg via nebulizer. 25 - 50 kg, 5.0 mg via nebulizer. > 50 kg, 10 mg via nebulizer. **OLMC REQUIRED:** See contraindications.

SUPPLIED: 150 mg / 3 ml pre-filled syringe or vial or 450 mg / 9 ml multiuse vial

PHARMACOLOGY AND ACTIONS:

Amiodarone depresses automaticity of the SA node. It slows conduction and increases refractoriness of the AV node. Amiodarone increases atrial and ventricular refractory period and prolongs the QTc interval. When given IV it is rapidly distributed. No dosage adjustments are needed for patients with renal, liver, heart failure, or advanced age.

INDICATIONS:

- A. Ventricular fibrillation.
- B. Ventricular tachycardia with pulses.
- C. Post resuscitation anti-dysrhythmic.
- D. PVCs in the setting of an acute ischemic event.

CONTRAINDICATIONS:

- A. None in cardiac arrest.
- B. Do not use in perfusing patients in the following situations without OLMC approval:
 - 1. Systolic BP is less than 90 mmHg.
 - 2. Heart rate is less than 50 beats per minute.
 - 3. Periods of sinus arrest are present.
 - 4. Second or third-degree heart blocks are present.

PRECAUTIONS:

- A. In high concentrations (> 3 mg/ml), amiodarone can cause phlebitis. Infusion concentrations should not exceed 2 mg/ml.
- B. Amiodarone will precipitate if administered in the same IV line as sodium bicarbonate or heparin.

SIDE EFFECTS AND NOTES:

- A. In perfusing patients, amiodarone may cause hypotension, prolonged QTc interval, proarrhythmic effects (Torsades and ventricular fibrillation), severe bradycardia, and atrioventricular block.
- B. Non-cardiac toxicities are usually related to chronic administration and include pulmonary infiltrates, hepatic and/or thyroid dysfunction, and peripheral neuropathy.

ADULT DOSING:

V Fib, pulseless V Tach - 300 mg IV/IO. May repeat once with 150 mg.

Unstable V Tach with a pulse (After unsuccessful cardioversion X 2) - 150 mg IV/IO slow IV push over 3 minutes.

Stable V Tach with a pulse - 150 mg IV/IO. Mix with 100 ml of NS (in Buretrol or 100 ml bag) and infuse over 10 minutes via drip or pump. May repeat once if no conversion.

Post resuscitation from VF/pVT - 150 mg IV/IO. Mix with 100 ml of NS (in Buretrol or 100 ml bag) and infuse over 10 minutes via drip or pump. (If amiodarone was last anti-dysrhythmic given prior to ROSC, wait 30 minutes after ROSC to re-dose). Max total arrest/post-ROSC dose is 450 mg. **PVCs (In the setting of an acute ischemic event) –** 150 mg IV/IO over 10 minutes.

PEDIATRIC DOSING:

V Fib, pulseless V Tach - 5 mg/kg IV/IO. May repeat once with 2.5 mg/kg. Unstable V Tach with a pulse (After unsuccessful cardioversion X 2) - 2.5 mg/kg IV/IO slow IV push over 3 minutes.

SUPPLIED: 81 mg chewable tablets (Children's aspirin)

PHARMACOLOGY AND ACTIONS:

Aspirin inhibits prostaglandins and disrupts platelet function for the life of the platelet. It is also a mild analgesic and anti-inflammatory agent.

INDICATIONS:

In unstable angina and acute myocardial infarction, aspirin has been shown to lower mortality and is indicated in patients with suspected ischemic chest pain.

CONTRAINDICATIONS:

- A. Allergy to aspirin or aspirin induced asthma.
- B. History of bleeding disorder (i.e. hemophilia).
- C. Current ulcer or GI bleeding.
- D. Suspected aortic dissection.

SIDE EFFECTS AND NOTES:

- A. High doses of aspirin can cause ringing in the ears.
- B. May cause heartburn, nausea, and vomiting.

ADULT DOSING:

Chest pain (acute myocardial infarction) - 324 mg orally.

PEDIATRIC DOSING: Not indicated for pediatric patients

SUPPLIED: 1 mg / 10 ml pre-filled syringe, 2 mg / 0.7 ml autoinjector, 8 mg / 20 ml vial

PHARMACOLOGY AND ACTIONS:

Atropine is a muscarinic-cholinergic blocking agent. As such, it has the following effects:

- A. Increases heart rate (by blocking vagal influences).
- B. Increases conduction through the AV node.
- C. Reduces motility and tone of the GI tract.
- D. Reduces action and tone of the urinary bladder (may cause urinary retention).
- E. Dilates pupils.

INDICATIONS:

- A. To increase the heart rate in bradycardia or pacemaker failure.
- B. To improve conduction in second- and third-degree heart block.
- C. As an antidote for some insecticide exposures (e.g. anti-cholinesterase, organophosphates) and nerve gases.
- D. To counteract excessive vagal influences causing some brady-systolic and asystole arrests.
- E. For bradycardia not due to hypoxia when using succinylcholine.

CONTRAINDICATIONS:

- A. Atrial fibrillation and atrial flutter because increased conduction may speed ventricular rate excessively.
- B. Not used in neonatal resuscitation.

PRECAUTIONS:

Bradycardia in the setting of an acute myocardial infarction is common and probably beneficial. Do not treat unless there are signs of poor perfusion (low blood pressure, mental confusion).

SIDE EFFECTS AND NOTES:

- A. Atropine blocks cholinergic (vagal) influences already present. If there is little cholinergic stimulation present, effects will be minimal.
- B. Remember in cardiac arrest situations, atropine dilates pupils.

ADULT DOSING:

Bradycardia (cardiac) -

1.0 mg IV/IO. May repeat every 3-5 minutes to max of 3 mg.

Bradycardia secondary to RSI/DSI -

0.5 mg IV/IO.

Organophosphate poisoning -

For mild to moderate poisoning (Headache, mild bronchorrhea, nausea, vomiting, diarrhea but normal mentation): 1-2 mg IV/IO/IM every 3-5 minutes until symptoms improve (e.g., decreased secretions).

For severe poisoning with unconsciousness (Altered mental status, unconsciousness, seizures): 3 - 5 mg IV/IO/IM every 3-5 minutes until symptoms improve (e.g., decreased secretions, ease of ventilation).

PEDIATRIC DOSING:

Bradycardia secondary to RSI/DSI -

0.02 mg/kg IV/IO. Minimum dose 0.1 mg. Do not exceed adult dose.

Organophosphate poisoning -

0.05 mg/kg IV/IO/IM. Contact OLMC for frequency of dosing.

SUPPLIED: 1 gram / 10 ml vial or 5 gram / 50 ml vial (vial size and concentration may vary depending on availability).

PHARMACOLOGY AND ACTIONS:

Calcium is the most common cation in the human body. The majority of the body stores of calcium are located in bone. It plays an important role in many physiologic functions and is essential for proper nerve and muscle function.

INDICATIONS:

- A. Suspected calcium channel blocker overdose.
- B. Hyperkalemia.
- C. Cardiac arrest from suspected hyperkalemia.
- D. Dermal hydrofluoric acid burn.

CONTRAINDICATIONS:

- A. Hypercalcemia and hypercalciuria (hyperthyroidism, Vitamin D overdose, bone metastases).
- B. Patients on digoxin.

PRECAUTIONS:

- A. Extravasation of calcium salts will cause necrosis of tissue. The IV should be secured and free blood return into the syringe should be checked 2-3 times during administration. If extravasation does occur, immediately stop administration.
- B. Administer slowly (no faster than 2 ml/min) and stop if patient complains of distress. Inject using a small needle in a large vein.
- C. Calcium gluconate will precipitate if mixed with sodium bicarbonate. Flush catheter completely before administering one medication after another.

SIDE EFFECTS AND NOTES:

- A. Rapid injection of calcium gluconate may cause vasodilatation, decreased blood pressure, bradycardia, cardiac arrhythmias, syncope, and cardiac arrest.
- B. One 10 ml vial of calcium gluconate 10% contains 1 gram of calcium gluconate salt (= 93 mg elemental calcium or 4.65 mEq calcium or 2.3 mmol calcium).

ADULT DOSING:

Hyperkalemia -

1-3 grams slow IV/IO over 5 – 10 minutes. Use a proximal port. Calcium channel blocker overdose (symptomatic)-

1-3 grams slow IV/IO over 5 – 10 minutes. Use a proximal port.

Cardiac arrest -

3 gram IV/IO push.

Dermal hydrofluoric acid burn-

3 grams mixed with 5 oz. water soluble lubricant applied directly to burn.

PEDIATRIC DOSING:

Hyperkalemia, calcium channel blocker overdose -

0.6 ml/kg slow IV/IO over 5 – 10 minutes. Use a proximal port. Max dose 10 ml.

SUPPLIED: 10 mg / 1 ml vial

PHARMACOLOGY AND ACTIONS:

Dexamethasone is a synthetic steroid that suppresses acute and chronic inflammation. In addition, it potentiates vascular smooth muscle relaxation by beta-adrenergic agonists and may alter airway hyperactivity.

INDICATIONS:

- A. Moderate to severe asthma/COPD.
- B. Severe allergic reaction.
- C. Croup.
- D. Chlorine Inhalation.

CONTRAINDICATIONS:

Do not use in patients with known hypersensitivity to corticosteroids.

PRECAUTIONS:

May cause hypertension and hyperglycemia.

SIDE EFFECTS AND NOTES:

May cause nausea, vomiting, headache, or dizziness.

ADULT DOSING:

Respiratory distress, severe allergic reaction, anaphylaxis -

10 mg IV/IO/IM/PO. Flavoring may be used if available for oral dosing. **Chlorine Inhalation-**10 mg IV/IO/IM/PO.

PEDIATRIC DOSING:

Respiratory distress, severe allergic reaction, anaphylaxis, croup -

0.6 mg/kg IV/IO/IM/PO to a max of 10 mg. Flavoring may be used if available for oral dosing.

SUPPLIED: 25 grams/50 ml pre-filled syringe 50%. 25 grams/250 ml bag 10%

PHARMACOLOGY AND ACTIONS:

Glucose is the body's basic fuel. It produces most of the body's quick energy. Its use is regulated by insulin which stimulates storage of excess glucose outside the bloodstream, and glucagon, which mobilizes stored glucose into the bloodstream.

INDICATIONS:

- A. Hypoglycemia.
- B. Unconscious patient when history is unobtainable.

CONTRAINDICATIONS: None

PRECAUTIONS:

- A. Extravasation of dextrose may cause necrosis of tissue and the patency of the IV should be secured during administration. If extravasation does occur, immediately stop administration.
- B. Report any extravasation to receiving hospital personnel and document on the Prehospital Care Report.

SIDE EFFECTS AND NOTES:

Hyperglycemia may complicate or worsen a number of medical conditions (e.g. myocardial infarction and stroke). Dextrose should be given whenever hypoglycemia is documented by blood glucose meters. If these findings are not available, the EMT should use judgement based on signs and history.

ADULT DOSING:

Hypoglycemia/Altered mental status -

10 - 25 grams slow IV/IO.

PEDIATRIC DOSING -

For infants < 10 kg (birth to 1 year) with CBG < 40 mg% and children 10 kg - 35 kg with CBG < 60 mg% give:

Dextrose 10% - 5 ml/kg IV by infusion to a maximum dose of 250 ml.

Dextrose 12.5% - 4 ml/kg by infusion to a maximum dose of 200 ml. (*if diluting D50*)

SUPPLIED: 50 mg/ml vial

PHARMACOLOGY AND ACTIONS:

Diphenhydramine is an antihistamine which blocks the action of histamines released from cells during an allergic reaction. It has direct CNS effects, which may be a stimulant, or more commonly a depressant, depending on individual variation. Diphenhydramine also has an anticholinergic and antiparkinsonian effect which is used to treat acute dystonic reactions to antipsychotic drugs (e.g. Haldol[®], Thorazine[®], Compazine[®], Inapsine[®]). These reactions include oculogyric crisis, acute torticollis, and facial grimacing.

INDICATIONS:

- A. The second-line drug in anaphylaxis and severe allergic reactions (after epinephrine).
- B. To counteract acute dystonic and dysphoric reactions to anti-psychotic drugs.

CONTRAINDICATIONS: None

PRECAUTIONS:

- A. May have an additive effect with alcohol or other CNS depressants.
- B. Although useful in acute dystonic reactions, it is not an antidote for anti-psychotic toxicity or overdose.
- C. May cause hypotension when given IV.

SIDE EFFECTS AND NOTES:

Diphenhydramine is rarely necessary in the field. It is not the first-line drug for allergic reactions but may be useful for long transports.

ADULT DOSING:

Anaphylaxis, extrapyramidal symptoms -

1 mg/kg IV/IM to a max of 50 mg.

PEDIATRIC DOSING:

Anaphylaxis, extrapyramidal symptoms -

1 mg/kg IV/IM to a max of 50 mg.

SUPPLIED: 5 mg / ml in 5 ml vial

PHARMACOLOGY AND ACTIONS:

Diltiazem is a calcium channel blocker. It slows conduction through the sinoatrial and AV node; thus, slows ventricular response to the stimuli of rapid atrial fibrillation and atrial flutter. IV diltiazem is used primarily for ventricular rate control in atrial fibrillation and atrial flutter. Conversion to normal sinus rhythm often occurs.

INDICATIONS:

As a secondary medication to adenosine in the setting of SVT if the patient has a contraindication to adenosine use, or the patient wishes not to receive adenosine based on past experience.

CONTRAINDICATIONS:

- A. Hypotension (systolic BP < 120)
- B. Wide complex tachycardia of uncertain origin
- C. WPW
- D. Left ventricular dysfunction/heart failure
- E. Sick Sinus Syndrome
- F. AV block without pacemaker
- G. Patient already taking a beta blocker

PRECAUTIONS:

- A. Slow administration is required to avoid hypotension.
- B. Dosing should be reduced by one half in setting of impaired hepatic or renal functions, the elderly and debilitated patients.
- C. Monitor for cardiac dysrhythmias
- D. Monitor for hyperthermia
- E. Be prepared to treat seizures

SIDE EFFECTS AND NOTES:

May produce hypotension, bradycardia, and decreased left ventricular activity

ADULT DOSING:

2.5 mg slow IV push over 1 min. May repeat up to 25 mg in 1-minute increments Or

2.5 mg/min IV infusion to a max of 25 mg

PEDIATRIC DOSING:

Not indicated for pediatric patients

OLMC REQUIRED: Patients < 12 years old

SUPPLIED: 5 mg / 2 ml vial and ampule

PHARMACOLOGY AND ACTIONS:

Droperidol is an antipsychotic that may be used as a chemical restraint by producing marked sedation and allaying apprehension. It also provides a state of mental detachment and indifference while maintaining a state of reflex alertness. Droperidol may potentiate the effects of other CNS depressants. It also causes mild alpha-adrenergic blockade which can lead to peripheral vasodilatation and hypotension, as well as a reduction of the pressor effect of catecholamines. It is also a very effective anti-emetic. Onset of action is from 3-10 minutes following administration and peak effect may not be apparent for up to 30 minutes. Duration is generally 2-4 hours.

INDICATIONS:

- A. Sedation of combative patients to facilitate restraint.
- B. Nausea and vomiting refractory to ondansetron.
- C. Vertigo

CONTRAINDICATIONS:

Unless directed by OLMC, do not administer droperidol in the following situations:

- A. Systolic BP < 90.
- B. Known allergy or prior reaction to droperidol.
- C. Pregnancy.
- D. Patients < 14 years old

PRECAUTIONS:

- A. Use caution when administering droperidol to patients who have taken other CNS depressant drugs (barbiturates, benzodiazepines, alcohol)
- B. Droperidol may induce Torsades De Pointes. <u>Continuously monitor the patient's ECG</u> <u>Q-T interval following use.</u>
- C. Use with caution in patients with a seizure disorder or a condition that causes seizures; droperidol and haloperidol are known to lower the seizure threshold. Consider use of midazolam instead.

SIDE EFFECTS AND NOTES:

- A. The most common side effects are hypotension and tachycardia which usually respond to a fluid bolus.
- B. Akathisia (restlessness) and dystonic reactions have been reported following administration. These symptoms can be treated with the administration of diphenhydramine.

ADULT DOSING:

Nausea & vomiting unresponsive to ondansetron -

1.25 mg IV 0.50 ml based on 5 mg/ 2 ml

Patient restraint -

2.5 mg IV/IO or 5 mg IM. May repeat once in 10 minutes.

For immediate threat (RASS +3 or +4, see patient restraint protocol 30.120): 2.5 - 5 mg IV/IO or 5 - 10 mg IM in addition to midazolam.

PEDIATRIC DOSING: Contact OLMC for patients < 14 years old.

SUPPLIED: 1:10,000 – 1 mg / 10 ml pre-filled syringe; 1:1000 – 30 mg / 30 ml vial or 1 mg/ml vials; racepinephrine 11.25 mg / 0.5 ml

PHARMACOLOGY AND ACTIONS:

Epinephrine is a catecholamine with both alpha and beta effects. In general, the following cardiovascular responses can be expected: increased heart rate, increased myocardial contractile force, increased systemic vascular resistance, increased arterial blood pressure, increased myocardial oxygen consumption, and increased automaticity. Epinephrine is also a potent bronchodilator.

INDICATIONS:

Epinephrine is indicated in the following situations: Ventricular fibrillation, asystole, pulseless electrical activity, symptomatic bradycardia, anaphylaxis, severe asthma, and nebulized in suspected croup (audible stridor at rest in children 6 months to 6 years)

CONTRAINDICATIONS: None

PRECAUTIONS:

- A. Epinephrine increases cardiac workload and can precipitate angina, MI, or major dysrhythmias in individuals with ischemic heart disease.
- B. Wheezing in an elderly person is pulmonary edema or pulmonary embolus until proved otherwise.

SIDE EFFECTS AND NOTES:

- A. May cause anxiety, tremor, or headache.
- B. Cardiac side effects include tachycardia, PVC's, angina and hypertension.

ADULT DOSING:

V Fib, Pulseless V-Tach, asystole, PEA -

1 mg 1:10,000 IV/IO every 3-5 minutes.

Asthma -

0.3 mg - 0.5 mg 1:1000 IM. May repeat once if patient is still in extremis. (Consider using lower doses (0.1 - 0.3 mg) for patients > 40 years old or known coronary artery disease).

Anaphylaxis -

- 1:1000, 0.3 0.5 mg IM. Repeat once in 5-15 minutes if patient is still in extremis. <u>Or, if IV established</u>,
- 1:10,000, 0.1 mg boluses IV/IO every 5 min titrated to effect. Max dose 0.5 mg. <u>OR</u>.
- Infusion IV at 2 mcg/min (2 mcg/ml) titrated to effect. (See drip preparation below)

Symptomatic Bradycardia -

Infusion at 2 mcg/min and increase as needed to a maximum of 10 mcg/min titrating to effect. (See drip preparation below)

Push Dose Epinephrine

INDICATIONS

- **A.** Severe shock (MAP < 50 mmHg or SBP < 60 mmHg) not responsive to fluids.
- **B.** A bridge to drip vasopressors.
- C. Short-lived hypotension (e.g., post-intubation or during sedation).

ONSET

• 1 minute

DURATION

• 5 - 10 minutes

MIXING INSTRUCTIONS:

- A. 10 ml syringe with 9 ml of normal saline.
- **B.** In this syringe, draw up 1 ml of epinephrine 1:10,000 (amp contains 100 mcg/ml epinephrine).
- C. Result is 10 ml of epinephrine with 10 mcg/ml (or 100mcg per syringe).

DOSE:

Adult Dosing: 10 - 20 mcg (1 - 2 ml) IV/IO every 1 - 5 minutes. Pediatric Dosing: 1 mcg/kg every 1 - 5 minutes up to 10 mcg.

Epinephrine Drip Preparation

Mix 1 mg of 1:1000 epinephrine in 500 ml of NS or LR (2 mcg/ml), deliver by micro-drip or infusion pump.

PEDIATRIC DOSING:

V Fib, Pulseless V-Tach, asystole, PEA -

0.01 mg/kg 1:10,000 IV/IO.

Symptomatic Bradycardia (cardiac) -

0.01 mg/kg 1:10,000 IV/IO. Repeat every 3 - 5 minutes.

Asthma -

0.01 mg/kg 1:1000 IM (max dose 0.5 mg). Contact OLMC for additional doses. **Anaphylaxis** -

- Epinephrine 1:1000, 0.01 mg/kg IM to a max of 0.5 mg. Repeat once in 5 -15 minutes if patient is still in extremis. <u>OR, if IV established</u>,
- Epinephrine 1:10,000, 0.01 mg/kg (max 0.1 mg) IV/IO boluses every 3 5 min titrated to effect. Max dose 0.5 mg. OR
- Epinephrine infusion IV at 0.01 mcg/kg/min (2 mcg/ml) titrated to effect.

Respiratory Distress with suspected croup (audible stridor at rest in patients 6 months to 6 years old)-

• See Racepinephrine dosing box below

Racepinephrine (Racemic Epinephrine) - Pediatric use only in suspected croup.

PHARMACOLOGY AND ACTIONS:

Racemic epinephrine is a mixture consisting of d-Epinephrine and I-Epinephrine enantiomers. Epinephrine causes smooth muscle relaxation on various tissues, including bronchial smooth muscles. It also results in vasoconstriction of airway soft tissues when nebulized.

CONTRAINDICATIONS:

Life-threatening cardiac arrhythmias (i.e. ventricular tachycardia, unstable SVT)

PRECAUTIONS:

- A. Monitor efficacy to nebulization by clinical status, oxygen saturation and respiratory rate and work of breathing.
- B. Monitor response to heart rate and blood pressure.
- C. Administer via nebulization ONLY.
- D. DO NOT administer IV/ IO/ IM/ IN.

PEDIATRIC DOSE:

Respiratory distress with audible stridor at rest (pts 6 months to 6 years old) - 0.5 ml (11.25 mg) of racepinephrine diluted with 2.5 ml of normal saline via nebulizer. May repeat once in 10 minutes if necessary. Contact OLMC for additional doses. **In the absence of Racepinephrine, you may substitute 5 ml (5 mg) of Epi 1:1000 via nebulizer.**

SUPPLIED: 40 mg / 20 ml pre-filled syringe or 2 mg/ml in 40 mg vial

PHARMACOLOGY AND ACTIONS:

Etomidate is a hypnotic drug without any analgesic activity. Intravenous injection of etomidate produces hypnosis characterized by rapid onset of action; usually within one minute. Duration of hypnosis is dose dependent but relatively brief, usually 3-5 minutes.

INDICATIONS:

- A. As an induction agent for use in rapid sequence intubation.
- B. As a sedation agent prior to synchronized cardioversion of unstable tachycardia.

CONTRAINDICATIONS:

Etomidate is contraindicated in patients who have a known hypersensitivity to the drug.

SIDE EFFECTS AND NOTES:

- A. The most frequent adverse reactions are transient injection site pain and transient skeletal muscle movements (myoclonus).
- B. Etomidate may also cause nausea and/or vomiting.

ADULT DOSING:

Induction agent for rapid sequence intubation -

0.3 mg / kg IV/IO slow push.

Synchronized cardioversion for unstable tachycardia -

0.15 mg / kg IV /IO push to a max of 10 mg. Wait 45-60 seconds for signs of sedation such as patient becoming verbally unresponsive or no longer following commands.

PEDIATRIC DOSING: Same as adult

SUPPLIED: 100 micrograms / 2 ml vial

PHARMACOLOGY AND ACTIONS:

Fentanyl is a potent synthetic opioid analgesic that produces analgesia and sedation. It is about 50-100 times more potent than morphine on a weight basis. Onset of action when given is 2-3 minutes. Peak effect occurs at 3-5 minutes and lasts 15-45 minutes.

INDICATIONS:

- A. Pain due to musculoskeletal injury or burns.
- B. Suspected ischemic chest pain.
- C. Post-intubation analgesia.
- D. CPR Induced Consciousness.

CONTRAINDICATIONS:

- A. Known allergy to fentanyl.
- B. Moderate to severe respiratory depression.

PRECAUTIONS:

- A. Fentanyl can cause respiratory depression that is reversible with naloxone. Respiratory depression can also be exacerbated by underlying lung disease and the use of other respiratory depressant drugs (benzodiazepines, alcohol, cyclic antidepressants). Have naloxone and respiratory support available when administering fentanyl.
- B. If administered rapidly and in very large doses, fentanyl can cause muscle spasm and chest wall rigidity. The only reliable treatment for this is neuromuscular blockade.
- C. The action of fentanyl is prolonged, and its elimination is slower in the elderly. Smaller maintenance doses are advisable.
- D. Fentanyl must be used cautiously in patients who have already received morphine for prehospital analgesia.

SIDE EFFECTS AND NOTES:

- A. If hypotension develops, it is usually responsive to naloxone administration and Trendelenburg position. If hypotension continues, follow Shock protocol.
- B. Check and document vital signs and patient response after each dose.
- C. The goal of fentanyl administration is patient comfort, not the total elimination of pain but the reduction in the perception of pain by the patient.

ADULT DOSING:

Pain Management -

IV/IN - 25-100 mcg. May repeat 25-50 mcg every 5 minutes as needed to a maximum of 500 mcg. IM – 25 -100 mcg. May repeat every 15 minutes as needed to a maximum of 400mcg. If BP < 100 mmHg or patient has minor altered mental status or respiratory depression - first dose fentanyl is 25 mcg, may repeat 25-50 mcg every 5 minutes to a maximum of 500 mcg. Monitor closely.

Post-Intubation analgesia -

After successful airway placement, administer fentanyl 25 to 100 mcg IV/IO if systolic BP \ge 100 mmHg (MAP is >65 mmHg). Repeat every 15 minutes as necessary to maintain analgesia.

CPR Induced Consciousness-

50 mcg IV/IO used in conjunction with midazolam. May repeat every 5 – 10 minutes as needed

PEDIATRIC DOSING: Pain Management -

1 mcg/kg IV. May repeat with 0.5 -1 mcg/kg every 5 minutes as needed to a maximum of 4 mcg/kg. Or, 2 mcg/kg IN repeated with 1 mcg/kg every 5 minutes as needed to a maximum of 4 mcg/kg. If no IV/IN, may give fentanyl 1-2 mcg/kg IM. May repeat every 15 minutes to a max of 4 mcg/kg. Do not exceed adult dosing. IN is preferred if no IV.

Post-Intubation analgesia -

After successful airway placement, administer fentanyl 1 mcg/kg IV/IO, not to exceed the adult dose, with repeat doses at 0.5 - 1 mcg/kg every 15 minutes.

SUPPLIED: 1 mg vial of powder / 1 ml vial of diluent

PHARMACOLOGY AND ACTIONS:

Glucagon is a hormone that causes glucose mobilization in the body. It works opposite to insulin, which causes glucose storage. It is released at times of insult or injury when glucose is needed and mobilizes glucose from body glycogen stores. Return to consciousness should be within 20 minutes of an IM dose if patient is hypoglycemic.

INDICATIONS:

- A. Known hypoglycemia (preferably demonstrated by blood glucose determination) when patient is confused or comatose and dextrose is not available or an IV cannot be started.
- B. Anaphylaxis in patients with beta-blockade when epinephrine is ineffective.

CONTRAINDICATIONS: None

PRECAUTIONS:

IV Dextrose is the treatment of choice for hypoglycemia in the patient who cannot tolerate oral glucose. The use of glucagon is restricted to patients who are seizing, comatose, combative, or with collapsed veins and in whom an IV cannot be started.

SIDE EFFECTS AND NOTES:

- A. Nausea and vomiting may occur with administration.
- B. Persons with no liver glycogen stores (malnutrition, alcoholism) may not be able to mobilize any glucose in response to glucagon.

ADULT DOSING:

Hypoglycemia -

1 mg IM.

Anaphylaxis (If epinephrine is ineffective in treating anaphylaxis in patients with beta-blockade)-

1 mg IM/IV.

PEDIATRIC DOSING:

Hypoglycemia -

0.02 mg/kg IM to a maximum of 1 mg.

SUPPLIED: 15 - 24 grams glucose in gel tubes

PHARMACOLOGY AND ACTIONS:

Glucose is the body's basic fuel and it produces most of the body's quick energy. Its use is regulated by insulin that stimulates storage of excess glucose from the bloodstream and glucagon that mobilizes stored glucose into the bloodstream.

INDICATIONS:

Oral glucose is indicated in the conscious patient where a suspicion of hypoglycemia exists, or a blood glucose measurement indicates a low blood glucose level.

CONTRAINDICATIONS:

Do not give to patients who cannot adequately protect their own airway.

PRECAUTIONS:

To give solutions orally, a patient must be continually assessed for the ability to protect his or her own airway.

SIDE EFFECTS AND NOTES:

- A. Research suggests that hyperglycemia may complicate, or worsen, a number of medical conditions (i.e. myocardial infarction, stroke). Oral glucose should be given to a conscious patient whenever hypoglycemia is documented by blood glucose meter. If these objective findings are not available, the EMT should use judgment based on signs and history.
- B. Effects will be delayed in the elderly and people with poor circulation.
- C. May be more tolerable if administered with liquid between dosages.
- D. Patient's condition may require more than one dose of oral glucose.

ADULT DOSING:

Hypoglycemia -

One tube or equivalent. Repeat as needed.

PEDIATRIC DOSING:

Same as adult

SUPPLIED: 5 mg / 1 ml vial

PHARMACOLOGY AND ACTIONS:

Haloperidol is an antipsychotic that may be used for pharmacological sedation by producing marked sedation and allaying apprehension. It also provides a state of mental detachment and indifference while maintaining a state of reflex alertness. Haloperidol may potentiate the effects of other CNS depressants. It also causes mild alpha-adrenergic blockade which can lead to peripheral vasodilatation and hypotension, as well as a reduction of the pressor effect of catecholamines. It is also a very effective anti-emetic. The onset of action of a single IV dose is from 5 - 15 minutes following administration, and the peak effect may not be apparent for up to 30 minutes. Duration is generally from 2 - 6 hours.

INDICATIONS:

- A. Sedation of combative patients to facilitate restraint.
- B. Nausea and vomiting

CONTRAINDICATIONS:

- A. Known allergy to haloperidol
- B. Known Parkinson's disease or use of dopamine agonists medications like carbidopa-levodopa (Sinemet), Pramipexole (Mirapex), or ropinirole (Requip).

PRECAUTIONS:

- A. Hypotension may occur; IV fluids and other measures to manage hypotension should be readily available.
- B. Use caution when administering haloperidol to patients who have taken other CNS depressant drugs (e.g., barbiturates, benzodiazepines, alcohol).
- C. Haloperidol may induce Torsade de Pointes. Monitor the patient's ECG Q-T interval following use.

SIDE EFFECTS AND NOTES:

- A. The most common side effects are hypotension and tachycardia, which usually responds to a fluid bolus.
- B. Dysphoric (restlessness) and dystonic reactions have been reported following administration. These symptoms can be treated with the administration of diphenhydramine.
- C. Haloperidol should be used with caution in patients with a seizure disorder or condition that causes seizures; other similar neuroleptics are known to lower the seizure threshold.

ADULT DOSING:

Pharmacological sedation -

5 - 10 mg IV, IO, IM. May repeat to a maximum of 20 mg (**For patients > 65:** 2 mg IV/ IO. May repeat after 15 mins. **or** 2.5 mg IM. May repeat after 15 - 20 mins.)

Nausea and vomiting

1.25 mg IV/IM

PEDIATRIC DOSING:

Haloperidol is not recommended for children.

Hydromorphone (Dilaudid®) – 20.144

OLMC REQUIRED: No

Santiam Ambulance - Only

SUPPLIED: 2 mg / 1 ml vial. Concentration and packaging may vary based on availability.

PHARMACOLOGY AND ACTIONS:

Hydromorphone is an opioid agonist that binds to several opioid receptors. Its analgesic characteristics are through its effect on the mu-opioid receptors. Hydromorphone has been shown to be 5 - 7 times more potent than morphine with a shorter duration of analgesia. Onset of action when given IV is 5 minutes and peak effect occurs at 8 - 20 minutes. Duration is 3 - 4 hours.

INDICATIONS:

- A. Pain due to burns or musculoskeletal injury.
- B. Suspected ischemic chest pain unresponsive to nitroglycerin.
- C. Post-intubation analgesia.

CONTRAINDICATIONS:

- A. Known allergy to hydromorphone.
- B. Blood pressure less than 100 mmHg systolic for pain management and post-intubation analgesia.
- C. Respiratory rate less than 14 breaths per minute or oxygen saturation less than 90%. For pediatric patients, vital signs should be maintained within the normal age-appropriate range.
- D. Patients < 12 months.

PRECAUTIONS:

- A. Hydromorphone causes respiratory depression that is reversible with naloxone. This respiratory depression is exacerbated by underlying lung disease (COPD, etc.) and other depressant drugs (Valium, alcohol, cyclic anti-depressants). Naloxone and respiratory support must be available when using hydromorphone.
- B. If hypotension develops, it is usually responsive to naloxone administration and Trendelenburg position. If hypotension persists, follow Shock protocol.
- C. The goal of hydromorphone administration is patient comfort (not the total elimination of pain but reduction in perception of pain by the patient).
- D. Use a 1 ml syringe for administration due to small volume.

SIDE EFFECTS AND NOTES:

- A. Common side effects include flushing, pruritus, diaphoresis, dry mouth, nausea and vomiting, asthenia, dizziness, headache, and somnolence.
- B. Serious adverse effects include hypotension, syncope, coma, increased intracranial pressure, seizures, respiratory depression, and apnea.

ADULT DOSING:

Pain Management -

0.25 - 0.5 mg IV. Repeat every 15 - 20 minutes up to a maximum of 2 mg. If no IV, give 0.5 - 1.0 mg IM. May repeat IM every 15 - 20 minutes to a maximum of 2 mg.

PEDIATRIC DOSING:

Pain - Musculoskeletal injuries-

For patients \geq 12 months: 0.01 mg/kg IV/IM not to exceed the adult dose. May repeat every 15 - 20 minutes to a maximum of 2 mg.

<u>NOTE:</u> Hydromorphone is not preferred in young infants and toddlers if fentanyl or morphine is available.

SUPPLIED: Liquid - 100 mg / 5 mL (Children's) ; 50 mg / 1.25 mL (Infant's); 200 mg tablets, capsules

PHARMACOLCOGY AND ACTIONS:

Ibuprofen, from isobutylphenyl propionic acid, is a nonsteroidal anti-inflammatory drug (NSAID) used for relieving pain, lowering fever, and reducing inflammation. Like other NSAIDs, it works by inhibiting the synthesis of prostaglandins, involved in mediating inflammation (swelling), pain, and fever. It achieves this effect on prostaglandin synthesis by inhibiting cyclooxygenase, an enzyme that is present in various tissues of the body.

INDICATIONS:

- A. Mild to moderate pain.
- B. Fever.

CONTRAINDICATIONS

- A. Known hypersensitivity to ibuprofen.
- B. Previous asthma, urticarial, or allergic reaction after taking aspirin or other NSAID.
- C. Recent heart surgery.
- D. Has taken ibuprofen in last 6 hours.
- E. Unable to take oral medication.
- F. Any signs of dehydration in pediatric patients.
- G. Patients less than 6 months old.

PRECAUTIONS:

Ibuprofen may cause a severe allergic reaction, especially in people who are allergic to aspirin. May cause stomach bleeding especially in patients:

- Older than 60 years
- Who have had stomach ulcers or bleeding problems
- Take blood thinners
- Take other medications containing NSAIDs.

ADULT & PEDIATRIC DOSING:

10 mg/kg PO to maximum of 600 mg.

Pediatric dosing using 100 mg/5 mL liquid

Weight	Dose	Volume
18 lbs / 7.5 kg	75 mg	3.75 mL
24 lbs / 10 kg	100 mg	5 mL
36 lbs / 15 kg	150 mg	7.5 mL
48 lbs / 20 kg	200 mg	10 mL
60 lbs / 25 kg	250 mg	12.5 mL
72 lbs / 30 kg	300 mg	15 mL

SUPPLIED: 0.5 mg / 2.5 ml vial individually or 0.5 mg packaged with 3 mg albuterol (Duo-Neb).

PHARMACOLOGY AND ACTIONS:

Ipratropium is an atropine derivative used for inhalation therapy. For severe asthma, ipratropium taken in addition to a short acting beta agonist (such as albuterol) can provide greater bronchodilation and clinical benefit than the beta agonist alone. It has no anti-inflammatory effects and does not decrease bronchial hyper-responsiveness.

INDICATIONS:

As a supplement to albuterol in patients with asthma and COPD.

CONTRAINDICATIONS: Do not use in patients with severe glaucoma.

PRECAUTIONS:

Ipratropium in the meter dose inhaler and auto-inhaler formulations should not be administered to individuals allergic to soy lecithin or related food products (e.g. soybeans, peanuts). The nebulized formulation may be administered to these patients.

SIDE EFFECTS AND NOTES:

- A. Dry mouth.
- B. Pharyngeal irritation.
- C. Increased intra-ocular pressure in glaucoma patients.

ADULT DOSING:

Asthma/ COPD -

3.0 ml DuoNeb (3.0 mg albuterol/0.5 mg ipratropium) via nebulizer. Repeat as needed X 2. Do not administer more than three total treatments.

PEDIATRIC DOSING: Same as adult dosing

SUPPLIED: 500 mg/10 ml vial.

PHARMACOLOGY AND ACTIONS:

Ketamine is a NDMA receptor antagonist, that is structurally similar to phencyclidine (PCP), that acts as a dissociative anesthetic agent by interrupting the connection between the thalamo-neocortical tracts and the limbic system in the brain, producing anesthesia. In addition, it has analgesic effects and can be used at lower doses for pain control – without causing anesthesia. It also stimulates catecholamine release from the adrenal glands causing an increase in heart rate, blood pressure, and cardiac output. Ketamine is also a bronchodilator and is a useful induction agent when intubating patients with severe bronchospasm.

INDICATIONS:

- A. As an induction agent for Drug Assisted Airway Management (DAAM).
- B. Post intubation sedation.
- C. Pain management.

CONTRAINDICATIONS:

- A. Known pregnancy.
- B. Non-traumatic chest pain.
- C. Patients with a history of schizophrenia or history of psychosis.

SIDE EFFECTS AND NOTES:

- A. Increased blood pressure due to catecholamine release.
- B. Emergence reaction can occur in 5 30% of patients. Duration of action is 10 20 minutes IV and continued sedation must be provided before the induction agent wears off.

ADULT DOSING:

Pain management -

A. 12.5 - 25 mg IV/IO slow push over 5 minutes, or by IV infusion over 15 minutes, or 25 - 50 mg IM. May repeat once after 30 min unless patient develops nystagmus, hallucinations, or dysphoric symptoms. Ketamine must be diluted prior to IV or IO administration for pain management. Either dilute 12.5 mg in 9.75 ml or 25 mg in 9.5 ml of Normal Saline for slow IVP or dilute 12.5 - 25 mg in 100 ml of Normal Saline and infuse over 15 minutes.*

OR

B. 1mg/kg VIA BREATH ACTUATED NEBULIZER (BAN). Add saline for total volume of 5 ml.

Induction agent for DAAM -

1 - 2 mg/kg IV/IO slow push over 1 minute. Dilute Ketamine with normal saline to a minimum of 10 ml total volume for a slower administration.

Post intubation sedation –

Initial dose is 1 mg/kg slow IV/IO push if not used for induction. If used for induction, initial dose is 0.5 mg/kg slow IV/IO push. May repeat 0.5 mg/kg every 15 minutes as necessary to maintain analgesia and sedation. <u>Ketamine should not be used for</u> sedation following ROSC in cardiac arrest patients.

*Ketamine should not be mixed with lactated ringers for dilution purposes due to compatibility concerns.

PEDIATRIC DOSING:

Pain management -

- A. For children ≥ 15, dose is 0.3 mg/kg IV slow push over 5 minutes, up to a max of 25 mg. Dose must be diluted in normal saline prior to administration.
 OR
- B. For children \geq 7, dose is 1 mg/kg **VIA BREATH ACTUATED NEBULIZER (BAN)**. Add saline for total volume of 5 ml.

Induction agent for DAAM - Same as adult

Nebulized ketamine with a concentration of 500 mg/10 ml

Pt. weight lbs/kg	Dose in mg (1 mg/kg)	mL of ketamine to add to nebulizer	mL of fluid to add to nebulizer	Concentration of ketamine in nebulizer after fluid is added
220 lbs/100 kg	100 mg	2 ml	3 ml	20 mg/ml
209 lbs/95 kg	95 mg	1.9 ml	3.1 ml	19 mg/ml
198 lbs/90 kg	90 mg	1.8 ml	3.2 ml	18 mg/ml
187 lbs/85 kg	85 mg	1.7 ml	3.3 ml	17 mg/ml
176 lbs/80 kg	80 mg	1.6 ml	3.4 ml	16 mg/ml
165 lbs/75 kg	75 mg	1.5 ml	3.5 ml	15 mg/ml
132 lbs/60 kg	60 mg	1.2 ml	3.8 ml	12 mg/ml
110 lbs/50 kg	50 mg	1 ml	4 ml	10 mg/ml
88 lbs/40 kg	40 mg	0.8 ml	4.2 ml	8 mg/ml
77lbs/35 kg	35 mg	0.7 ml	4.3 ml	7 mg/ml
66 lbs/30 kg	30 mg	0.6 ml	4.4 ml	6 mg/ml
55 lbs/25 kg	25 mg	0.5 ml	4.5 ml	5 mg/ml
44 lbs/20 kg	20 mg	0.4 ml	4.6 ml	4 mg/ml

(5 minus volume left in BAN) X Concentration of ketamine in BAN = mg administered to patient.

Example for 80 kg patient (80 mg dose) with 1.5 ml of volume left in BAN: $(5 - 1.5 \text{ ml}) \times 16 \text{ mg/ml} = 56 \text{ mg}$ administered. Waste amount in BAN is 24 mg + the 420 mg left in vial for a total waste of 444 mg.

SUPPLIED: 30 mg /1 ml vial

PHARMACOLOGY AND ACTIONS:

Ketorolac works by inhibiting cyclooxygenase-1 and 2 enzymes to block the synthesis of prostaglandins and reduces inflammation and pain.

INDICATIONS:

- A. Musculoskeletal pain.
- B. Flank pain from suspected kidney stone.

CONTRAINDICATIONS:

- A. Age < 2 or > 80.
- B. Multisystem trauma
- C. History of renal disease or kidney transplant.
- D. History of liver disease.
- E. Allergies to aspirin or other NSAIDs.
- F. Pregnancy, or lactating females.
- G. On anticoagulant, such as vitamin K antagonists (e.g. warfarin) or directing agents such as rivoraxaban, apixaban, edoxaban, lovenox, and dabigatran.
- H. Bleeding or clotting disorder or history of ulcer.
- I. Suspected cardiac chest pain.

SIDE EFFECTS AND NOTES:

- A. Burning or pain at the injection site
- B. Nausea and vomiting
- C. Dizziness
- D. Headache
- E. Itching
- F. Flushing

ADULT DOSING:

Pain management -

30 mg IM or 15 mg IV. Single dose only

PEDIATRIC DOSING (age 2-16 years):

Pain management -

1 mg/kg IM to a max of 30 mg or 0.5 mg/kg IV to a max of 15 mg.

SUPPLIED: 100 mg / 20 ml vial

PHARMACOLOGY AND ACTIONS:

Labetalol combines both selective, competitive alpha1-adrenergic blocking and nonselective, competitive beta-adrenergic blocking activity in a single substance. These actions decrease blood pressure without reflex tachycardia and without a significant reduction in heart rate.

INDICATIONS:

The treatment of uncontrolled, and sustained, hypertension in pregnant and postpartum women.

CONTRAINDICATIONS:

- A. Bronchial Asthma.
- B. Overt cardiac failure.
- C. Greater than first degree heart block.
- D. Cardiogenic shock.
- E. Severe bradycardia.

SIDE EFFECTS AND NOTES:

- A. Cardiovascular: Symptomatic postural hypotension, ventricular dysrhythmia, syncope, bradycardia, and heart block.
- B. CNS: Dizziness, tingling of the scalp/skin, numbness, vertigo.
- C. Respiratory: Wheezing, bronchospasm.
- D. GI: Nausea and vomiting.

ADULT DOSING:

For sustained elevation in BP > 160 mmHg systolic and/or \ge 110 mmHg diastolic (either one or both) that persists for at least 15 minutes or more in pregnant or post-partum women.

10 mg slow IV push over 1 - 2 minutes. May be repeated twice (3 doses total) every 15 minutes if BP is not within target range. Depending on effect of preceding dose, double remaining doses (e.g. 1st dose 10 mg, 2nd dose 20 mg, 3rd dose 40 mg. Maximum total dose 70 mg.) Target systolic BP 140 - 150 mmHg and diastolic BP 90 - 100 mmHg. Stop administration if HR < 60 bpm or other adverse effects.

PEDIATRIC DOSING: Not indicated for pediatric patients

OLMC REQUIRED: See Contraindications.

SUPPLIED: 100 mg / 5 ml of 2% solution in pre-filled syringe

PHARMACOLOGY AND ACTIONS:

Lidocaine depresses the automaticity of Purkinje fibers, raising stimulation threshold in the ventricular muscle fibers which makes the ventricles less likely to fibrillate. It has little antiarrhythmic effect on the atrial muscle in normal doses. The effect of a single bolus on the heart disappears in 10-20 minutes due to redistribution in the body. The metabolic half-life of lidocaine is about 2 hours.

INDICATIONS:

- A. Recurrent ventricular fibrillation or pulseless ventricular tachycardia.
- B. Following successful defibrillation from ventricular fibrillation or pulseless ventricular tachycardia.
- C. PVCs in a suspected ischemic event.
- D. Pain management following IO placement.

CONTRAINDICATIONS:

- Do not use in perfusing pts in the following situations without OLMC approval:
- A. Systolic BP is less than 90 mmHg.
- B. Heart rate is less than 50 beats per minute.
- C. Periods of sinus arrest are present.
- D. Second or third-degree heart block are present.

PRECAUTIONS:

- A. Lidocaine is not recommended in the treatment of supra-ventricular arrhythmias.
- B. If administering maintenance dosing and the patient begins seizing, stop the lidocaine dosing and treat per Seizure protocol.

SIDE EFFECTS AND NOTES:

- A. Side effects include drowsiness, dizziness, disorientation, confusion, and seizures.
- B. Hypotension.
- C. Lidocaine is metabolized in the liver and, therefore, patients with hepatic disease, shock or congestive heart failure will have decreased metabolism. All doses after the initial dose must be decreased to one-quarter of the initial dose.
- D. Toxicity is more likely in elderly patients.

ADULT DOSING:

V-Fib/Pulseless VT -

Bolus dose - 1.5 mg/kg IV/IO. Repeat to a max of 3 mg/kg if needed.

ROSC (from V-Fib/Pulseless VT arrest) -

If no antidysrhythmic is given prior to ROSC - 1.5 mg/kg repeated with 0.75 mg/kg every 10 minutes up to 3 mg/kg. If Lidocaine was the last anti-dysrhythmic given - 0.75 mg/kg every 10 minutes up to 3 mg/kg

PVCs (In the setting of an acute ischemic event only) -

Bolus 1.5 mg/kg IV/IO over 1 – 2 minutes. If no change, give 0.75 mg/kg IV/IO every 5 minutes, up to 3 mg/kg. When PVCs are suppressed, give 0.75 mg/kg every 10 minutes.

Pain management for IO placement -

40 mg (2cc's of 2% Lidocaine slowly over 2 minutes).

PEDIATRIC DOSING:

Same as adult for V-Fib/Pulseless VT, ROSC, and PVC's.

Pain management for IO placement- 0.5 mg/kg slowly, not to exceed 40 mg.

Seizures in eclampsia/pre-eclampsia. Asthma in pediatric patients.

SUPPLIED: 1 gram (50%) / 2 ml vial or 5 grams (50%) / 10 ml vial

PHARMACOLOGY AND ACTIONS:

Magnesium is a cation that is present in human cells and intercellular fluids. It acts as an antiarrhythmic agent and is useful in the treatment of polymorphic ventricular tachycardia due to an underlying prolonged QT interval, ventricular fibrillation and ventricular tachycardia

INDICATIONS:

- A. Polymorphic Ventricular Tachycardia (stable wide complex, irregular tachycardia associated with an underlying prolonged QT [Torsade de Pointes]).
- B. For the treatment of seizures in women with pre-eclampsia/eclampsia with OLMC approval.
- C. In severe asthma as a smooth muscle relaxant and inhibitor of histamine.

CONTRAINDICATIONS: None in the emergency setting.

PRECAUTIONS:

In the non-arrest patient, magnesium sulfate may cause hypotension, bradycardia, decreased reflexes, and respiratory depression.

ADULT DOSING:

Wide complex, irregular tachycardia -

2 grams IV/IO over 1-2 minutes.

Eclampsia/Pre-eclampsia -

Contact OLMC for dosing in this situation. Normal dose is 4 grams IV over 15-20 minutes.

Asthma -

Dose is 2 grams IV over 15-20 minutes.

PEDIATRIC DOSING:

Asthma -

Contact OLMC for dosing in this situation.

DILUTING FOR IV ADMINISTRATION

- A. Dilute each gram of magnesium sulfate in 8 ml of normal saline. (Example: Mix 1 gram in 8 ml of NS; mix 2 grams in 16 ml of NS)
- B. For use with IV pump, dilute either 2 grams or 4 grams of magnesium sulfate in 100 ml of normal saline or lactated ringers (in Buretrol or 100 ml bag).

SUPPLIED: 10 mg / 2 ml vial

PHARMACOLOGY AND ACTIONS:

Midazolam is a benzodiazepine with potent sedative, anti-anxiety, and anticonvulsant properties. It also causes significant antegrade amnesia when administered IV.

INDICATIONS:

- A. Patient actively seizing upon EMS arrival or having repetitive seizures without regaining consciousness.
- B. To relieve anxiety and produce amnesia during cardioversion, pacing, or following Drug Assisted Airway Management (DAAM).
- C. To facilitate restraint in patients whose cause of agitation is likely drug ingestion (especially stimulants), withdrawal, or from a postictal state.
- D. CPR Induced Consciousness.

CONTRAINDICATIONS:

In seizures, do not give unless patient is actively seizing.

PRECAUTIONS:

Midazolam causes respiratory depression and/or hypotension especially if administered rapidly. Monitor patient closely.

SIDE EFFECTS AND NOTES:

- A. Common side effects include drowsiness, hypotension, respiratory depression, and apnea. These are more likely to occur in the very young and the elderly. Rarely, patients may experience paradoxical agitation.
- B. Respiratory depression is more likely in patients who have taken other CNS depressant drugs such as opioids alcohol, sedative-hypnotics, or when given rapidly.
- C. Midazolam is metabolized in the liver and excreted by the kidney. Doses should be adjusted accordingly in patients with underlying hepatic or renal diseases and low flow states such as congestive heart failure.

ADULT DOSING:

Seizures -

2.5 - 5 mg IV/IO or 5 mg IM/IN. Repeat every 5 minutes until seizure stops. **Pharmacological sedation -**

2.5 mg IV/IO or 5 mg IM. May repeat once in 10 minutes.

For immediate threat (RASS +3 or +4): 2.5 - 5 mg IV/IO or 5 - 10 mg IM with repeat doses of 1 - 2 mg IV/IO every 5 minutes as needed.

Induction medication for DAAM (Least desirable option) -

10 mg IV/IO if systolic BP is \geq 100 mmHg.

5 mg IV/IO if systolic BP < 100 mmHg.

Sedation after intubation & for induced hypothermia shivering -

2.5 - 5 mg IV/IO if systolic BP is \geq 100 mmHg. Repeat every 15 minutes as necessary to maintain sedation.

Sedation before cardioversion (with no IV) -

5 mg IM/IN

Sedation for Pacing -

2.5 - 5 mg IV/IO or 5 mg IM/IN, may repeat once. Call OLMC for additional orders.

CPR Induced Consciousness-

Up to 2.5 mg IV/IO used in conjunction with fentanyl. May repeat every 5 - 10 minutes as needed.

PEDIATRIC DOSING:

Seizures -

0.1 mg/kg IV/IO to a max of 5 mg, or

- 0.3 mg/kg IM/IN to a max of 10 mg
- * Repeat every 5 minutes until seizure stops.

Pharmacological sedation (Age > 12 only) -

0.1 mg/kg IV/IM, max 10 mg IM and 6 mg IV. May repeat every 10 mins. as needed. (Refer to pediatric pharmacological sedation flow chart in the Agitated Patient Protocol 10.015).

Induction medication for DAAM (Least desirable option) -

0.2 mg/kg IV/IO not to exceed adult dose.

Sedation after intubation with or without paralytics -

0.1 mg/kg IV/IO, max dose 2.5 mg, repeat every 15 minutes as necessary to maintain sedation.

Sedation before cardioversion -

0.2 mg/kg IM/IN to a max of 5 mg

Sedation for pacing -

0.1 mg/kg IV/IO, max dose 5 mg. May repeat once after 5 minutes.

* Call OLMC for additional orders.

SUPPLIED: Varies

PHARMACOLOGY AND ACTIONS:

Morphine is a narcotic analgesic that induces drowsiness, mental clouding, and mood changes. It also increases venous capacitance, decreases venous blood return

(preload), and reduces systemic vascular resistance at the arteriolar level (afterload). This may lead to decreases in myocardial oxygen demand. Onset of action when given IV is 2-3 minutes and peak effect occurs at 7-10 minutes. Duration is 3-5 hours.

INDICATIONS:

- A. Suspected ischemic chest pain unresponsive to nitroglycerin.
- B. Pain due to burns or musculoskeletal injury.

CONTRAINDICATIONS:

- A. Known allergy to morphine or sulfates (Sulfa drugs are not sulfates).
- B. Blood pressure less than 100 mmHg systolic.
- C. Respiratory rate less than 14 breaths per minute or oxygen saturation less than 90%. For pediatric patients, vital signs should be maintained within the normal age-appropriate range.

PRECAUTIONS:

- A. Morphine causes respiratory depression that is reversible with naloxone. This respiratory depression is exacerbated by underlying lung disease (COPD, etc.) and other depressant drugs (Valium, alcohol, cyclic anti-depressants). Naloxone and respiratory support must be available when using morphine.
- B. If hypotension develops, it is usually responsive to naloxone administration and Trendelenburg position. If hypotension persists, follow Shock protocol.
- C. Trauma or pain of the head or abdomen.

SIDE EFFECTS AND NOTES:

- A. The goal of morphine administration is patient comfort (not the total elimination of pain but reduction in perception of pain by the patient).
- B. Morphine is a Schedule II controlled substance. Follow your agencies Controlled Substance policy or procedure for control and monitoring of use.

ADULT DOSING:

Pain - Musculoskeletal injuries, burns, chest pain -

2-8 mg IV. Repeat every 5 to max of 20 mg. If no IV give 5-10 mg IM. May repeat IM with 5 mg every 15 minutes to a maximum of 20 mg.

PEDIATRIC DOSING (< 20kg):

Pain - Musculoskeletal injuries, burns, chest pain -

0.1 mg / kg IV/IM. May repeat IM after 15 minutes. Do not exceed adult dosing.

SUPPLIED: 2 mg / 2 ml pre-filled syringe

PHARMACOLOGY AND ACTIONS:

Naloxone is an opioid antagonist that competitively binds to opioid receptor sites, but which exhibits almost no pharmacologic activity of its own. Duration of effect is 1-4 hours.

INDICATIONS:

- A. Reversal of opioid effects, particularly respiratory depression, due to opioid drugs either ingested or injected or administered during treatment. Opioid drugs include fentanyl, morphine, meperidine, hydromorphone, oxycodone, hydrocodone, and codeine.
- B. Diagnostically in coma of unknown etiology to rule out or reverse opioid depression.

CONTRAINDICATIONS: Do not use in neonates.

PRECAUTIONS:

- A. In patients physically dependent on opioids, violent withdrawal symptoms may occur. Be prepared to restrain the patient.
- B. Some opioid intoxications may require up to 8 mg of naloxone to reverse symptoms (e.g. methadone, carfentanil).

SIDE EFFECTS AND NOTES:

- A. The duration of some opioids is longer than naloxone, repeat doses may be necessary. <u>Monitor the patient closely</u>. Patients who have received naloxone must be transported to the hospital because coma may reoccur when naloxone wears off.
- B. Side effects are rare. Do not hesitate to use if indicated.
- C. If no effect is seen from naloxone administration, consider other causes of coma.

ADULT DOSING:

Reversal of opioid effects, coma of unknown etiology -

0.5 mg IV. Repeat every 2 minutes up to 2 mg titrating to respiratory rate. If no IV, give 2 mg IM/IN. If no response to initial dose and opiate intoxication is still suspected, repeat 2 mg IV/IM/IN every 3-5 minutes up to a maximum of 8 mg total.

PEDIATRIC DOSING:

Reversal of opioid effects, coma of unknown etiology -

0.1 mg / kg IV/IM/IN up to 2 mg. May repeat q 3 - 5 minutes up to 2 mg / dose. Max total dose 8 mg. Do not use in neonates.

OLMC REQUIRED: See Contraindications (B).

SUPPLIED: 0.4 mg metered dose spray, 0.4 mg tablets, 50 mg/10 ml vial, paste 2%

PHARMACOLOGY AND ACTIONS:

Nitroglycerin is a vasodilator. It is a smooth muscle relaxant that reduces venous tone causing pooling of blood in the peripheral veins, decreasing peripheral resistance, and thereby decreasing cardiac preload. It also causes mild dilation of the coronary arteries.

INDICATIONS:

- A. Presumed ischemic chest pain.
- B. Decompensated heart failure.
- C. SCAPE (Sympathetic Crashing Acute Pulmonary Edema).

CONTRAINDICATIONS:

- A. Blood pressure less than 100 mmHg systolic.
- B. Patients who have taken Viagra® (sildenafil citrate), Levitra® (vardenafil HCI) or other similar drugs within 24 hours, or who have taken Cilais® (tadalafil) within 48 hours. Contact OLMC for direction.

PRECAUTIONS:

- A. Nitroglycerine can cause hypotension in 10% of patients.
- B. Nitroglycerin should be used with caution in patients with an inferior myocardial infarction (ST elevation in II, III and AVF) as this can result in hypotension due to an associated right ventricle infarction (RVI).
- C. RVI may be present in up to 50% of inferior myocardial infarctions. 12-lead ECG clues to RVI include STE in III > II. RVI can also be confirmed with a right sided 12 lead ECG and STE ≥ 1 mm in V₄R. RVI patients are preload dependent and may benefit from IV fluids.
- D. Generalized vasodilatation can cause profound hypotension and reflex tachycardia.
- **E.** IV should be established prior to administration in patients who have not taken nitroglycerin previously, or who have a potential for hemodynamic instability.

SIDE EFFECTS AND NOTES:

- A. Common side effects are headache, flushing, and dizziness.
- B. Because nitroglycerin causes generalized smooth muscle relaxation, may relieve chest pain caused by esophageal spasm.

ADULT DOSING:

Chest pain-

0.4 mg SL every 5 minutes until pain is relieved as long as systolic BP is greater than 100 mmHg. **Decompensated heart failure –**

Nitroglycerine 0.4 mg SL; repeat every 3 - 5 minutes. (<u>Do not administer nitroglycerine without OLMC</u> approval if patient has taken sildenafil (Viagra®), vardenafil (Levitra®) or other similar drugs in the last 24 hours, or tadalafil (Cialis®) within the last 48 hours).

SCAPE - Sympathetic Crashing Acute Pulmonary Edema (extreme respiratory distress, systolic BP > 160 mmHg, diaphoresis, tachycardia, decreased oxygen saturation).

- 0.4 mg SL; repeat every 3 5 minutes.
- Once CPAP/BiPAP is established; Nitro paste 1"-2" to chest if respiratory distress persists and

systolic BP remains > 120 mmHg.

PEDIATRIC DOSING:

Contact OLMC for dosing.

SUPPLIED: 4 mg/4 ml ampules or vials

PHARMACOLOGY AND ACTIONS:

Norepinephrine stimulates alpha receptors in the peripheral vasculature, producing vasoconstriction related increase in systemic blood pressure. Concurrent beta receptor stimulation may produce increases in heart rate and mild bronchodilation.

INDICATIONS:

Obstructive, cardiogenic and distributive shock unresponsive to fluid administration.

CONTRAINDICATIONS: Hypovolemic shock.

PRECAUTIONS:

- A. Norepinephrine should be given in a large, patent vein (i.e. antecubital or larger). Do not administer through a hand or leg vein, as these are more likely to be affected by vaso-occlusive diseases and more prone to ischemic complications.
- B. Extravasation of norepinephrine into tissue may cause necrosis. The IV should be checked for patency prior to administration and monitored continuously.
- C. Norepinephrine is a potent vasoconstrictor and may cause hypertension. The rate of flow should be carefully monitored, and blood pressures checked often.
- D. Consider hypovolemia and treat this with appropriate fluids before administration of norepinephrine.

SIDE EFFECTS AND NOTES:

- A. Symptoms may include headache, palpitations, tachycardia, chest pain, and eventual hypertension.
- B. Reflex bradycardia can result from an increase in blood pressure.

ADULT DOSING:

Cardiogenic/Distributive/Obstructive shock -

Begin at 4 mcg/min. If no response, increase every 5 minutes in 4 mcg/min increments to max of 12 mcg/min. Goal is a systolic blood pressure of \geq 90 mmHg.

PEDIATRIC DOSING:

Begin at 0.1 mcg/kg/min. If no response in 5 min, increase to 0.2 mcg/kg/min. If still no response after 5 more minutes may increase to 0.4 mcg/kg/min. Goal is age appropriate systolic blood pressure.

MIXING/ADMINISTRATION:

Add one 4 mg ampule or vial to 500 ml of NS or LR, or two 4 mg ampules or vials to 1000 ml of NS or LR for a concentration of 8 mcg/ml. Administer via infusion pump or 60 gtt/ml infusion set (**Infusion pump preferred**).

Adults (8 mcg/ml concentration)

Mcg/min	4	8	12
Drops/min	30	60	90

SUPPLIED:

10 mg orally dissolving tablets (ODT)

PHARMACOLOGY AND ACTIONS:

- A. Dopamine and serotonin (5-HT) antagonist, along with anticholinergic, antihistaminic, and anti-alpha-adrenergic effects.
- B. Has anxiolytic properties.
- C. Low incidence of extrapyramidal effects.

INDICATIONS:

To avoid the need for physical restraint in the mildly agitated patient between the ages of 18 – 65 who is willing to take an oral agent. (**RASS +1, see patient restraint protocol 30.120).**

CONTRAINDICATIONS

A. Patients less than 18 or greater than 65 years of age.

PRECAUTIONS:

- A. May cause QT prolongation, but unlikely in single dose. Obtain ECG before administration if known history or suspicion for prolonged QT or cardiovascular disease.
- B. Known hypersensitivity.
- C. Use with caution in suspected drug overdose.

ADULT DOSING (Age 18 – 65):

10mg PO - Administer tablet immediately once it is removed from the blister unit or bottle. Tablets disintegrate in the mouth and can be swallowed subsequently with saliva or liquid.

Patients who have received olanzapine (ODT) may be transported directly to Unity Hospital (see Behavioral Health Emergencies protocol 30.025).

Patients < 6 months except for children in spinal immobilization or children receiving chemotherapy.

SUPPLIED: 4 mg oral tablet; 4 mg / 2 ml vial

PHARMACOLOGY AND ACTIONS:

Ondansetron is a potent, highly selective serotonin (5-HT3) receptor agonist. Its precise mode of action in the control of nausea is not known. Pharmacologic agents and other triggers may cause release of 5-HT3 receptors. Ondansetron blocks the initiation of this reflex. Ondansetron is commonly used in the treatment of nausea in patients who are receiving chemotherapy or as a postoperative nausea treatment. Peak plasma concentrations of the drug occur 10 minutes after IV administration, and 40 minutes after IM injection. Both routes have the same elimination half-life of 4 hours.

INDICATIONS:

Prevention and control of uncomplicated nausea and vomiting.

CONTRAINDICATIONS:

Known hypersensitivity to Zofran or similar medications.

PRECAUTIONS:

- A. Hypersensitivity reactions have been reported in patients who have exhibited hypersensitivity to other 5-HT3 medications (Anzemet®, Kytril®).
- B. Patients with bowel obstruction should be monitored closely following administration.
- C. Ondansetron may precipitate if mixed with alkaline solutions.
- D. ECG changes including QT interval prolongation and Torsade de Pointes have been observed in patients receiving ondansetron. Monitor patient's ECG closely.

SIDE EFFECTS AND NOTES:

- A. The most common side effects include headache, dizziness, drowsiness, and shivers.
- B. Body aches, agitation, dysuria, hypotension, and rash have also been reported in a very small number of patients.

ADULT DOSING:

Nausea & vomiting -

8 mg oral dissolving tablet or 4 mg IV/IM. Give slowly over two minutes if giving IV. If nausea and/or vomiting are inadequately controlled after 10 minutes, may repeat dose once.

PEDIATRIC DOSING:

Nausea & vomiting (children 6 months - 2 yrs) 2 mg oral dissolving tablet.
Nausea & vomiting (children 2 yrs - 12 yrs) 4 mg oral dissolving tablet or 0.1 mg/kg IV/IM. Give slowly over two minutes if giving IV. Do not exceed 4 mg.

SUPPLIED: Various. D cylinder contains 415 liters at 2,000 psi.

PHARMACOLOGY AND ACTIONS:

Oxygen added to the inspired air raises the amount of oxygen in the blood and the amount delivered to the tissues. Breathing in most persons is regulated by small changes in acid/base balance and CO₂ levels and it takes a large drop in oxygen concentration to stimulate respiration.

INDICATIONS:

- A. Suspected hypoxemia or respiratory distress from any cause.
- B. Acute chest pain in which cardiac ischemia or myocardial infarction is suspected.
- C. Shock from any cause.
- D. Major trauma.
- E. Carbon monoxide poisoning.

CONTRAINDICATIONS: None

PRECAUTIONS:

- A. If the patient is not breathing adequately on their own, the treatment of choice is ventilation with oxygen, not just supplemental oxygen.
- B. In a small percentage of patients with chronic lung disease, administration of oxygen will decrease respiratory drive. Do not withhold oxygen because of this possibility. Be prepared to assist ventilations if needed.
- C. Titrate oxygen to the lowest level required to achieve an SpO₂ \ge 94%.

SIDE EFFECTS AND NOTES:

- A. Non-humidified oxygen is drying and irritating to mucous membranes.
- B. Restless may be an important sign of hypoxia.
- C. Oxygen toxicity is not a risk in acute administration.
- D. Nasal cannula prongs work equally well on nose and mouth breathers.

DELIVERY:		
Nasal cannula -		
2-8 lpm	24 –	40% inspired O ₂
Simple face mask	-	
6 lpm	50 –	60% inspired O ₂
Rebreather mask -		
10-12 lpm		90% inspired O ₂
Bag valve mask -		
Room air		21% inspired O ₂
12 lpm		40% inspired O ₂
With reservoir	+	90% inspired O ₂

Oxymetazoline Hydrochloride (Afrin®) – 20.245

OLMC REQUIRED: No

SUPPLIED: 0.5 fl oz nasal solution or other various sizes

PHARMACOLOGY AND ACTIONS:

Oxymetazoline hydrochloride is a selective alpha 1 adrenergic receptor agonist and alpha 2 adrenergic receptor partial agonist that causes localized vasoconstriction.

INDICATIONS:

Epistaxis uncontrolled by direct pressure.

CONTRAINDICATIONS:

- A. Allergy to oxymetazoline hydrochloride.
- B. Monoamine Oxidase Inhibitor (MAOI) use within the past 14 days.
- C. Diastolic blood pressure > 110 mmHg.

SIDE EFFECTS AND NOTES:

- A. Avoid administration into the eyes, which will dilate pupils.
- B. Temporary burning, stinging, dryness in the nose, runny nose, and sneezing may occur.

ADULT DOSING:

Epistaxis -

Instill two sprays to each affected nostril.

PEDIATRIC DOSING:

- A. Follow adult dosing.
- B. Oxymetazoline hydrochloride should be avoided if child cannot follow instructions to blow their nose or are unable to tolerate the administration of a nasal medication.

Oxytocin (Pitocin®) – 20.246

OLMC REQUIRED: No

SUPPLIED: 10 unit/ml, 1 ml vial

PHARMACOLOGY AND ACTIONS:

Synthetic pituitary hormone which stimulates uterine contractions to assist with control of postpartum bleeding or atony.

INDICATIONS:

Prophylactic use to reduce risk of postpartum hemorrhage after delivery.

CONTRAINDICATIONS:

- A. Uterine Rupture
- B. Incomplete delivery

PRECAUTIONS:

When administered IV, must be given slowly over 1 minute. Rapid infusion may lead to hypotension and dysrhythmias.

SIDE EFFECTS AND NOTES:

Confusion, seizures, difficulty breathing, fast or irregular heartbeat, headache, hives, pelvic or abdominal pain, skin rash, or itching.

ADULT DOSING:

Postpartum-

10 IU IV/IM if the neonate is a singleton. For multiple births, administer only after last neonate has been delivered.

PEDIATRIC DOSING:

Not indicated for pediatrics. Consider OLMC.

SUPPLIED: 100 mg in 10 mL vial

PHARMACOLOGY AND ACTIONS:

Rocuronium is a non-depolarizing neuromuscular blocking agent causing skeletal muscle relaxation. Rocuronium produces a pure reversible competition between antagonist molecules and acetylcholine (Ach) for occupancy at the Ach binding site. Neuromuscular blockade occurs within 90 seconds for induction dose and 1 to 3 minutes for maintenance dose. Time to recovery is 20 to 30 minutes. Metabolism is 5 to 35% renal and the remainder by the liver.

INDICATIONS:

- A. For sustained neuromuscular blockade in the intubated patient.
- B. For induction intubation (RSI/DSI) in the patient when succinylcholine is contraindicated or unavailable.

CONTRAINDICATION: Maintainance of paralysis in patients in status epilepticus

PRECAUTIONS:

- A. Use of pulse oximetry is required.
- B. Rocuronium does not substantially affect heart rate or rhythm, systolic or diastolic blood pressure, mean arterial pressure, cardiac output, or systemic vascular resistance.
- C. Rocuronium has no effect on consciousness and must be used with a sedative or induction agent.
- D. Rocuronium should not be administered simultaneously with furosemide, methylprednisolone, or sodium bicarbonate.

ADULT AND PEDS DOSING:

Maintenance of post-intubation paralysis - 0.5 mg/kg IV/IO.

Induction for intubation – 1.2 mg/kg IV/IO.

OLMC REQUIRED: Pediatric hyperkalemia and crush injury

SUPPLIED: 50 mEq / 50 ml pre-filled syringe

PHARMACOLOGY AND ACTIONS:

Sodium bicarbonate is an alkalotic solution which neutralizes acids found in the blood. Acids are increased in the blood when body tissues become hypoxic due to cardiac or respiratory arrest. Acidosis depresses cardiac contractility and cardiac response to catecholamines and makes the heart more likely to fibrillate and less likely to defibrillate.

Current guidelines no longer recommend routine use of sodium bicarbonate, except in cases of arrest secondary to hyperkalemia, cyclic antidepressant overdose, or acidosis.

INDICATIONS:

- A. Acidosis associated with PEA and asystole.
- B. To control arrhythmias or asystole in cyclic antidepressant overdose or hyperkalemia.
- C. Chlorine inhalation injury.

CONTRAINDICATIONS: None

PRECAUTIONS:

- A. Addition of too much bicarbonate may result in alkalosis that is difficult to reverse and may cause as many problems in resuscitation as acidosis.
- B. May increase cerebral acidosis, especially in diabetic ketoacidosis.
- C. Do not mix sodium bicarbonate with calcium preparations. Slowly flush one drug from the catheter before administering the other.

SIDE EFFECTS AND NOTES:

Each amp of sodium bicarbonate contains 50 mEq of sodium. This may increase intravascular volume and hyperosmolarity resulting in cerebral impairment.

ADULT DOSING:

Sodium Channel Blockade overdose (Tricyclic Antidepressants,

Diphenhydramine, propranolol, Type 1a or 1c anti-dysrhythmics) -

1 mEq/kg IV or IO.

PEA, asystole -

1 mEq/kg IV or IO. May repeat q 10 minutes at 0.5 mEq/kg.

Hyperkalemia -

50 mEq IV or IO.

Crush injury -

50 mEq IV or IO.

Chlorine Inhalation - 2.5 ml of 8.4% Sodium Bicarbonate via nebulizer

PEDIATRIC DOSING:

- A. Use same dosing as for adult with exception of hyperkalemia and crush injury; call OLMC for dosing in that situation.
- B. For children less than 10 kg (1 yr.), dilute by one-half with normal saline prior to administration.

SUPPLIED: 200 mg / 10 ml vial

PHARMACOLOGY AND ACTIONS:

Succinylcholine is a short acting motor nerve depolarizing skeletal muscle relaxant. It competes with acetylcholine to combine with cholinergic receptors in the motor end plate causing depolarization inhibiting neuromuscular transmission. After intravenous injection, paralysis is obtained within 1-2 minutes and persists for approximately 4-6 minutes. Effects then start to fade and return to normal. It has no effect on consciousness. Muscle relaxation begins in the eyelids and jaw, then progresses to the limbs, abdomen, diaphragm and finally intercostal muscles. Succinylcholine is hydrolyzed by plasma pseudocholinesterase and is excreted by the kidneys.

INDICATIONS:

To achieve temporary paralysis where endotracheal intubation is indicated.

CONTRAINDICATIONS:

- A. Hypersensitivity to the drug.
- B. Major burns and crush injuries between 48 hours and 6 months old.
- C. Stroke or spinal cord injuries with profound residual defecits between 48 hours and 6 months old.
- D. Neuromuscular disease (e.g. muscular dystrophy, multiple sclerosis).
- E. Suspected hyperkalemia (e.g. end-stage renal disease patients who have missed dialysis).

PRECAUTIONS:

- A. Succinylcholine shall not be administered unless personnel trained and authorized in this procedure are present and ready to perform the procedure.
- B. Oxygen, ventilation equipment, and resuscitation drugs should be readily available.
- C. Succinylcholine produces paralysis but does not alter a person's level of consciousness. Paralysis in the conscious patient is very frightening, therefore, sedation should be provided to the patient during the procedure. Verbal explanations should be provided to the patient during the procedure, even if you do not think they can hear you.

SIDE EFFECTS AND NOTES:

In rare individuals, because of pseudocholinesterase deficiency, paralysis may persist for a prolonged period. Be prepared to continue to assist ventilations as needed.

ADULT DOSING:

Rapid or delayed sequence intubation -1.5 mg/kg IV/IO

PEDIATRIC DOSING:

Rapid or delayed sequence intubation -1.5 mg/kg IV/IO for patients > 6 years old.

2 mg/kg IV/IO for patients < 6 years old.

Tranexamic Acid (TXA)- 20.277

OLMC REQUIRED: No

SUPPLIED: 1000 mg (1 gram) / 10 ml vial

PHARMACOLOGY AND ACTIONS:

Tranexamic acid (TXA) is a synthetic analog of the amino acid lysine. It reversibly binds to lysine receptor sites on plasminogen to decrease the conversion of plasminogen to plasmin. This antifibrinolytic effect reduces breakdown of fibrin and helps to stabilize clots to reduce bleeding. TXA also has anti-inflammatory properties.

INDICATIONS:

- A. Moderate to severe head trauma, either blunt or penetrating, in patients with a $GCS \le 12$ and with a reactive pupil.
- B. Hemorrhagic shock from blunt or penetrating trauma with a systolic blood pressure < 70 mmHg.
- C. Significant postpartum hemorrhage (> 500 ml).

CONTRAINDICATIONS

- A. Patients less than 15 years old (or weight < 50 kg if age unknown)
- B. > 2 hours from time of injury for hemorrhagic shock or TBI
- C. GCS of 3 with no reactive pupil
- D. EMS chest compressions (manual or mechanical)
- E. Patients with clinical concern for epilepsy/seizures, MI, stroke, PE, DVT, renal failure, or dialysis
- F. Known or suspected pregnancy
- G. Drowning
- H. Hanging
- I. Burns > 20% TBSA

PRECAUTIONS AND SIDE EFFECTS:

- A. Hypotension has been observed with rapid IV injection.
- B. TXA, by causing clots to get stronger, can make MI, stroke, PE, and DVTs more challenging to manage.
- C. TXA is renally cleared, so its use in patients with known renal failure or dialysis should be avoided.
- D. Reported side effects have included seizures, nausea, vomiting, and chest pain.

ADULT DOSING (Age \geq 15):

Head trauma, hemorrhagic shock, or postpartum hemorrhage: 2 g slow IV/IO push.

PEDIATRIC DOSING:

Not indicated in patients < 15 years of age suffering from head trauma, hemorrhagic shock, or postpartum hemorrhage. Consider OLMC.

SUPPLIED: 10 mg vial of powder and 10 ml vial of diluent solution

PHARMACOLOGY AND ACTIONS:

Vecuronium is a non-depolarizing neuromuscular blocking agent causing skeletal muscle relaxation. It reversibly binds the acetylcholine receptor, blocking the action of acetylcholine. Neuromuscular blockade occurs within 2-3 minutes. Time to recovery is 30-45 minutes. Vecuronium metabolism is 5-35% renal with the remainder done in the liver.

INDICATIONS:

- A. For sustained neuromuscular blockade in the intubated patient.
- B. As the first line agent for rapid sequence intubation (RSI) or delayed sequence intubation (DSI) in the patient where succinylcholine is contraindicated.

CONTRAINDICATIONS:

Patients in status epilepticus who require intubation.

PRECAUTIONS:

- A. Patients with renal or hepatic failure may experience prolonged paralysis.
- B. Vecuronium has no effect on consciousness and must be used with a sedative or induction agent.

SIDE EFFECTS AND NOTES:

- A. Vecuronium exhibits minimal side effects and does not substantially affect heart rate or rhythm, systolic or diastolic blood pressure, mean arterial pressure, cardiac output, or systemic vascular resistance.
- B. Vecuronium can be used to maintain paralysis even if intubation was performed without Succinylcholine.

ADULT DOSING:

0.1 mg/kg IV/IO.

PEDIATRIC DOSING: Same as adults.

Procedures

Regional EMS Patient Treatment Protocols - 2023

DEFINITION:

An AICD is an implanted defibrillator device that consists of a lead system that senses cardiac activity, logic circuitry to analyze sensed signals, a power supply for device function and generating high voltage, and a capacitor that stores and delivers shocks. This device activates when brady and/or tachyarrhythmias are detected within programmed parameters.

INDICATIONS:

Consider application of a magnet to deactivate an implanted cardioverter defibrillator that is firing inappropriately. **Call OLMC prior to application**. Inhibition of AICD devices should be considered when continuous ECG monitoring verifies malfunction and ACLS is readily available.

PROCEDURE:

- A. Contact OLMC.
- B. Monitor ECG and verify sinus rhythm AND inappropriate defibrillator discharge.
- C. Locate the position of the AICD device.
- D. Place doughnut magnet directly over the device.
- E. After proper positioning and AICD deactivation, tape magnet securely in place and transport.

NOTES & PRECAUTIONS:

- A. It is very important to make the correct diagnosis before utilizing this protocol. Be sure that the ECG is showing a normal sinus rhythm without ectopy AND indications of recurrent AICD discharges.
- B. Some AICD devices will emit varying beeping or continuous tones when magnets are applied, other will not. Disregard these tones.
- C. If the magnet placement is successful in overriding the pulse generation of the AICD, **DO NOT REMOVE THE MAGNET**. Some units will return to normal operation after removal from the magnetic field.
- D. Magnets should be stored so as not to come into contact with magnetic sensitive materials, i.e., monitor screens, tapes, credit cards, magnetic door entry cards, and other electronic equipment.
- E. A small percentage of AICDs are impervious to magnetic fields (AICD recipients who normally work around magnetic fields have these special units). These will not be deactivated with the doughnut magnet. In such cases, advise OLMC and transport.
- F. Consider use of the AICD magnet in deactivating cardiac pacemaker malfunctions. Application of a magnet to a pacemaker changes the pacing to asynchronous mode but will not turn off the pacemaker. Call OLMC prior to application.
- G. Identification information of the AICD type, date implanted, and location of implantation should accompany the patient to the hospital. This information is typically found on a wallet card that the patient has.

Breath Actuated Nebulizer (AEROECLIPSE® II BAN®) – 30.030

DEFINITION:

The AEROECLIPSE[®] II BAN[®] Nebulizer only creates aerosol when the patient breathes in. This means medication is not wasted between breaths This puts the patient in control of their aerosol treatment and creates a safer environment. Other nebulizers continuously produce aerosol whether you are inhaling, exhaling or resting. This means the medication may be lost into the room instead of delivered to the lungs. In some cases, this can be hazardous to the health of others who may be nearby during the treatments.

INDICATIONS:

Nebulized ketamine administration for pain management. The BAN[®] <u>must</u> be used when nebulizing ketamine to avoid the risk of secondary exposure to aerosol medications.

CONTRAINDICATIONS:

Only to be used by patients \geq 7 years of age.

PROCEDURE:

- A. Unscrew and remove the top of the nebulizer.
- B. Carefully place the prescribed medication into the nebulizer cup and replace the nebulizer top.
- C. Make sure that the quarter turn valve on top of the nebulizer is pointed towards the dotted arrow which indicates that it is in breath actuated mode. (see picture below).
- D. Connect the nebulizer to an oxygen source.
- E. Have the patient place the mouthpiece into their mouth and instruct them to breathe slowly and deeply and to exhale normally through the device as desired.
- F. The green button on the top of the BAN[®] will go down when the patient breathes in and will go up when the patient breathes out.
- G. Follow your agency's controlled medication disposal process for any remaining medication left in the BAN[®].



Continuous Positive Airway Pressure – 30.032

DEFINITION:

Continuous Positive Airway Pressure (CPAP) has been shown to rapidly improve vital signs, gas exchange, and to decrease the work of breathing, the sense of dyspnea, and the need for endotracheal intubation in patients who suffer from shortness of breath secondary to CHF/Pulmonary edema, COPD, or asthma. In patients with CHF, CPAP improves hemodynamics by reducing preload and afterload.

INDICATIONS:

Medical patients complaining of <u>moderate to severe</u> respiratory distress meeting <u>ALL</u> the following criteria:

- A. Is awake, oriented and has the ability to maintain an open airway.
- B. Has signs and symptoms consistent with either CHF/pulmonary edema, COPD, or asthma.
- C. Has a systolic blood pressure above 90 mmHg.
- D. Is over 12 years old and is able to fit the CPAP mask.

CONTRAINDICATIONS:

- A. Respiratory arrest.
- B. Non-cooperative patient.
- C. Suspected pneumothorax.
- D. Hemodynamically unstable.
- E. Inability to maintain mask seal.
- F. Active vomiting.

PROCEDURE:

- A. EXPLAIN and COACH THE PATIENT ON THE PROCEDURE.
- B. Ensure adequate oxygen supply to ventilation device.
- C. Place the patient on continuous pulse oximetery and end-tidal CO2.
- D. Turn on device. Set device to minimum flow (2-5 cmH₂O).
- E. Place the CPAP over the patient's mouth and nose (consider having the patient hold the mask against their face initially to reduce anxiety).
- F. Secure the mask with the provided straps.
- G. Check for air leaks.
- H. Monitor and document the patient's respiratory response to the treatment.
- I. Continue to coach patient to keep mask in place and readjust as needed to a maximum of 10 cmH_2O .
- J. <u>IF RESPIRATORY STATUS DETERIORATES, REMOVE THE DEVICE AND CONSIDER BAG</u> VALVE MASK VENTILATION AND/OR ENDOTRACHEAL INTUBATION.

REMOVAL PROCEDURE:

CPAP therapy needs to be continuous and should not be removed unless the patient cannot tolerate the mask or experiences continued or worsening respiratory failure.

SPECIAL NOTES:

- A. If unable to maintain oxygen saturation > 90%, administer positive airway pressure via BVM and PEEP valve.
- B. Contact the receiving hospital as soon as possible that a patient with CPAP is enroute to their hospital so they can be prepared for the patient.
- C. Reassessment of the patient's status is critical, and documentation should be performed every 5-10 minutes until patient is stable.
- D. CPAP mask may be removed temporarily to administer nitroglycerin.
- E. Suctioning of secretions may be required on some patients.
- F. Watch for gastric distention and/or nausea.
- G. The CPAP monometers should be used to determine and adjust CPAP pressures as this will vary depending on the device used and whether nebulization is occurring simultaneously.
- H. Monitor mean arterial blood pressure closely in all patients with CPAP.

Procedures - Revised 9/17/21

INDICATIONS:

This technique is to be used only when other attempts to establish an airway have been unsuccessful (i.e., you are unable to intubate or ventilate using BVM) and respiratory obstruction exists. Such conditions are most likely to be found with foreign-body obstruction, facial and laryngeal trauma, inhalation, thermal, or caustic injury to the upper airway, angioneurotic edema, upper airway bleeding, epiglottitis, and severe croup.

PROCEDURE:

Place the patient in a supine position with support under the shoulders and mild hyperextension of the neck. Palpate the neck in the midline and locate the slight depression just below the notch of the thyroid cartilage. This is the position of the cricothyroid membrane.

Control-Cric ™

- A. Position the patient supine and identify the cricothyroid membrane.
- B. Stabilize the larynx with thumb and middle finger with the non-dominant hand.
- C. Utilizing the Cric-Knife[™], incise the skin making a vertical incision from the mid-thyroid cartilage to the cricoid cartilage (about 2 finger breadths in length). A longer incision may be needed if the patient has a thick neck. If landmarks are clearly visible, a horizontal incision may be used.
- D. After palpating the cricothyroid membrane, tum the Cric-KnifeTM to a horizontal position over the cricothyroid membrane.
- E. Push the blade downward, perpendicular to the trachea, until the blade is fully inserted and the airway is entered.
- F. While maintaining a downward force, slide the tracheal hook down the handle with your thumb until the hook is felt to enter the trachea, and disengages from the handle.
- G. Grab the hook with the non-dominant hand, lifting up on the thyroid cartilage.
- H. Insert the Cric-KeyTM through the incision. Placement can be confirmed by moving the device along the anterior wall of the trachea to feel for the tracheal rings. Tenting of the skin, difficulty advancing the Cric-KeyTM, or lack of tactile feedback from the tracheal rings suggests incorrect placement.
- I. Once placement has been confirmed, advance the Cric-Key[™] to the flange. Stabilize the Cric-Key[™] tube and pivot the tracheal hook toward the patient's shoulder to remove from the airway.
- J. While stabilizing the Cric-Key[™] tube, remove the Cric-Key[™] introducer. Inflate the cuff until resistance is met.
- K. Confirm proper placement of the airway device utilizing standard methods (presence of breath sounds, absence of gastric sounds) and quantitative waveform capnography.
- L. Secure the device with the stabilizing strap.

Needle Cricothyrotomy - (pediatric patients 12 years and younger).

- A. Assemble equipment: 14g or 16g angiocath, 3 cc syringe, 3.0 ETT adapter, oxygen, BVM.
- B. Place the patient in a supine position with support under the shoulders and mild hyperextension of the neck unless C-Spine injury is suspected.
- C. Palpate the neck in the midline and locate the slight depression just below the notch of the thyroid cartilage. This is the position of the cricothyroid membrane.
- D. Prepare the area with antiseptic solution.
- E. Stabilize the airway between thumb and forefingers.
- F. Insert the needle with catheter into the cricothyroid membrane at a 30-degree angle caudally (toward the patient's feet).
- G. When the needle is through the membrane. Stop and aspirate for air to ensure tracheal entry.
- H. Advance the catheter over the needle and then remove the needle.
- I. Attach the 3.0 ETT adapter to the hub of the catheter and begin ventilations with the BVM.
- J. Secure the cannula with tape after confirming correct placement by auscultation for breath sounds (5-point check). Observe for kinking of cannula.
- K. Consider sedation with midazolam as with RSI if not already given.

NOTES & PRECAUTIONS:

- A. Hazards in performing this procedure are primarily those of damage to nearby structures; major vessels to either side of the midline, to the vocal cords if the puncture is made too high, or a through and through injury of the trachea if the puncture is made too deeply. The latter is more commonly seen in infants and children whose tracheas may be deceptively narrow.
- B. Palpation of the cricothyroid membrane is very difficult in the infant and young child. The key to success is immobilization of the trachea throughout the procedure.
- C. Needle cricothyrotomy is only a temporizing measure providing oxygenation not adequate ventilation.

INDICATIONS:

- A. Airway obstruction
- B. Need for airway protection
- C. Respiratory failure

PROCEDURE:

Cardiac Arrest Patients:

- A. Patients in cardiac arrest can typically be intubated without the use of an induction agent and paralytics. Pre-oxygenation and apneic oxygenation are not indicated.
- B. Assemble and check all equipment:
 - 1. Cardiac monitor
 - 2. Suction
 - 3. EtCO₂
 - 4. Pulse Oximeter
 - 5. O₂ tank w/regulator
 - 6. Mask and BVM
 - 7. Intubation equipment (VL, DL)
 - 8. Backup devices ready: Bougie, supraglottic airway, surgical airway (cric kit)
- C. Intubate in a controlled, but timely manner. (Consider use of a supraglottic airway to minimize CPR interruptions or when ALS resources are limited.)
- D. Use of the bougie is encouraged for endotracheal intubation to facilitate first pass success.
- E. Verify placement of ET tube using waveform capnography and a careful five-point check. <u>Monitor waveform capnography continuously.</u>
- F. Secure the tube utilizing ETT securing device. Record ET Tube depth at the teeth or gum line. Depth in adults is height based. Reasonable targets are 21 cm for women, and 23 cm for men at the teeth.
- G. Avoid interruptions to CPR when securing a patient's airway. Once secured, deliver 1 breath every 6 secs. (10 breaths/min) asynchronous with compressions. About 1 second per breath, with visible chest rise. <u>Optional method</u>: 30:2 compression/ventilation ratio with advanced airway until ROSC. Post-ROSC, deliver 1 breath every 6 seconds.
- H. Ventilate and monitor patient's vital signs including SpO₂.
- I. If signs of "CPR Induced Consciousness" are present, administer up to 2.5 mg of midazolam IV/IO **and** 50 mcg of fentanyl. May repeat as needed every 5 10 minutes.
- J. Consider orogastric tube placement.

Drug Assisted Airway Management (DAAM) in Perfusing Patients:

- A. DAAM is the technique of using medications to overcome the body's protective airway reflexes to facilitate airway insertion using sedatives and paralytics.
- B. Two DAAM techniques are Rapid Sequence intubation (RSI) and Delayed Sequence Intubation (DSI).
- C. RSI and DSI choice should be based on paramedic discretion and/or medical director preference.
- D. If the patient is agitated and difficult to preoxygenate, consider DSI with ketamine to optimize oxygenation and facilitate resuscitation.

- E. Assemble and check equipment: Two O₂ tanks with regulators, nasal cannula, BVM with mask, EtCO₂, intubation equipment, suction, back up devices (bougie, SGA, cric kit).
- F. Attach pulse oximeter, cardiac monitor, BP cuff, and waveform capnography.
- G. Establish IVs or IOs, if not already done.
- H. Verbalize missed airway plan to the entire team and verify/mark surgical airway landmarks.
- I. Physiologically optimize patient prior to intubation with a MAP > 65 mmHg (systolic BP > 100 mmHg). **Preoxygenation and denitrogenation are essential steps in every DAAM.**
- J. Treat hypotension with fluids and Push Dose epinephrine 10 -20 mcg every 1- 5 minutes, with a goal MAP > 65 mmHg (SBP ≥ 100 mmHg).
- K. Place nasal cannula and administer oxygen at 15 lpm. Continue apneic oxygenation during the procedure.

Delayed Sequence Intubation procedure	Rapid Sequence Intubation procedure
 Administer Induction Agent: Ketamine 1 - 2 mg/kg IV/IO slow push over 60 seconds for sedation and analgesia prior to paralysis 	 Positioning: Ensure patient is positioned ear to sternal notch with head of bed/backboard elevated ≥ 15°. Maintain apneic O₂ via NC at 15 lpm throughout
 2. Positioning: Ensure patient is positioned ear to sternal notch with head of bed/backboard elevated ≥ 15°. Maintain apneic O₂ via NC at 15 lpm throughout 	 2. Preoxygenation and Denitrogenation: If breathing adequately, administer oxygen via NRB at 15 lpm If breathing inadequately, use a BVM at 15 lpm with OPA/NPA. Perform two-person BVM
 3. Preoxygenation and Denitrogenation: If patient is breathing adequately, Hold BVM (NO VENTILATIONS) using 15 lpm of oxygen and NPA/OPA with two-handed mask seal and PEEP @ 10. Increase PEEP if unable to achieve SpO₂ ≥ 94% 	 ventilations with two-handed thumbs-down seal on mask Ensure the patient has a SpO₂ ≥ 94% for at least 3 minutes before medication administration. If unable to achieve a SpO₂ ≥ 94%, consider DSI
 If patient is breathing inadequately, VENTILATE with BVM using 15 lpm of oxygen and OPA/NPA with two-handed mask seal and PEEP @ 10. Increase PEEP if unable to achieve SpO₂ ≥ 94% Upon reaching SpO₂ ≥ 94%, begin 3-minute countdown to allow for complete denitrogenation. See letter K below 	 3. Administer Induction Agent: Etomidate 0.3 mg/kg IV/O OR Ketamine 1 - 2 mg/kg IV/IO slow push over 60 seconds OR Midazolam 0.2 mg/kg IV/IO (least desirable option) If systolic BP ≥ 100 mmHg- max dose 10 mg If systolic BP < 100 mmHg- max dose 5 mg
 4. Paralysis Administer one of the following paralytics: Succinylcholine ≥ 6 years or > 20 kg - 1.5 mg/kg IV/IO < 6 years or < 20 kg - 2 mg/kg IV/IO <u>OR</u> Rocuronium 1.2 mg/kg IV/IO <u>OR</u> Vecuronium 0.1 mg/kg IV/IO 	 4. Paralysis Immediately following induction agent, administer one of the following paralytics: Succinylcholine ≥ 6 years or > 20 kg - 1.5 mg/kg IV/IO < 6 years or under 20 kg - 2 mg/kg IV/IO OR Rocuronium 1.2 mg/kg IV/IO OR Vecuronium 0.1 mg/kg IV/IO

- L. If unable to achieve $SpO_2 \ge 94\%$, consider failed airway plan, including use of a supraglottic airway.
- M. Perform intubation approximately 60 seconds after succinylcholine or rocuronium, and 2 3 minutes after vecuronium.
- N. Use of the bougie is encouraged to facilitate first pass success.
- O. If SpO₂ drops to < 94% during intubation attempt, ventilate with BVM using 100% oxygen before next attempt.
- P. If intubation unsuccessful, consider use of BVM and/or backup supraglottic airway device.
- Q. If unable to ventilate with BVM or backup airway, proceed to surgical airway (cricothyrotomy).
- R. If bradycardia occurs, first ensure adequate oxygenation and ventilation, and if persistent, administer atropine 0.5 mg IV/IO (Pediatric patients: 0.02 mg/kg IV/IO. Minimum dose 0.1 mg. Do not exceed adult dose.)
- S. Verify placement of ET tube using waveform EtCO₂ and a careful five-point check.
- T. Continue cardiac, waveform EtCO₂, and pulse oximetry monitoring at all times.
- U. Following intubation, titrate PEEP down to lowest setting to maintain $SpO_2 \ge 94\%$.
- V. Insert an oral airway or compatible bite-block device if needed.
- W. Secure the endotracheal tube and record the depth at the teeth/gums.
- X. Recheck and document ET tube placement after every patient movement or change in vital signs. For sudden hypoxia, consider DOPE:
 - 1. **D**islodgement
 - 2. Obstruction
 - 3. Pneumothorax
 - 4. Equipment issue
- Y. After successful airway placement, administer fentanyl <u>PLUS</u> midazolam, <u>OR</u> ketamine for analgesia and sedation:
 - 1. Fentanyl and midazolam:
 - a. Fentanyl 50 100 mcg IV/IO if SBP ≥ 100 mmHg (MAP > 65 mmHg), repeat every 15 minutes as necessary to maintain analgesia. (Pediatric dosing, 1 mcg/kg, not to exceed the adult dose with repeat doses at 0.5-1 mcg/kg)
 - b. Midazolam 2.5 5 mg IV/IO if SBP ≥ 100 mmHg (MAP > 65 mmHg).
 Repeat every 15 minutes as necessary to maintain sedation. (Pediatric dose of midazolam is 0.1 mg/kg IV/IO up to 2.5 mg), <u>OR</u>
 - c. Analgesia should be addressed first. Opioids are preferred first line agents before benzodiazepines. Ensure hemodynamic stability before giving a second agent to facilitate analgesia and sedation.
 - Ketamine: Initial dose is 1 mg/kg slow IV/IO push if not used for induction. If used for induction, initial dose is 0.5 mg/kg slow IV/IO push. May repeat 0.5 mg/kg every 15 minutes as necessary to maintain analgesia and sedation.
 Ketamine should not be used for sedation following ROSC in cardiac arrest patients.

Endotracheal Intubation – 30.040

- Z. Consider ketamine for ongoing sedation in airway management if:
 - 1. Non-depolarizing neuromuscular blockade (e.g. vecuronium, rocuronium) is used at any point as a paralytic agent, or
 - 2. Ketamine is used for DAAM.
- AA. If additional paralysis is needed, administer vecuronium 0.1 mg/kg <u>or</u> rocuronium 0.5 mg/kg IV/IO.
- AB. Consider orogastric tube placement.

NOTES & PRECAUTIONS:

- A. If unable to establish and/or maintain an adequate airway, transport patient, <u>including trauma patients</u>, to the nearest hospital to obtain definitive airway control.
- B. An attempt is defined as the insertion of the laryngoscope blade or rescue airway past the teeth. In most situations, intubation attempts should be limited to 2 per paramedic (with a maximum of 4 attempts prior to/during transport).
- C. <u>DO NOT</u> rely solely on monitoring equipment. Auscultate for lung sounds and/or revisualize with laryngoscope (VL or DL) if there is any doubt about tube placement.
- D. Continuously monitor the patient's overall condition including vital signs, SpO₂, EtCO₂, cardiac rhythm, perfusion, and ease of ventilation post-intubation.
- E. Succinylcholine, rocuronium and vecuronium do not affect the level of consciousness and should be used with etomidate/ketamine/midazolam.
- F. Succinylcholine is contraindicated in the following:
 - 1. Known hypersensitivity.
 - 2. Major burns and crush injuries between 48 hours and 6 months old.
 - 3. Stroke or spinal cord injuries with profound residual deficits between 48 hours and 6 months old.
 - 4. Neuromuscular disease (e.g., muscular dystrophy).
 - 5. Suspected hyperkalemia (ESRD patients on dialysis).
- G. Avoid vecuronium and rocuronium in patients suspected of having underlying status epilepticus (seizures).
- H. In DSI, start with 1 mg/kg of ketamine for induction. If disassociation is not achieved, administer a second 1 mg/kg dose of ketamine.
- I. Rapid administration of ketamine can lead to apnea. Ketamine should be administered slowly over 60 seconds. Dilute ketamine with normal saline to a minimum of 10 ml total volume for a slower administration.
- J. Ketamine can cause laryngospasm and may cause an emergence reaction with vivid dreams.
- K. Preoxygenation and denitrogenation can be challenging in some instances (e.g., ARDS, pneumonia). Consider a BVM with a PEEP valve or non-invasive positive pressure ventilation (e.g., CPAP/BiPAP).
- L. Patients dependent on sympathetic tone may develop profound hypotension post intubation. This should be treated with fluids and/or push dose pressors per the shock protocol. It is always best to have push dose epinephrine available.

DOCUMENTATION:

Visualization of the cords (if applicable), size and depth of tube at the teeth/gums, number of attempts, 5-point check and equal chest expansion, EtCO₂ waveform device used/reading, SpO₂, any other devices/ techniques used, and reconfirmation of placement after each patient movement.

Indications:

1. Capnography shall be used when available with the use of all advanced airway procedures and as required by treatment guidelines.

Procedure:

- 1. Attach capnography sensor to the monitor first to allow for room air calibration, then attach to the advanced airway or any other oxygen delivery device, including bag-valve mask and nasal cannula.
- 2. Note that $EtCO_2$ level and waveform changes. Normal $EtCO_2$ levels range from 30s and 40s, but this may vary based on the patient's underlying, respiratory, and metabolic status. EtCO₂ levels that rise from a normal baseline to or above 50 may indicate hypoventilation is occurring.
- Respiratory and metabolic values shall be documented in the ePCR.
- 4. The capnometer shall remain in place and be monitored throughout prehospital care and transport.
- 5. Any loss of EtCO₂ detection or waveform may indicate an airway problem and should be immediately addressed and thoroughly documented.
- Document the procedure and results in the ePCR. 6.

Notes:

- 1. EtCO₂ readings may be unreliable if the patient is in shock or has poor perfusion.
- 2. Patient stimulation, use of a BVM, or use of Naloxone may be appropriate based on the situation.

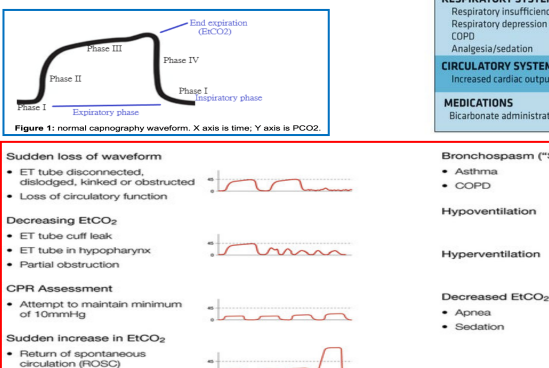
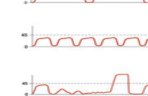


Table 1: Factors Affecting EtCO₂

CAUSES OF ELEVATED EtCO ₂	CAUSES OF DECREASED
METABOLISM Pain Hyperthermia Shivering	METABOLISM Hypothermia Metabolic acidosis
RESPIRATORY SYSTEM Respiratory insufficiency Respiratory depression COPD Analgesia/sedation	RESPIRATORY SYSTEM Alveolar hyperventilation Bronchospasm Mucus plugging
CIRCULATORY SYSTEM Increased cardiac output MEDICATIONS Bicarbonate administration	CIRCULATORY SYSTEM Hypotension Sudden hypovolemia Cardiac arrest Pulmonary emboli



Hyperventilation



DEFINITION:

The i-gel[®] is a disposable supraglottic airway created as an alternative to endotracheal intubation or mask ventilation. The i-gel[®] is designed for positive pressure ventilation as well as for spontaneously breathing patients.

INDICATIONS:

The i-gel[®] supraglottic airway device can be used as an alternative to endotracheal intubation in those patients who need a secure airway.

CONTRAINDICATIONS:

- A. Trismus (clenched jaw), limited mouth opening.
- B. Suspected upper airway obstructions secondary to laryngeal edema, smoke inhalation, foreign body, tumor, mass, or abscess.

i-gel Size	Patient Size	Patient Weight (kgs)	Patient Weight (Ibs)
1	Neonate	2.5	4-11
1.5	Infant	5-12	11-26
2	Small pediatric	10-25	22-55
2.5	Large pediatric	25-35	55-77
3	Small adult	30-60	66-132
4	Medium adult	50-90	110-198
5	Large adult	90+	198+

SIZES:

Size should be determined on lean body mass

PROCEDURE:

- A. Identify correct size i-gel[®].
- B. Lubricate i-gel[®] prior to insertion with water soluble gel and only to the back side of the device.
- C. If equipped, ensure that the supplemental oxygen port is capped.
- D. Position the patient. The patient should always be in the "sniffing position" prior to insertion unless head/neck movements are considered inadvisable or are contraindicated.
- E. If needed, use tongue depressor or curved laryngoscope blade to facilitate passage of i-gel[®] through the oropharynx.
- F. Grasp the lubricated i-gel[®] firmly along the integral bite block.
- G. Position the device so that i-gel® cuff outlet is facing towards the chin of the patient.
- H. Introduce the leading soft tip into the mouth of the patient in a direction toward the hard palate. The leading edge of the i-gel's[®] tip must follow the curvature of the patient's hard palate upon insertion. Glide the device downward and backward along the hard palate with a continuous but **gentle** push until a definitive resistance is felt.
- I. Determine appropriate depth of insertion. When placed correctly, the tip of the i-gel[®] will be within the upper esophageal opening and the cuff will be against the laryngeal framework. The incisors will be resting on the integral bite block. There is a horizontal black line on sizes 3, 4, and 5 indicating optimal position.(Fig. 1)

i-gel® Supraglottic Airway Device – 30.072



- J. Secure i-gel[®] to maxilla with approved holder, strap, or tape.
- K. If gastric distention is present or fluid is present in the gastric channel of i-gel[®], an appropriately sized lubricated orogastric tube (Fig. 2) may be passed down the gastric channel.
- L. Attach capnography per protocol.

i-gel Size	Maximum Size of Orogastric Tube (French Gauge) or French Suction Catheter	Fig. 2
1	N/A	
1.5	10	
2	12	
2.5	12	
3	12	
4	12	
5	12/14	

NOTES & PRECAUTIONS:

- A. Do not use excessive force to insert the device or orogastric tube.
- B. Sometimes a feel of "give-way" is felt before the end point resistance is met. This is due to the passage of the i-gel[®] bowl through the faucial pillars (pharyngo-epiglottic folds).
- C. Once resistance is met and the teeth are located on the integral bite block, do not repeatedly push the i-gel[®] down or apply excessive force during insertion.
- D. Do not allow peak airway pressure of ventilation to exceed 40 cm H₂O (Zoll Series 731 EMV+ or equivalent).
- E. Patients with any condition which may increase the risk of a full stomach (e.g. hiatal hernia, sepsis, morbid obesity, pregnancy, or a history of upper gastrointestinal surgery), may increase the risk of aspiration.

DEFINITION:

Intraosseous cannulation is an alternative technique for establishing vascular access in critical adult and pediatric patients when peripheral IV access is difficult or time sensitive.

INDICATIONS:

- A. Intraosseous infusion is indicated in emergency situations when lifesaving fluids or drugs should be administered, and IV cannulation is difficult, impossible, or too time-consuming to perform.
- B. If a peripheral IV cannot be established after two attempts or within 60–90 seconds of elapsed time *and* in:
- C. Adult and pediatric patients, within the proper weight range, who present with one or more of the following clinical conditions:
 - 1. Cardiac arrest.
 - 2. Hemodynamic instability (BP < 90 mmHg and clinical signs of shock).
 - 3. Imminent respiratory failure.
 - 4. Status epilepticus with prolonged seizure activity greater than 10 minutes, and refractory to IM anticonvulsants.
 - 5. Toxic conditions requiring immediate vascular access for antidote.
- D. Intraosseous placement may be considered prior to peripheral IV attempts in cases of cardiopulmonary or traumatic arrest, in which it may be obvious that attempts at placing an IV would likely be unsuccessful and/or too time consuming, resulting in a delay of life-saving fluids or drugs.

EZ-IO[®] PROCEDURE:

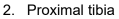
- A. Determine patient's weight.
- B. Assemble all necessary equipment:
 - 1. The 25 mm (Blue) $EZ-IO^{\circ}$ needle can be utilized for patients who weigh \geq 3 kg.
 - The 45 mm (Yellow) EZ-IO[®] needle can be used for adult insertions (larger individuals) where the 25 mm (Blue) needle is not adequate. The 45 mm needle should be used for all humeral IOs.
 - 3. EZ-Stabilizer[®] should be used to secure the needle.
- C. Site selection:
 - 1. Proximal humerus is preferred in adult patients to achieve the following:
 - a. Increased flow rates
 - b. Decreased pain
 - c. Closer access to central circulation (heart) during cardiac arrest and for resuscitation
 - 2. Proximal Tibia
 - Distal Tibia
- D. Site landmarks:
 - 1. Proximal humerus (contraindicated in children)
 - a. Ensure that the patient's hand is resting on the abdomen and that the elbow is adducted (close to the body).

Intraosseous Access & Infusion - 30.080

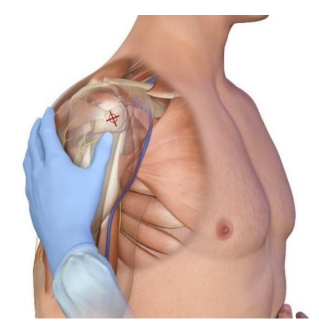
D. Site landmarks:

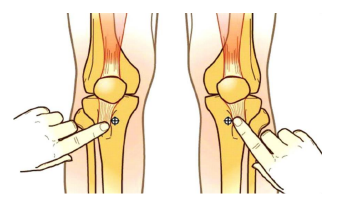
1. Proximal humerus (contraindicated in children)

- a. Ensure that the patient's hand is resting on the abdomen and that the elbow is adducted (close to the body).
- b. Insertion site is located directly on the most prominent aspect of the greater tubercle. Slide thumb up the anterior shaft of the humerus until you feel the greater tubercle, this is the surgical neck. Approximately 1 cm (depending on patient anatomy) above the surgical neck is the insertion site.



- a. Palpate the landmarks at the proximal tibia (patella and tibial tuberosity).
- Insertion site should be approximately one finger width (2 cm) medial to the tibial tuberosity, along the flat aspect of the tibia.







3. Distal tibia

-Two finger widths proximal to the medial malleolus along the midline of the tibia.

- E. Needle insertion
 - 1. Prep the surface with povidone-iodine or chlorohexidine and wipe dry with a sterile gauze pad.
 - 2. Stabilize patient's extremity and begin insertion from a 90-degree angle to the insertion site. Push the needle set through the skin until the tip touches the bone.
 - 3. With the needle tip against the bone, assure adequate needle length by ensuring at least one black line (5 mm) is visible outside the skin.
 - 4. Gently advance the needle set into position-<u>do not force</u>. Stop when you feel the "pop" or "give" on smaller patients.
 - 5. When needle is in proper position, remove stylet, place the EZ-Stabilizer[®] on the hub, but do not secure EZ-Stabilizer[®] yet.
 - 6. Connect EZ-Connect tubing, primed with saline, to IO hub.
 - 7. Rapid bolus or "power" flush with approximately 10 ml normal saline (administer lidocaine to the awake patient prior to flushing).
 - 8. Confirm the catheter position
 - a. Catheter is stable at a 90-degree angle to the bone, able to aspirate blood (not always able to aspirate even with the line in the proper position), and fluids flow without evidence of extravasation).
 - b. If insertion fails, leave the needle in place and clamp the EZ-Connect; do not attempt second insertion on same extremity.
 - 9. Secure the EZ-Stabilizer[®] when patency is confirmed.
 - 10. Consider additional bolus of saline if flow rates slower than expected.
 - 11. Utilize a blood pressure cuff or pressure bag around the IV bag to help infuse fluids.
 - 12. Monitor for patency frequently.
- F. Pain Management
 - 1. If the procedure is performed on a conscious patient, immediately following placement of the IO needle **and before saline flush**, administer 2 ml (40 mg) of 2% lidocaine slowly over 2 minutes (rule of 2 ml of 2% over 2 min). Wait approximately 30–60 seconds before flushing with normal saline.
 - 2. In the event a patient regains consciousness and complains of severe pain secondary to the IO insertion, temporarily stop infusing the fluids, and administer lidocaine as in F.1 above. Wait approximately 30–60 seconds before continuing fluid administration.
 - 3. If fluids do not flow freely, flush IO site with an additional 10 ml normal saline.

PEDIATRIC EZ-IO[®] PROCEDURE (patients weighing 3-39 kg)

- A. Assemble all equipment
 - 1. The 15 mm (Pink) EZ-IO[®] needle or 25 mm (Blue) EZ-IO needle should be used for patients who weigh less than 3kg (approximately 6 lb.). The 15 mm needle, if carried, is used primarily on neonates.
 - 2. The 25 mm (Blue) EZ-IO[®] needle should be utilized for pediatric patients who weigh \geq 3 kg or when the 15 mm (Pink) is deemed inadequate or not carried.
 - 3. EZ-Stabilizer should be used to secure the needle.
- B. Site selection (Patients weighing 3-39 kg)
 - 1. Proximal Tibia
 - a. Palpate the landmarks at the proximal tibia (patella and tibial tuberosity).
 - b. Insertion site should be one finger width below and one finger width medial of the tibial tuberosity.
 - c. If the tibial tuberosity cannot be identified on the child, then the insertion site may be two finger widths below the patella, then medial along the flat aspect of the tibia.
 - 2. Distal femur
 - a. Secure the leg outstretched to ensure the knee does not bend.
 - b. Locate upper edge of the patella. Insertion site is one finger width above and then one finger width medial (towards the inner leg) from the upper patella edge. This location will avoid the growth plate of the distal femur.



- C. Needle insertion
 - 1. Prep the surface with povidone-iodine or chlorohexidine and wipe dry with a sterile gauze pad.
 - 2. Stabilize patient's leg and begin insertion from a 90-degree angle to the insertion site. Push the needle set through the skin until the tip touches the bone.
 - 3. With the needle tip against the bone, assure adequate needle length by ensuring at least one black line (5 mm) is visible outside the skin.
 - 4. Gently advance the needle set into position–do not force. Stop when you feel the "pop" or "give".
 - 5. When needle is in proper position, remove stylet, place the EZ-Stabilizer[®] on the hub, but do not secure EZ-Stabilizer[®] yet.
 - 6. Connect EZ-Connect tubing, primed with saline, to IO hub.
 - 7. Rapid bolus or "power" flush with approximately 5 ml normal saline.
 - 8. Confirm the catheter position:
 - a. Catheter is stable at a 90-degree angle to the bone, able to aspirate blood, and fluids flow without evidence of extravasation).
 - b. If insertion fails, leave the needle in place and clamp the EZ-Connect; do not attempt second insertion on same extremity.
 - 9. Secure the EZ-Stabilizer[®] when patency is confirmed.
 - 10. Consider additional bolus of saline if flow rates slower than expected, no more than 2-3 ml normal saline
 - 11. Consider a blood pressure cuff or pressure bag to help infuse fluids.
 - 12. Monitor for patency frequently.
 - D. Pain Management
 - 1. If the procedure is performed on a conscious patient, immediately following placement of the IO needle, administer 0.5 mg/kg of 2% lidocaine slowly over 2 minutes, not to exceed adult dose of 40 mg. Wait approximately 30–60 seconds before flushing with normal saline.
 - 2. If fluids do not flow freely, flush IO site with an additional 2-3 ml normal saline.

PEDIATRIC PROCEDURE WITH MANUAL IO DEVICE:

- A. Assemble equipment
 - 1. Approved bone marrow needles, 15- or 18-gauge size (Jamshidi)
 - 2. Povidone-iodine or chlorohexidine preps
 - 3. Two small syringes (3-5 ml)
 - 4. One large Luer-lock® syringe (35-50 ml)
 - 5. Flush solution
 - 6. Sterile gauze pads and tape
- B. Site Selection Proximal tibia. Palpate the landmarks and note the entry point that is the anteromedial flat surface 1-3 cm below the tibial tuberosity.
- C. Prep the surface with povidone-iodine or chlorhexidine prep and wipe dry with a sterile gauze pad.

- D. Needle Insertion
 - 1. Insert the needle at the proximal tibial site, directing the needle caudally. The needle should penetrate the skin and subcutaneous tissue and be pushed through the cortex of the bone using rotation (avoid rocking the needle) until a "pop" or "give" is felt.
 - 2. Confirm placement of the needle by:
 - a. Firm fixation of the needle and free aspiration of marrow/blood.
 - b. Infusion of 2-3 ml of NS, palpating for extravasation or noting significant resistance. If extravasation occurs, further attempts at the site should be avoided.
 - c. It is not always possible to aspirate blood/marrow, but the line may be patent.
- E. Tape and secure IO needle firmly in place.
- F. Start Infusion
 - 1. Although gravity drainage may suffice, pressurized infusions may be needed during resuscitation.
 - 2. When infusing medications via an IO route, pressure must be applied to the fluid bag in order to maintain flow rates. The EMT must continually monitor the rate of infusion.

CONTRAINDICATIONS:

- A. Suspected fracture of the bone selected for IO insertion.
- B. Prior prosthetic joint replacement involving bone selected for IO insertion.
- C. Previous significant orthopedic procedures (IO within 48 hours, surgery, etc.).
- D. Infection at the site of insertion.
- E. Excessive tissue at insertion site with the absence of landmarks.

- A. Osteomyelitis, growth plate injury (in pediatric patients), and extravasation of fluid with compression of popliteal vessels or the tibial nerve may occur.
- B. Airway and breathing should be established first in accordance with other protocols.
- C. Do not perform more than one attempt in each tibia.
- D. Any ALS medication may be administered IO.
- E. Do not give hypertonic saline through an IO line.
- F. In the event of driver failure, EZ-IO needle may be inserted manually.
- G. All EZ-IO needles are 15 gauge regardless of length.

BACKGROUND:

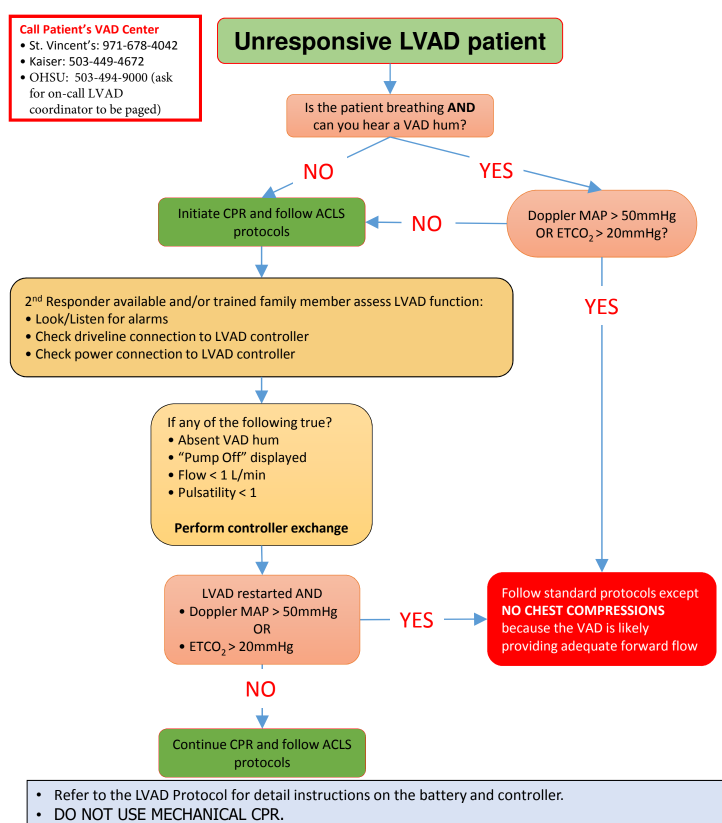
Left ventricular assist devices (LVADs) are designed to assist the pumping function of the patient's left ventricle. The HeartWare HVAD[®],HeartMate II[®], and HeartMate III[®] devices attach to the apex of the left ventricle (pump inflow) and propel blood to the ascending aorta (pump outflow). These devices utilize an external wearable system that includes a small controller connected to the internal pump by an external driveline and is powered by two batteries. They may also be "plugged in" to 110 or 12 V power, depending on the device. When managing an LVAD patient, follow these general assessment guidelines.

ASSESSING PATIENT WITH LVAD:

- A. Establish airway and provide supplemental oxygen if any respiratory signs or symptoms are present.
- B. If a patient with an LVAD is having a medical emergency, it does not necessarily mean that it is a device issue. Consider the whole clinical picture and perform a thorough patient assessment, including device function. Infection, volume depletion, stroke, bleeding, and dysrhythmias may be the cause of patient's symptoms. Most LVAD patients are anticoagulated and are at risk for bleeding complications.
- C. Auscultate heart sounds to determine if the device is functioning. Both the HeartWare HVAD[®] and HeartMate II[®], are continuous flow devices and you should hear a "whirring" sound". Because these devices diminish pulsatile flow in the circulation, peripheral pulses may not be palpable. The HeartMate III[®], although continuous flow, may provide artificial pulsatility (as well as a pulsatile hum) due to the addition of intermittent speed reduction which was designed into the device. Since this artificial pulse is not synchronized with the patient's heart rate, it may augment or diminish the native pulse. If a pulse is palpable, a BP can be attempted. Assess other signs of circulation—capillary refill, absence or presence of dizziness, temperature/ moisture of skin, end-tidal CO₂, and mental status to determine perfusion status.
- D. Standard blood pressure devices may not work. If unable to obtain a blood pressure consider using the following, if available, to estimate perfusion pressure:
 - 1. End-Tidal CO_2 Expected values should be between 35 45 mmHg.
 - 2. Doppler cuff pressure Estimates the mean arterial pressure. The goal range for Doppler MAP is > 60 and less than 90.
 - 3. Other clinical signs Capillary refill, mental status.
- E. Locate the device to identify which type is in place and follow the device specific troubleshooting guidelines. Intervene appropriately based on the type of alarm and device.
- F. Start Large Bore IV and treat with fluids as needed.
- G. Pulse oximetry may not be accurate due to the continuous flow nature of the device. You may not get an accurate reading in the field.
- H. Your cardiac monitor <u>will</u> work, and a reliable ECG may be obtained. Because the LVAD creates continuous flow independent of left heart function, not all arrhythmias will be symptomatic, including ventricular arrhythmias. If a patient requires defibrillation, leave the pump running and all components in place. The LVAD does not interfere with electrical conduction. In general, LVAD patients also have an AICD/Pacemaker. Do not place defibrillation pads directly over the pump or AICD/Pacemaker (consider

anterior/posterior placement).

- I. All ACLS medications may be administered if necessary.
- J. If suspected cardiac arrest, proceed to following flow chart:



- The 2 most common causes of pump failure are disconnection of the power and failure of the controller.
- Transport LVAD patient in circulatory arrest to the nearest VAD hospital; otherwise transport the patient to their designated VAD center.
- Patients on LVAD support frequently do not have a palpable pulse or recognizable BP yet have adequate perfusion.
- In the non-invasive assessment of the BP, use a manual BP cuff with Doppler when available, with ETCO2 as the second option.
- Assess and treat non-LVAD pathology:
 - 5 H's: Hypovolemia, hypoxia, hydrogen ion (acidosis), hypo/hyperkalemia, hypothermia
 - ➤ 5 T's: Toxins, tamponade, tension pneumothorax, thrombosis-heart, thrombosis-lung
- Keep all back-up equipment with the patient during transport!

Left Ventricular Assist Devices LVAD – 30.107

TRANSPORTING AN LVAD PATIENT:

- A. Consider transporting the LVAD patient in circulatory arrest to the nearest VAD hospital; otherwise transport the patient to their designated VAD center. <u>Call the number on the device and follow</u> <u>advice of the LVAD Coordinator on call for troubleshooting the device.</u>
- B. For all other concerns contact OLMC.
- C. The patient must be supported by battery power. <u>Remember to also transport the backup</u> <u>controller and the spare batteries.</u>
- D. The controller should be kept close to the patient, and care taken to not kink the leads.
- E. If removing or cutting patients clothing, use caution as not to sever the driveline.
- F. Do not put external pressure on any area of the LVAD system.
- G. Place gurney straps underneath the leads, and keep the batteries easily accessible.
- H. Allow the trained caregiver to ride in the transport vehicle if possible to act as an expert on the device in the absence of consciousness in the patient.
- I. Bring all of the patient's equipment.

- A. LVAD patients who are anticoagulated have a higher risk of bleeding and hemorrhage.
- B. There are no valves on an LVAD, so there is the risk of retrograde flow and stagnation of blood if the device stops, or flow is impeded.
- C. These patients are pre-load and afterload dependent, so hypovolemia can have a profound effect.
- D. If a patient is **hypertensive**, flow through the device may be reduced.

Orogastric Tube Insertion and Maintenance – 30.115

OVERVIEW:

While a patient is being ventilated with a BVM, trapped air can gather in the stomach increasing the risk of vomiting and aspiration. In addition, an enlarged stomach pushes against the diaphragm to increase intrathoracic pressure, decrease venous return, and interferes with lung ventilation.

INDICATIONS:

To alleviate gastric distention, reduce aspiration, and facilitate ventilation in intubated patients.

CONTRAINDICATIONS:

- A. Known alkali or acid ingestion.
- B. Known esophageal varices.
- C. Esophageal obstruction.
- D. Suspected epiglottitis or croup.

PROCEDURE:

- A. Assemble equipment:
 - 1. Proper size orogastric tube
 - 2. Lubricant
 - 3. 30 or 60 cc syringes
 - 4. Suction unit

Gastric Tube Size Guide			
Age	Size		
Less than 1 year	Refer to Pediatric Guide		
1 yr. to 16 yrs.	10 – 14 French		
Older than 16 yrs.	Up to 18 French		

- B. With patient's head in a neutral position measure tube length from xiphoid process to angle of jaw to corner of the mouth. Place a mark on the tube to indicate how far to advance the tube.
- C. Lubricate end of tube; about 3 4 inches.
- D. Gently insert tube and advance toward posterior oropharynx.
- E. For non-traumatic patients, repositioning the head into a slightly flexed forward position may facilitate OG tube passage past the hypopharynx and into stomach.
- F. Continue to insert tube to the measured mark). Secure tube with tape.
- G. Attach syringe to the distal end of the OG tube.
- H. Confirm tube placement by placing stethoscope over epigastrium and auscultate while inserting 30 60 ml of air in tube. You should hear gastric gurgling.
- I. Secure tube in place with tape.
- J. Place the tube to low continuous suction as needed, gastric contents should be visible in tubing.
- K. Document tube size and depth, color, consistency, and amount of gastric contents.

NOTES AND PRECAUTIONS:

- A. OG tube placement can cause bradycardia.
- B. Do not delay transport for this procedure.
- C. Monitor SpO₂ and EtCO₂ continuously.

Procedures – New 8/10/16

INDICATIONS:

Isolated non-traumatic lateral patellar dislocation.

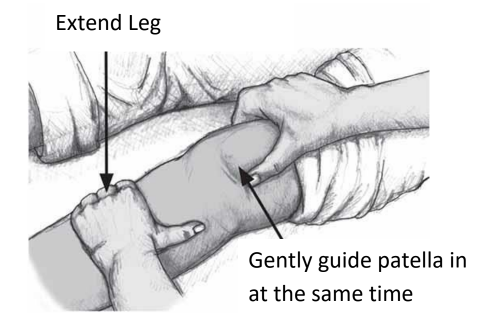
CONTRAINDICATIONS:

- A. Direct traumatic mechanism of injury (impact directly to the knee).
- B. Any sign of associated patella fracture (crepitus).
- C. Any associated injury to same extremity (femur fracture, tibia/fibula fracture, pelvic fracture).

PROCEDURE:

- A. Follow Pain Management protocol.
- B. Patient will usually present with the knee flexed and an obviously laterally displaced patella.
- C. Gently apply pressure to the lateral aspect of the patella (directing it medially) while extending the leg.

- A. Reductions should not be attempted for medial dislocations, as these commonly have associated fractures.
- B. Patients should be splinted and transported regardless of success of reduction attempt. If a patient does not want transport after successful reduction, OLMC contact is mandatory as part of the refusal process.



PURPOSE:

Escalation control of agitation or aggressive behavior should be attempted prior to physical or chemical restraint. However, if escalation control is not feasible, physical and chemical restraint should be used to protect the safety of patients and responders. Patient restraints should be utilized only when necessary and because the patient is exhibiting behavior that presents a danger to themselves and/or others based on an objective assessment of agitation and risk of violence such as the Broset Violence Assessment.

PROCEDURE:

Escalation Control Guideline:

Respect Personal Space: allowing extra physical space between you and the patient helps the patient not feel crowded or threatened and allows for provider safety. Always try to position near an exit. Do not rush in. Assess the patient's body language, listen to their tone of voice, and make a plan.

Do not go in alone: but limit the number of responders in the room or area with the patient so as not to overwhelm them. All communication with the patient should channel through one person.

Appear Calm: use a low, monotonous voice when speaking to the patient. Use good body language: avoid crossed legs/hands, hands in pockets, etc. Try to be at the patient's eye level.

Listen Without Judgement: acknowledge what the patient is saying. Confirm the legitimacy of their perceived problem, not their behavior.

Use Reflective Statements/Responses: for example, if the patient says, "I can't believe no one in my family has called me or visited" a reflective response would be, "sounds like you are frustrated and feeling unsupported by your family."

Tolerate Silences

Set Limits: it is ok to set parameters around the patient's behavior so that everyone is safe.

Give the Patient Appropriate Choices: this can help the patient regain a sense of control. For example, "Would you like to sit up on the gurney or lay down?"

Offer Optimism However Do Not Make Promises That Cannot Be Kept

Do Not Argue with the Patient: stick to the goal of escalation control. Remember they do not know you so do not take insults personally.

Physical Restraint Guidelines:

- A. Perform the Broset Violence Assessment checklist:
- B. Use the minimum level of physical restraints required to accomplish patient care and ensure safe transportation (soft restraints may be sufficient). If law enforcement or additional manpower is needed, call for assistance prior to attempting restraint procedures. Do not endanger yourself or your crew.
- C. Do not place restraints in such a way as to preclude evaluation of the patient's medical status or interfere with management of the airway.

Physical Restraint Procedure:

- A. Place the patient face up on gurney, NOT PRONE. Closely monitor the patient's respiratory status.
- B. Secure ALL extremities to gurney. Try to restrain lower extremities first using restraints around both ankles. Next, restrain the patient's arms with one arm anchored to the head of the bed and the other arm anchored to the frame of the gurney.
- C. Soft restraints are the restraint of choice. Zip ties or flex cuffs should not be used.
- D. Hand cuffed patients should be transitioned to soft restraints as soon as feasible.
- E. If it is not feasible to transition a patient out of hand cuffs, law enforcement must be present in the ambulance with a handcuff key for transport.
- F. Evaluate the patient's respiratory and cardiac status continually. Monitor SpO₂ if possible.
- G. DO NOT tighten the gurney's chest straps to the point that they restrict breathing.

Pharmacological Sedation Procedure:

- A. Document the Broset violence assessment checklist score pre and post sedation.
- B. Evaluate the number/ability of personnel needed to safely restrain the patient.
- C. Patients who are agitated but cooperative should be offered Olanzapine 10 mg SL
- D. Patients who are disruptive without danger should be treated with Midazolam 5 mg IM or Olanzapine 10 mg SL.
- E. Patients who are severely agitated, considered a danger to self or staff, who are physically violent or pose credible threats should be treated with Midazolam 10 mg IM AND Droperidol 10 mg IM or Haloperidol 10 mg IM. Be prepared to manage the patient's airway if necessary.
- F. Multiple medications cannot be given in the same syringe.
- G. Consider and treat medical and traumatic causes of combativeness-hypoxia, head injury, hypoglycemia, etc.
- H. If the patient remains a safety threat to self/others 10 minutes after the initial dose of pharmacological sedation, administer Droperidol 5 mg IM or Haloperidol 5 mg IM.
- I. Assess vital signs in first 5 minutes and at least every 10 minutes and before each additional medication if possible.
- J. Monitor patient's EKG and obtain a 12-lead if possible. Be alert for prolonged QT. If prolonged QT is identified, <u>Droperidol and Haloperidol are contraindicated</u>. Olanzapine may be administered in standard doses.

BACKGROUND:

A Peripherally Inserted Central Line (PICC) is a common method of maintaining long-term venous access in select patients. PICC lines are typically inserted into the antecubital fossa, and then threaded into central circulation. PICC lines are flushed with heparin to maintain patency and therefore it is imperative to aspirate 5 ml of blood from the line prior to use.

INDICATIONS:

- A. PICC lines may be accessed when there is a need for drug or fluid administration and traditional means of venous access are unsuccessful.
- B. Patient or patient's caregiver requests use of PICC line.

CONTRAINDICATIONS:

- A. Inability to aspirate or infuse through the catheter.
- B. Catheter located in any place other than the patient's upper arm.
- C. Need for rapid fluid resuscitation.

PROCEDURE:

- A. Use clean gloves and maintain sterility as much as possible.
- B. If there is a needleless type port on the distal end of the catheter, perform the following: (figure 1)
 - 1. Scrub the port with an alcohol pad for at least 15 seconds and allow to dry for at least 5 seconds.
 - 2. Attach a 10 ml syringe (without saline) to the port.
 - 3. Unclamp if necessary (needleless port may not have a clamp).
 - 4. Attempt to aspirate at least 5 ml of blood. Blood should draw freely. If it does not, remove the syringe and DO NOT use the catheter for access.
 - 5. If blood aspirates freely, remove the 10 ml syringe with blood and discard.
 - 6. Attach a 10 ml syringe with NS and gently flush the line. Never use a smaller syringe. If line does not flush, remove the syringe and DO NOT use the catheter for access.
 - 7. If line flushes, remove the syringe and attach the catheter to the end of the IV tubing and begin infusion of NS or LR. Adjust the rate to the needs of the patient within the limits of the catheter.
 - 8. Administer medications though IV tubing port if indicated.
- C. If there is a capped needle-type port on the distal end of the catheter, perform the following: *(figure 2)*
 - 1. Scrub the cap with an alcohol pad for at least 15 seconds and allow to dry for at least 5 seconds.
 - 2. Clamp the catheter tubing using ONLY the existing clamp on the catheter and then remove the cap. <u>Never allow a central line to be open to air.</u>
 - 3. Attach a 10 ml syringe on the catheter end.
 - 4. Unclamp the catheter.
 - 5. Attempt to aspirate at least 5 ml of blood. Blood should draw freely. If it does not, reclamp the line and remove the syringe. DO NOT use the catheter for access.
 - 6. If blood aspirates freely, clamp the catheter again.
 - 7. Remove the 10 ml syringe with blood and discard.

PICC Line Access – 30.140

- 8. Attach a 10 ml syringe with NS.
- 9. Unclamp and gently flush the line. Never use a smaller syringe. If line does not flush, re-clamp the line and remove the syringe. DO NOT use the catheter for access.
- 10. If line flushes, re-clamp and remove the syringe.
- 11. Attach the catheter to the end of the IV tubing.
- 12. Unclamp the catheter and begin infusion of NS or LR. Adjust the rate according to the needs of the patient within the limits of the catheter.
- 13. Administer medications though IV tubing port if indicated.

- A. <u>Do not administer medications, flush, or aspirate with less than a 10-cc syringe.</u> <u>Smaller</u> <u>size syringes generate too much pressure and can damage the catheter.</u>
- B. Do not attempt to reinject aspirated blood as it may contain clots.
- C. The maximum flow rates for a PICC line is 125 ml/hr for less than size 2.0 French, and 250 ml/hr for catheters over 2.0 size French.
- D. Keep patient's arm straight to avoid kinking the PICC line and obstructing flow.
- E. Ensure all line connections are secure.
- F. PICC lines access the patient's central circulation and the risk of infection is high. Avoid contamination to ports and connections while accessing.
- G. Do not administer the following medications through a PICC line:
 - 1. <u>Adenosine</u> The line may rupture during rapid infusion due to over pressurization.
 - 2. Dextrose 50% The catheter can be damaged due to the viscosity of the fluid

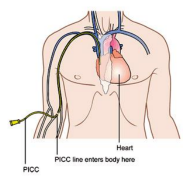




Figure 1- Needleless port

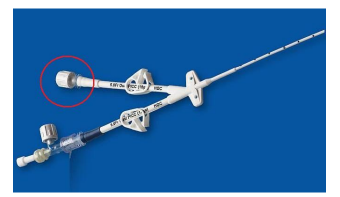


Figure 2 – Non-needleless type port with cap

Positive End-Expiratory Pressure (PEEP)– 30.145

DEFINITION:

Positive end-expiratory pressure (PEEP) is a method of ventilation in which airway pressure is maintained above atmospheric pressure at the end of exhalation by means of a mechanical impedance (PEEP valve). PEEP has some similarity to CPAP although it is delivered through bag instead of a facemask. It can be delivered via bag-valve-mask or bagging into an endotracheal tube. At the end of exhalation PEEP prevents alveolar collapse (i.e. the alveoli stay open) and improves oxygen exchange across the alveolar membrane. Additionally, PEEP may recruit more alveoli that have collapsed, which may further improve oxygenation. **ADDING PEEP IS DONE TO IMPROVE OXYGENATION.** The disadvantage to PEEP is that it may increase intrathoracic pressure, which may reduce blood flow in cardiac arrest or a shock state.

INDICATIONS:

Hypoxia, either prior to or post intubation despite appropriate bag ventilation with 100% oxygen.

CONTRAINDICATIONS:

- A. Cardiac arrest (absolute).
- B. Hypotension or shock state (relative). May still choose to apply PEEP when preparing to RSI a hypoxic/hypotensive patient.

PROCEDURE:

- A. If not already applied, apply PEEP valve to bag device.
- B. Dial PEEP value to 5cm H_2O and bag per usual.
- C. Increase PEEP by 5cm H₂O every 3-5 minutes until hypoxia resolves (oxygen saturation > 95%).
- D. Maximum PEEP is $15 \text{ cm H}_2\text{O}$.

- A. Increasing bagging rate will not necessarily improve oxygenation but can cause hyperventilation, which can be detrimental to patients.
- B. PEEP valve may come out of the package set to five or zero. Be aware of valve settings.
- C. Maximum PEEP in pediatrics is 5cm H₂O.

DEFINITION:

To provide direction on the safe removal of protective sports equipment that includes helmet and shoulder pads. <u>This procedure page uses football gear as an example, but these guidelines can be used with other sports equipment as well.</u>

PROCEDURE:

A. Initial Evaluation

- 1. The initial evaluation should begin by assessing level of consciousness, breathing, and circulation. If the athlete is breathing and stable, but a neck injury is suspected, a quick sensory and motor nerve exam should be initiated.
- 2. After the quick neurological exam on a stable athlete, the facemask should always be removed.

B. Face Mask Removal

- 1. Stabilize the head.
- 2. Release side attachments first by quick release or screwdriver. Second unscrew top loops to remove facemask. Cutting should be the last resort if quick release or screwdriver does not work.
- 3. Quick release face masks are also in use and found on newer helmets. One popular device looks like a "rivet" instead of a screw (pictured below). The release mechanism can be activated by pressing it down with a pen or tip of a screwdriver.
- 4. Athletic trainers and coaching staff are familiar with this and can provide assistance.



C. General equipment removal guidelines:

- 1. Equipment should be removed on the field if an athletic trainer and/or 3+ individuals trained in technique are present. If no athletic trainer is present, and individuals are not comfortable with removal, leave gear intact, but attempt to remove facemask should be performed for airway access.
- 2. Equipment removal should be performed by at least three trained and experienced rescuers.
- 3. If removing equipment, always remove the helmet and the shoulder pads, never just one or the other. Leaving the helmet on or just the shoulder pads on by itself creates head, neck, or spinal cord flexion.

D. Removal of helmet and shoulder pads as a unit:

- 1. Gear removal starts from the head and proceeds down the body.
- 2. Remove the helmet first and then remove the shoulder pads, and leg gear. **Do not start with the shoulder pads.**
- 3. Cut chin straps.
- 4. Remove ear pad or deflate pad bladder.
- 5. Use a two-person technique to remove the helmet.
 - a. Person at the top firmly holds manual c-spine at the top using two hands to stabilize the patient's helmet.
 - b. The other responder, starting at the chin, slides his or her hands inside the patient's helmet "firmly" gripping the head and sliding their hands inside the helmet.
 - c. Responders transition manual c-spine responsibility from the person at the top of the head/ helmet to the person supporting the patients head from underneath.
 - d. Firm control of the head and neck is the goal. The person at the top proceeds to remove the helmet off the patient's head in a coordinated and smooth manner. **DO NOT SPREAD APART SIDES OF HELMET.**
 - e. Once helmet is removed, the person at the top of the head resumes manual c-spine until full c-spine precautions are in place.
- 6. Cut shoulder pad straps.
- 7. Cut both the jersey and shirt up sleeves towards midline of body.
- 8. Person at head stabilizes maxilla and occiput and gives commands.
- 9. Position three people on each side, with one stabilizing the head. Another person removes the equipment as a unit.

While backboard and straps are being prepared:

E. Chest access:

- 1. Cut jersey and front laces of shoulder pads.
- 2. Flip out shoulder pads. Some newer systems allow the shoulder pads to come apart prior to removal. Athletic trainers and coaching staff are familiar with these systems and can provide assistance.
- 3. Place hands on shoulders with thumbs grasping the clavicle and fingers surrounding the upper trapezius muscles.
- 4. Secure the athlete's head between the responder's forearms.

F. Backboard utilization:

- 1. If an athletic trainer is present an 8-person lift and slide technique is preferred as it causes the least amount of cervical movement. If no athletic trainer is present and the athlete is too big for lift and slide, a log roll technique will be performed.
- 2. The person at head initiates commands and oversees proper placement and techniques.
- 3. Position three responders on each side of body; one at shoulders, one at hips, and one at legs.
- 4. One other person is in charge of the backboard and slides it into place.
- 5. If the helmet is not resting on board, padding can be added to fill space.
- 6. Fasten straps and tape helmet to board.
- 7. Chinstrap remains in place unless it interferes with airway.
- 8. Recheck sensory and motor nerve vitals for changes and document.
- 9. If C-Spine injury is suspected with neurological deficits, spine board should be utilized in route to the hospital.
- 10. If athlete is sitting or standing, a c-collar can be utilized, and athlete can be carefully placed on the gurney.

NOTES & PRECAUTIONS:

Athletic Trainers and coaching staff are subject matter experts when it comes to the gear regardless of the sport. Collaborate with them early and often.

INDICATIONS:

When patient is exhibiting respiratory difficulty secondary to secretions in airway or the potential for aspiration exists.

PROCEDURE:

- A. Oral Suctioning
 - 1. Pre-oxygenate patient with 100% oxygen.
 - 2. Assemble equipment: Suction unit with tonsil tip or dental tip, personal protective equipment (gloves, goggles, gown).
 - 3. Attach required monitoring equipment.
 - 4. Turn suction unit on and confirm mechanical suction is present.
 - 5. Insert tip without suction.
 - 6. Cover thumbhole to begin suction if using a tip other than dental tip.
 - 7. Apply suction for < 15 seconds.
 - 8. Monitor patient's oxygen saturation.
 - 9. Re-oxygenate patient for at least 2 3 minutes between suctioning attempts.

B. <u>Tracheal Suctioning</u>

- 1. Pre-oxygenate patient with 100% oxygen.
- 2. Assemble equipment: Suction unit, correct size suction catheter, sterile rinse, personal protective equipment (gloves, goggles, gown).
- 3. Attach required monitoring equipment.
- 4. If patient is being ventilated with BVM through an endotracheal tube prior to suctioning, have someone else remove the bag from end of ET tube prior to suction attempt.
- 5. Insert catheter into the ET tube without applying suction.
- 6. Advance catheter as far as possible.
- 7. Withdraw slowly using intermittent suctioning while rotating catheter.
- 8. Do not suction more than 15 seconds.
- 9. Monitor patient's oxygen saturation.
- 10. Rinse catheter in sterile saline.
- 11. Re-oxygenate patient for at least 2 3 minutes between suction attempts.
- C. Suctioning with Meconium Aspirator

Tracheal suctioning is not indicated in the vigorous infant born with meconium stained fluid, whatever the consistency. Simply use a bulb syringe or large bore catheter to clear secretions from the mouth and nose as needed.

- 1. Assemble equipment: Suction unit, appropriate size ET tube, personal protective equipment (gloves, goggles, gown).
- 2. Attach required monitoring equipment.
- 3. Turn suction unit on and confirm mechanical suction is present.
- 4. After infant has been intubated, attach meconium aspirator to end of ET tube.
- 5. Cover thumbhole to begin suctioning while slowly withdrawing the ET tube. Do not suction for more than 15 seconds.
- 6. Monitor patient's oxygen saturation and heart rate and stop if patient becomes bradycardic.
- 7. Re-oxygenate patient for at least 2 3 minutes between suctioning attempts.

- 8. If patient has not been intubated and meconium is thick, at the least, aggressive oropharyngeal suctioning should be carried out with the largest diameter suction device available.
- D. Suctioning with Nasal Aspirator Device
 - 1. Assemble equipment: Bulb syringe, suction unit with nasal aspirator, personal protective equipment.
 - 2. If nasal secretions are thick consider instilling 1-4 drops of NS into nares to loosen prior to suctioning.
 - 3. If using electric suction be sure vacuum is set less than 100 mmHg.
 - 4. Gently place device tip into nostril. Avoid placing against inside walls of nostril.
 - 5. Apply suction (< 15 seconds if using electric suction)
 - 6. Repeat as needed

- A. Oral and tracheal suctioning can cause trauma to the oropharynx and airway, bradycardia, or hypoxia. It should not delay other resuscitation.
- B. Suction pressure should be set as low as possible and yet effectively clear secretions. Negative pressure of less than 80-100 mmHg in neonates and less than 150 mmHg in adults are recommended.
- C. When suctioning the intubated patient, the diameter of the suction catheter should not exceed one half of the internal diameter of the endotracheal tube.

INDICATIONS:

TASER® barbs should be removed at the request of law enforcement if:

- A. The patient has been adequately subdued so as not to pose a danger to Fire/EMS personnel. AND,
- B. The barbs are not embedded in the face, neck, or groin areas.

PROCEDURE:

- A. Perform patient assessment.
- B. Monitor vital signs and LOC. Ensure that vital signs are in the normal limits for the situation.
- C. Expose the area where TASER[®] barb has implanted under the skin.
- D. Cut wires from the barb if still attached.
- E. Place thumb and forefinger above and below the barb parallel to the portion of the shaft implanted in the patient's skin.
- F. Spread your thumb and forefinger apart to stretch the skin tightly over the barb.
- G. Holding tension, use needle-nose pliers (or similar tool) with gripping strength and grasp the end of the barb protruding out of the skin near the wire lead and firmly pull out the barb with one quick jerking motion.
- H. If probe removal tool is available (see TASER[®] 7 picture below)
 - 1. Place one hand on the patient in the area where the probe is embedded and stabilize the skin surrounding the puncture site.
 - 2. Slide the safety clip notch between the probe and the subject, catching the probe between the dart body and the dart point.
 - 3. In one uninterrupted motion, pull the safety clip, and probe with it, straight out of the puncture site maintaining a 90-degree angle to the skin (avoid twisting or bending the probe).
- I. Assess the skin where the barb was removed. The skin should be cauterized from the electrical current. Dress the wound to prevent infection.
- J. Contact OLMC if unsure whether to transport.

- A. Patients should be in police custody and monitored by police for the safety of medical personnel.
- B. Do not remove TASER[®] Barbs from the face, neck, or groin area. Stabilize the barbs and transport to the Emergency Department.
- C. TASERS[®] emit two barbs. Make sure both are removed. Treat all barbs as a biohazard and dispose as you would any other sharps. Some law enforcement agencies may direct you to place the probe back into the cartridge as evidence.
- D. Potential trauma may have occurred before (during a struggle) or after the patient was hit by the TASER[®] (e.g., patient falls and hits head).
- E. Consider whether the patient meets criteria for Altered Mental Status or Poisonings and Overdoses protocols.
- F. CAUTION: Where barbs have wires still connected to the TASER[®] Gun, shock can still be delivered.



Tension Pneumothorax Decompression – 30.170

DEFINITION:

The emergency decompression of a tension pneumothorax using an over-the-needle catheter.

INDICATIONS:

To warrant chest decompression in the field, the patient must be <u>significantly</u> <u>symptomatic or</u> <u>in extremis (at risk of death)</u> with:

- A. High clinical suspicion and;
- B. Progressive respiratory distress and;
- C. Shock symptoms with low or rapidly decreasing blood pressure.

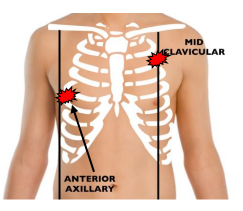
and at least one of the following:

- A. Decreased or absent breath sounds.
- B. Consistent history (i.e., chest trauma, COPD, asthma).
- C. Distended neck veins.
- D. Tracheal shift away from affected side (late sign).
- E. Asymmetrical movement on inspiration.
- F. Hyper-expanded chest on affected side.
- G. Drum-like percussion on affected side.
- H. Increased resistance to positive pressure ventilation, especially if intubated.

EMS witnessed traumatic arrest patients with abdominal or chest trauma for whom resuscitation is indicated should have bilateral chest decompression performed even in the absence of the above signs.

PROCEDURE:

- A. Expose the entire chest.
- B. Establish landmarks:
 - 1. Anterior 2nd intercostal mid clavicular or if unavailable.
 - 2. Lateral 4th intercostal space anterior axillary (above nipple).
- C. Clean chest vigorously with appropriate antiseptic.
- D. On affected side, locate the landmark and insert a large gauge over-the-needle catheter with syringe attached along <u>the superior margin</u> of the rib below (e.g. top of third rib to enter second intercostal space).
- E. If the air is under tension, the barrel will pull easily and "pop" out of the syringe.
- F. Remove syringe, advance catheter, and remove needle.
- G. Secure from movement.



- A. Patient's chest should be auscultated often for return of tension or other respiratory complications.
- B. Tension pneumothorax is a rare condition, but can occur with trauma, spontaneously, or as a complication of intubation. Tension takes time to develop, but forceful positive ventilation may increase the rate of development.
- C. Simple or non-tension pneumothorax is not life threatening and should not be decompressed in the field.
- D. The ideal decompression catheter length is three inches.
- E. Possible complications:
 - 1. Creation of pneumothorax if none existed previously.
 - 2. Laceration of lung or pericardium. Stop needle advancement once it has popped through the pleura and advance the catheter only.
 - 3. Laceration of blood vessels. (Always slide the needle above the rib).
 - 4. Infection. Clean rapidly but vigorously; use sterile gloves if possible.
- F. Tension pneumothorax can be precipitated by the occlusion of an open chest wound. If the patient deteriorates after dressing an open chest wound, remove the dressing.

DEFINITION:

Placement of a circumferential band around a limb to occlude arterial blood flow distal to the band.

INDICATIONS:

Extremity hemorrhage that is uncontrollable by less aggressive means (direct pressure, bandaging, or pressure dressing) OR a wound that could cause life threatening extremity hemorrhage during an ongoing tactical problem (e.g. potential building collapse, mass casualty event, amputation).

PROCEDURE:

- A. Fully expose and evaluate the wound.
- B. Apply tourniquet directly to the skin, 2 3 inches proximal to the most proximal limb wound, not over a joint.
- C. Tighten until all bleeding stops and no distal pulse is palpable.
- D. Secure the windlass per manufacturer instructions.
- E. If one properly placed tourniquet does not control bleeding, a second should be placed proximal to the first and tightened appropriately.
- F. Endeavor to keep all tourniquets exposed.
- G. Mark with time of application and communicate this to receiving providers.
- H. Re-evaluate tourniquets frequently to ensure they have not loosened.

- A. If an improvised tourniquet is present before medical provider arrival, place a commercial tourniquet per protocol and remove the improvised tourniquet if operationally feasible.
- B. Properly applied tourniquets will rarely damage tissue if removed within two hours.
- C. If unable to fully expose a limb and identify all wounds on that limb place the tourniquet as high on the limb as possible. Once all wounds on that limb can be identified, every effort should be made to move the tourniquet to 2 3 inches proximal to the most proximal wounds, and not on a joint.
- D. Intermittently loosening and tightening a tourniquet to "reperfuse" a limb is of no benefit and dangerous as it encourages additional bleeding.
- E. A single commercially available tourniquet completely occludes femoral artery blood flow about 70% of the time. Two tourniquets placed side by side completely occlude about 80% of the time.
- F. The ability of the tourniquet to completely occlude arterial flow is dependent on limb circumference. Larger limbs are more difficult to occlude.
- G. A persistent pulse, continued venous congestion / distention, re-bleeding after initial hemorrhage control, and expanding hematoma are all indications of an ineffective tourniquet.
- H. Clothing, padding under the tourniquet, and limb movement all cause tourniquets to loosen over time and should be avoided.
- I. Tourniquets can cause significant pain and may require narcotics for pain control.
- J. Proper placement of a CAT® tourniquet on a lower extremity requires threading the circumferential band through both slits of the buckle.
- K. Proper placement of the SOFTT tourniquet requires tightening the knurled screw on the buckle before tightening the windlass.

DEFINITION:

Transcutaneous pacing is the technique of electronic cardiac pacing accomplished by using skin electrodes to pass repetitive electrical impulses through the thorax.

INDICATIONS:

Transcutaneous pacing should be considered in bradycardia with evidence of inadequate perfusion, (e.g. altered mental status, chest pain, hypotension, other signs of shock).

PROCEDURE:

- A. Ensure ECG pads are attached, and monitor displays a rhythm.
- B. Attach pacing electrodes to anterior and posterior chest just to the left of the sternum and spinal column, respectively. Alternatively, pads may be placed in the standard anterior and lateral position as with defibrillation. If there is difficulty in obtaining capture, try alternative position.
- C. Begin pacing at a heart rate of 80 beats per minute and 30 mA current output.
- D. Increase current by increments of 10mAs while observing monitor for evidence of electrical capture. Confirm mechanical capture by checking pulses and BP.
- E. If patient is comfortable at this point, continue pacing. If patient is experiencing discomfort, consider analgesia per pain management protocol and/or sedation with a benzodiazepine per appropriate medication protocol if blood pressure allows.
- F. If the patient remains unconscious during pacing, assess capture by observing the monitor and evaluating pulse and blood pressure changes. In the event of electrical capture and no pulses, follow PEA protocol.
- G. If there is no response to pacing <u>and</u> drugs, consult with OLMC. If a change in pacing rate is desired, contact OLMC.

PEDIATRIC PATIENTS:

Use above guidelines except:

- A. Use anterior/posterior pad placement first for patients less than 1 year.
- B. Begin pacing at smallest mA output.
- C. Increase current in increments of 10 mA while observing monitor for evidence of electrical capture.
- D. Confirm mechanical capture by checking pulses and BP.
- E. Contact OLMC for adjustments to rate based on age and response to pacing.

NOTES & PRECAUTIONS:

Transcutaneous pacing should not be used in the following settings:

- A. Asystole.
- B. Patients meeting Death In The Field criteria.
- C. Patients in traumatic cardiac arrest.

Operations

PURPOSE:

Law enforcement agencies stress that their first priority on any crime scene is the preservation of life with reconstruction of the crime scene second. EMS personnel can be of assistance by adhering to the following guidelines regarding crime scene response.

PROCEDURE:

- A. <u>Response and Arrival</u>
 - 1. Be conscious of physical and weather conditions around the site. Tire tracks of suspect vehicles are often located in or adjacent to a driveway.
 - 2. Limit the number of personnel allowed onto the scene. Consult with police on the scene to direct placement of vehicles and route of personnel onto the scene.

B. Access and Treatment

- 1. Select a single route to the victim. Maintaining a single route decreases the chance of altering or destroying evidence or tracking blood over a suspect's footprints.
- 2. Note the location of furniture, weapons, and other articles, and avoid disturbing them. If they need to be moved, someone should note the location the article was moved from, by whom it was moved, and where it was placed.
- 3. Remove from the scene all EMS generated debris that is contaminated with blood or body fluid and dispose of through established channels.
- 4. Be conscious of any statements made by the victim or other persons at the crime scene. Write down what these statements were and report to the investigating officers.
- 5. Note the specific garments worn by the patient at the time of treatment. It is also important not to tear the clothing off or cut through any holes, whether made by a knife, bullet, or other object.
- 6. The victim should be placed on a clean sheet when ready for transport. At the hospital, please try to obtain the sheet once the victim is moved off it. Fold it carefully in on itself and give it to the investigating officers. This is especially important in close contact crimes such as rape, serious assault, and death cases.

C. Documentation

- 1. A detailed report is important in case you are later called to testify in court. An incident report should be completed and should cover your observations, conversations with family or witnesses, location of response vehicles and equipment, furniture, weapons, clothing that has been moved, items that were handled, and your route to the victim.
- 2. An Unusual/Supplemental Event Report may be helpful for you to complete. This is a protected document and if you are called to court may be used by you to refresh your memory of aspects of the call that are not included in the Patient Care Report.
- 3. Do not offer your opinions or evaluations about the crime scene.

REMINDER:

Any location can be, or become, a crime scene. When responding, and upon arrival, if something does not appear to be right, notify police. If you suspect a crime scene and police are not present, secure area and document what you see.

PURPOSE:

To define under what conditions treatment can be withheld or stopped.

PROCEDURE:

A. DEATH IN THE FIELD

Resuscitation efforts may be withheld if:

- 1. The patient has a Do Not Attempt Resuscitation (DNAR)/Do Not Resuscitate (DNR) order.
- 2. The patient is pulseless and apneic in a mass casualty incident or multiple patient scene where the resources of the system are required for the stabilization of living patients.
- 3. The patient is decapitated.
- 4. The patient has rigor mortis in a warm environment.
- 5. The patient is in the stages of decomposition.
- 6. The patient has skin discoloration in dependent body parts (dependent lividity).

Medical Cardiac Arrest:

- 1. If the initial ECG shows asystole or agonal rhythm confirmed in 3 leads, and the patient, in the responder's best judgment would not benefit from resuscitation:
 - a. The PIC may determine death in the field, OR
 - b. Begin BLS procedures, and contact OLMC with available patient history, current condition, and with a request for advice regarding discontinuing resuscitation.
- 2. If after the airway is established and the asystole protocol has been exhausted the patient persists in asystole (confirmed in 3 leads) the PIC may determine the patient to be dead in the field.
- 3. Death in the field may be determined with EtCO₂ of 10 or less in patients with PEA after 30 minutes of ACLS resuscitation. For patients with EtCO₂ greater than 10 either continue resuscitation or contact OLMC to stop resuscitation.
- 4. Patients in VF should be treated and transported.

Traumatic Cardiac Arrest:

- 1. Traumatic arrest carries high rates of mortality, but improved outcomes have been seen in EMS witnessed arrest. Causes of arrest that may be amenable to prehospital resuscitation include severe hypovolemia, hypoxia, and tension pneumothorax.
- 2. A cardiac monitor may be beneficial in determining death in the field.
- 3. Trauma patients who have arrested prior to EMS arrival can by declared dead in the field.
- 4. Witnessed traumatic arrest patients and patients who deteriorate to PEA or asystole may benefit from "HAT" resuscitation. Follow the Traumatic Cardiac Arrest Protocol (10.050).

Death and Dying – 50.025

Pediatric Non-Traumatic Cardiac Arrest:

- 1. Death in the field may be determined **<u>if all</u>** the following conditions are met:
 - a. Patient remained in asystole for duration of resuscitation. If at any time there was PEA or a shockable rhythm, they should be transported.
 - b. Resuscitation has been ongoing for at least 30 minutes.
 - c. At least 3 doses of epinephrine have been administered.
 - d. There is adequate safety/support on scene.
- 2. In unclear circumstances, call OLMC or initiate transport.

Notes & Precautions:

- 1. ORS allows a layperson, EMT or paramedic to determine "Death in the Field".
- 2. Consult OLMC with any doubt about the resuscitation potential of the patient.
- 3. A person who was pulseless or apneic and has received CPR and has been resuscitated is not precluded from later being a candidate for solid organ donation.

B. POLST ORDERS AND DECISION MAKING

- 1. In the pulseless and apneic patient who <u>does not meet</u> DEATH IN THE FIELD criteria but is suspected to be a candidate for withholding resuscitation, begin CPR and contact OLMC.
- 2. A patient with decision-making capacity or the legally authorized representative has the right to direct his or her own medical care and can change or rescind previous directives.
- 3. EMS providers may honor a DNAR/DNR order signed by a physician, nurse practitioner or physician assistant. DNAR/DNR orders apply only to the patient in cardiopulmonary arrest and do not indicate the types of treatment that a person not in arrest should receive. POLST was developed to convey orders in other circumstances.
- 4. Portable Orders for Life-Sustaining Treatment (POLST):
 - The POLST was developed to document and communicate patient treatment preferences across treatment settings. While these forms are most often used to limit care, they may also indicate that the patient wants everything medically appropriate done. **Read the form carefully!** When signed by an allopathic physician (MD or DO), naturopathic physician, nurse practitioner, or physician assistant, POLST is a medical order and EMS providers are directed to honor it in their Scope of Practice unless they have reason to doubt the validity of the orders or the patient with decision-making capacity requests change. If there are questions regarding the validity or enforceability of the health care instruction, begin BLS treatment and contact OLMC [OAR 847-035-030 (7)] If the POLST is not immediately available, a POLST form as documented in the Electronic POLST registry hosted at MRH (503-494-7333) may also be honored.
 - Section A: <u>Applies only when patient is in cardiopulmonary arrest.</u>
 - Section B: Applies in all other circumstances.
 - For a POLST form to be valid it must include:
 - i. Patient's name
 - ii. Date signed (forms do not expire)
 - iii. Health care professional's signature (patient signature is optional)

- Consider providing pain/symptom management and not transporting patient if they are Comfort Measures Only, the symptoms can be managed, and the patient and caregivers on scene do not want transport to the hospital. Consider OLMC contact for advice.
- 5. The legally authorized representative may make decisions for the patient who is unable to make medical decisions. However, when in doubt or for unresolved conflict on the scene contact OLMC. The order is:
 - a. A legal guardian
 - b. A power of attorney for health care as designated by the patient on the Oregon Advance Directive
 - c. Spouse or legal domestic partner
 - d. Adult children
 - e. Parent
- 6. <u>Death with Dignity</u>:

If a person who is terminally ill and appears to have ingested medication under the provisions of the Oregon Death with Dignity Act, the EMS provider should:

- a. Provide comfort care as indicated.
- b. Determine who called 9-1-1 and why (i.e., to control symptoms or because the person no longer wishes to end their life with medications).
- c. Establish the presence of DNAR/DNR orders and/or documentation that this was an action under the provisions of the Death with Dignity Act.
- d. Contact OLMC.
- e. Withhold resuscitation if DNAR/DNR orders are present, <u>and</u> there is evidence that this is within the provisions of the Death with Dignity Act <u>and</u> OLMC agrees.

C. PATIENTS ENROLLED IN HOSPICE AND DYING PATIENTS

- 1. Look for POLST forms (contact Registry if needed) and attempt to honor patient preferences. Always provide comfort measures.
- 2. If the patient is enrolled in hospice or receiving palliative care, refer to Hospice and Palliative Care protocol 50.062.

D. CARE OF GRIEVING PERSONS

Resuscitation phase:

- 1. As time allows, give accurate and truthful updates about the patient's prognosis. If available, assign one person to interact with and support family members.
- 2. Consider gently removing children from the resuscitation area.
- 3. Depending upon the emotional state of family members, consider allowing them to watch and/or participate in a limited and appropriate way.
- 4. If family or friends were doing CPR prior to your arrival, commend their efforts.

- 5. If family or friends are disruptive consider removing them or try assigning simple tasks, such as helping bring in the stretcher, holding doors open, telling other family about the event, and calling the doctor or clergy member.
- 6. Be respectful. Make requests. Don't give orders.

Once death is determined:

- 1. Treat the recently dead with respect.
- 2. Tell family and friends of the death honestly. Use the words "death" or "dead". Avoid using euphemisms such as "passed away" or "gone".
- 3. Avoid using past tense terms when speaking to survivors of the recently dead.
- 4. Allow family and friends to express their emotions. Listen to them if they want to talk but don't push them.
- 5. Give factual information.
- 6. Genuine warmth and compassion will be more helpful than almost anything else for survivors. Don't feel it necessary to say the "right" things. Listening often provides grieving people with the most comfort.

Focusing on survivors:

- 1. See to it that survivors have a support system present before you leave. Consider calling TIP through EMS Dispatch, if available in your jurisdiction. Call friends, family, clergy, or neighbors to be with them. Respect the survivor's wishes to be alone.
- 2. Explain the next steps to them after you have pronounced death. This will include the police coming to make reports, possibly the medical examiner, and the possible need for an autopsy.
- 3. Contact the Medical Examiner's office as soon as possible before moving or altering the body.
- 4. Allow family and friends to say their good-byes if possible.
- 5. A chaplain may be helpful in assisting with survivors. It is advisable to call early, as the chaplains do not have code-3 capabilities.
- 6. Help survivors make decisions such as which people should be called. If they ask you to make calls, try to comply, mention the need to find a funeral home, if one has not already been chosen. Clergy may also be helpful with this decision.

E. DEATH OF A CHILD:

- 1. Do not accuse the parents of abuse or neglect but take careful note of the patient's surroundings and the general physical condition of the child.
- 2. Do not be overly silent, which may imply guilt to the parents.
- 3. Ask the parents only necessary questions and do not judge or evaluate them. Do not tell them what they "should have" been doing before your arrival.
- 4. Remind parents to arrange for childcare of other children.
- 5. Listen carefully to their statements and answer only with accurate information.
- 6. If there is a police investigation, tell the parents that this is routine.
- 7. Successful management of child deaths requires supportive, compassionate, and tactful measures.

PURPOSE:

To establish guidelines for the handling of the body and required notification following a declaration of death as outlined in ORS Chapter 146. The goal of an investigation by the medical examiner's office is to determine the cause and manner of death.

PROCEDURE:

- A. If the patient appears to meet obvious death in the field criteria, have only one person enter the scene to verify death; limit access if possible. Don't move the decedent unless necessary. Document anything that was altered by your examination (e.g. unbuttoned/removed clothing, movement of the decedent, etc.).
- B. Contact police for all deaths in the field except for hospice and skilled nursing facilities.
- C. Upon declaration of death, the medical examiner (ME) must be contacted. Until contact is made with the ME:
 - 1. Do not move the body.
 - 2. Do not cover the body unless necessary (outside, public place). If covering the body is necessary, use a new/clean non-cloth disposable sheet or blanket such as an emergency blanket.
 - 3. Do not remove clothing or cleanse the body or otherwise alter the appearance of the state of the body.
 - 4. Do not remove any of the effects of the deceased or instruments or weapons related to the death.
 - 5. Do not let anyone in the area where the deceased is located.
 - 6. If resuscitation was attempted, do not remove IV's, advanced airways, or defib/ECG pads. Circle all IV attempts or any trauma or marks that you caused to the body with an ink pen if possible.
- D. Depending on the circumstances, the ME will either respond to the scene for a full investigation or release the body to a funeral home with a limited investigation. Generally, it is best to turn the scene over to law enforcement once you have given a report.
- E. You should not leave the scene without passing the scene off to law enforcement or until the ME has released you over the phone or the ME arrives at the scene and has released you.
- F. The following documentation is required for declaration of death calls:
 - 1. Location and position the body was found.
 - 2. Location of evidence if moved for safety concerns (gun, knife, bat, etc.).
 - 3. Anything suspicious (e.g. bruises on the body, deformed arm, black eye, comments made by bystanders/relatives/friends, etc.).
 - 4. Name and title of individual the scene is turned over to (law enforcement, ME, another crew) and the disposition of the body.
 - 5. The name of the ME if the body is released with a limited investigation.
 - 6. Follow your individual agency's medical records policy for listing witnesses or possible witnesses with contact information.

NOTES:

- A. Once the person is declared dead, your jurisdiction ends. Even law enforcement is not allowed to touch or move the body. Only the ME, Deputy ME (also referred to as a Medicolegal Death Investigator), or District Attorney, has lawful authority over the body. Any of these individuals can grant access or removal of the body.
- B. Not all deaths are under the jurisdiction of the ME (e.g. patient on hospice care longer than 24 hours, patient who dies in a skilled nursing facility). However, EMS calls should be considered an ME case and reported to the ME. It is best to let the ME decide if this is their case or not.
- C. Your chart may be read by the ME's office and if read, will become part of the report for cause and manner of death.
- D. In smaller counties and jurisdictions, law enforcement officers may be appointed as Deputy ME's or medicolegal Death Investigators, who under the direction of the ME's office, can investigate deaths and authorize the removal of a body of a deceased person from the apparent place of death.
- E. If you suspect a COVID-19 death, document the names and contact information of everyone who had contact with the person that is on scene.
- F. The following information should be available, if possible, prior to contacting the ME. The ME may not ask for all this information but be ready with this information.

•	Your name	•	Any evidence of drug use
•	Unit number	•	Name of deceased
•	What you were dispatched on	•	Address of deceased
•	How you found the patient	•	Age of deceased
•	Brief description of your actions	•	Gender of deceased
•	Whether you suspect foul play	•	Medical history
•	Whether death occurred at work	•	Medications
٠	Whether death occurred while in custody	•	Primary caregiver and phone number
٠	Whether death was the result of a crime	•	Family contact
•	Whether death was unattended	•	Funeral home
•	Whether cause of death might be from a contagious disease		

PROCEDURE:

- A. A patient care report shall be generated for each identified patient and shall be completed on an approved State EMS patient care form.
- B. Documentation shall include, at least:
 - 1. The patient's presenting problem.
 - 2. Vital signs with times.
 - 3. History and physical findings as directed by individual protocols.
 - 4. Treatment(s) provided, and time(s).
 - 5. If monitored, ECG strip, 12-lead ECG, and interpretation.
 - 6. Any change in the condition of the patient.
 - 7. OLMC contact:
 - a. Include physician name
 - b. Time of contact
 - c. Orders received from physician
- C. An electronic Prehospital Care Report must be submitted to a hospital or facility receiving the patient with 24 hours of the patient being transported per ORS 333-250-0310.
- D. If a patient refuses treatment and/or transport, refer to Refusal and Informed Consent protocol.

PURPOSE:

The transfer of care is an activity that has the potential for medical error. Patient hand-off reports between either EMS personnel on scene or between EMS personnel and hospital staff during transfer of care, needs to be delivered in a consistent and clear format to ensure accuracy and completeness of information. As many agencies are transitioning to paperless in-field reporting, the passage of detailed information from one agency to another or to the hospital becomes critically important.

PROCEDURE:

The following "DMIST" format is a guideline for both oral and/or written communications when passing information from one agency to the next as well as for reports to receiving facilities. It is understood that not all information may be available at the time of the handoff.

DEMOGRAPHICS:

- Name
- Legal Name (If Different)
- Code Status/POLST
- Age, DOB, Phone Number
- Weight in Lbs./Kg

MEDICAL COMPLAINT/MECHANISM OF INJURY:

- Chief Complaint/OPQRST
- Background/Time of Injury

ILLNESS/INJURY:

- ECG
- Stroke assessment (PPSS, C-STAT), Last Known Well
- PMHX
- Medications
- Allergies

SIGNS:

- GCS/LOC
- Lowest and Last Blood Pressure
- SpO₂
- CBG
- EtCO₂
- Temperature

TREATMENT:

- IV Site and Size
- Medications and Response to Treatments

To allow paramedics to provide immunizations and infectious disease testing for both seasonal outbreaks and during public health emergencies.

Community immunization and other public health applications are important duties that paramedics and EMTs may perform as determined necessary in cooperation with the Oregon Health Authority (OHA) and the local public health department. Training will be approved by the EMS Medical Director and OHA and may be accomplished under the direction of the OHA and/or local public health department.

PROCEDURE:

- 1. Indications for immunization and/or infectious disease testing:
 - a. The public or EMS agency personnel may be immunized or tested under guidelines developed by OHA or the local public health department.
 - b. Age groups for immunization will be determined by the OHA or public health department as appropriate for the immunization clinic setting or infectious disease testing requirements as determined necessary by the local public health department or agency infection control guidance.
 - c. Timing of immunizations or infectious disease testing will be determined by OHA, EMS agency and public health department to comply with public health needs or agency immunization requirements as determined by agency infection control guidance.
 - d. Immunizations or testing may be performed in clinic, mass immunization or agency setting as approved by OHA and/or local public health department.
- 2. Immunization or infectious disease testing:
 - a. Immunizations or testing may be administered via IM, SQ or intranasal route in dosing determined by guidance provided by the MCA or local public health department as required for the agent administered.
 - b. Other testing methods/procedures will follow guidance and training provided by OHA or the local health department.
 - c. Screening will be performed as determined appropriate for the agent administered by OHA or local health department.
- 3. Training: Training for immunization or infectious disease testing will be provided by local public health department personnel or under an approved OHA program.
- 4. Personnel requirements: Immunizations or infectious disease testing may only be performed by paramedics or EMTs trained by local public health department personnel or under approved OHA training programs.
- 5. Record keeping: A record of public or agency personnel receiving immunizations or infectious disease testing will be maintained by the agency performing the immunizations or testing as determined by the local public health department or OHA.

Hazardous Materials Response – 50.060

PURPOSE:

Non-hazardous materials trained EMS personnel may be first on the scene of a hazardous materials situation because of shorter response times or no knowledge of dispatch that hazardous materials are involved. This protocol is intended to guide personnel who do not normally function in hazardous materials scenes. If the scene you are responding to is a known or <u>suspected</u> (based on information from dispatch) hazardous materials situation, stage and wait for the hazardous materials personnel. When you have arrived at the scene and find out during scene assessment that hazardous materials are involved, stage and wait for the hazardous materials personnel. All scenes (MVA, Industrial, etc.) should be considered as being a potential hazardous materials situation. The following approach procedure should be used:

PROCEDURE:

A. Approach

- 1. All scenes:
 - a. Be cautious all times.
 - b. The reported location may be inaccurate, response into a contaminated area might occur.
 - c. Approach upwind and upgrade if possible.
 - d. Position vehicle well away from the incident.
 - e. Communicate your actions to the 9-1-1 Center.
 - f. Remember: Contaminated and/or exposed response personnel may add to the overall problem and reduce their effectiveness to help.
- 2. If at any time you suspect a hazardous materials situation:
 - a. Confirm that fire and police have been notified. The agency responsible for hazardous materials response may respond with different levels of personnel and equipment based upon the information received. Do not always expect a hazardous materials team to respond.
 - b. If you are a first-in responder, the first priority is scene isolation.
 - c. If you believe that you or your vehicle is contaminated, stage in an isolated area. KEEP OTHERS AWAY! KEEP UNNECESSARY EQUIPMENT FROM BECOMING CONTAMINATED.
- B. Person in Charge
 - If a "non-hazardous materials trained" paramedic is the first medical person on the scene, he/ she should assume the role of PIC (medically) until a "hazardous materials trained paramedic" (HMP) arrives. If possible, the Incident Command Structure should be implemented.
 - 2. The HMP will direct all care.
 - 3. The HMP will determine the method of transport of the exposed patient (air vs. ground).
 - 4. The HMP will determine who will provide care during transport (HMP may remain in that position during transport).

Hazardous Materials Response – 50.060

C. Patient Care for the Contaminated Patient

- 1. Types of incidents which may require decontamination of the patient:
 - a. Radiation
 - b. Biological hazards
 - c. Chemical
 - d. Toxic substances
- 2. Contamination can occur though:
 - a. Smoke
 - b. Vapor
 - c. Direct contact
 - d. Run-off
- 3. Determine the hazardous substance involved and provide treatment as directed by HMP. In the absence of an HMP, consult Poison Control through OLMC.
- 4. The hazardous materials team must be contacted about removal of contaminated clothing and packaging of the patient with regard to your protection and the patient's.
- CI. Ambulance Preparation
 - 1. The HMP shall determine the process needed for ambulance preparation.
 - 2. Remove any supplies and equipment that will be needed for patient care.
 - 3. Seal cabinets and drape interior, including floor and squad bench, with plastic (available from hazardous materials team).
- CII. Transport and Arrival at the Hospital (if requested by HMP)
 - If an ambulance has transported a patient from an incident that is subsequently determined to involve hazardous materials exposure, scene personnel must immediately relay all relevant information to the transporting unit(s) and/or receiving facility(s) involved (via EMS dispatch or OLMC).
 - 2. OLMC and the receiving hospital should be contacted as soon as possible. The EMS providers should communicate the material involved, degree of exposure, decontamination procedures used and patient condition.
 - 3. The ambulance should park in an area away from the emergency room or go directly to a decontamination center or area.
 - 4. Patient(s) should not be brought into the emergency department before the EMS providers receive permission from the hospital staff.
 - 5. Once the patient(s) has been released to the hospital, follow the HMP's direction and if necessary double bag the plastic sheeting used to cover the gurney and the floor. Double bag any equipment, which is believed to have become contaminated.
 - After unloading the patient from the ambulance, check with the HMP to see where the ambulance can be safely decontaminated and whether or not there is equipment available for this purpose. Do not begin decontamination without direction from the HMP. After consultation with the Hazardous Materials Team leader, the HMP may recommend that the ambulance be decontaminated.
 - 7. Following decontamination recommendations from the HMP, decontaminate the ambulance and personnel before returning to the incident scene. When returning to the incident scene, bring bags containing contaminated materials, equipment, clothing, etc., and turn them over to the HMP.

F. EMS Personnel Exposure

- 1. If an EMS provider is exposed or is concerned with the possibility of exposure, medical help should be sought immediately.
- 2. Report all exposures to the HMP, Poison Center, and supervisor, and the on-call OHDP nurse.
- 3. Follow your agencies guidelines for appropriate Personnel Exposure Report.
- 4. Do not return to service until cleared to do so by the HMP or Poison Center.

FOR ADDITIONAL INFORMATION SEE THE HAZMAT PROTOCOL

To provide guidance to the EMS provider in the care of a patient who has a life-limiting or terminal illness and prefers comfort-focused treatment.

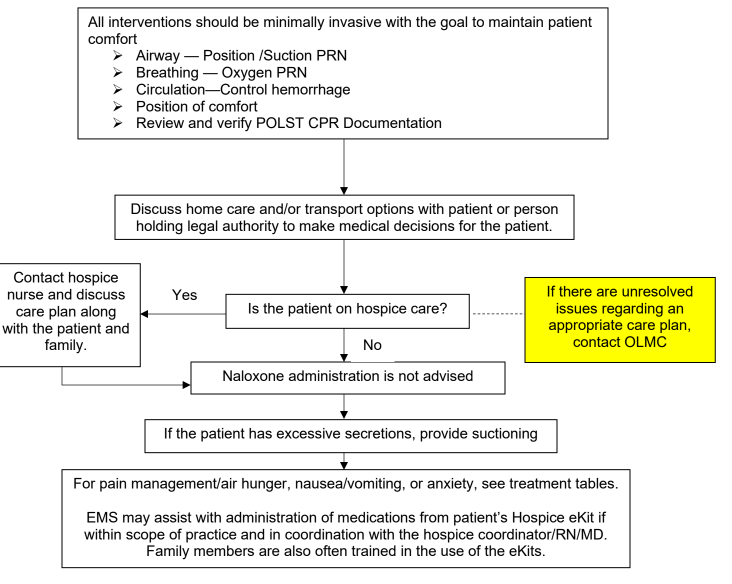
INDICATIONS:

Patient has a life limiting or terminal illness, prefers comfort-focused treatment, and has one of the following:

- A. POLST form specifying DNAR/DNR and comfort focused treatment and/or
- B. Patient is enrolled in hospice care

GOALS:

- A. Reduce patient symptom distress
- B. Maintain patient dignity by aligning care with stated end-of-life preferences
- C. Affirm dying as a normal process



Patients with decision making capacity: If the patient can communicate and has the capacity to make decisions regarding treatment and transport, consult directly with the patient before treatment or transport.

Patients without decision making capacity:

If the patient lacks the capacity to make decisions regarding treatment or transport, identify any advanced care planning in place for information relating to advanced care planning and consent for treatment, including

- Advanced care directives
- POLST
- Guardian, healthcare power of attorney, or other accepted healthcare proxy

NOTES AND PRECAUTIONS:

- A. Palliative care is specialized care for patients with a chronic and/or terminal illness which focuses on managing symptoms exacerbation and the stress of illness.
- B. Hospice care is specialized care, like palliative care, for patients within the last 6 months of life.
- C. Patients may not have a DNAR/DNR or POLST form completed and still be enrolled in hospice care.
- D. Careful and thorough assessments should be performed to identify complaints not related to the illness for which the patient is receiving hospice or palliative care.
- E. Treat dying persons with warmth and understanding. Do not avoid them. Allow them to discuss their situation, but do not push them to talk. Ask the person how you might help.
- F. Many dying people are not upset by discussions of death as long as you do not take away all of their hope. Touching a dying person is important. Use words like "death". Do not use meaningless synonyms. Give factual information.
- G. Be aware of your own fears regarding death and admit when a dying person reminds you of a loved one. If a situation is too disturbing, have your partner or other members of the responding team take over.
- H. Social interactions with the family may affect end-of-life care, including psychological and spiritual aspects of patient care.
- I. Offer support system to help the family cope during the patient's illness and in their own bereavement.
- J. Care should be delivered with the utmost patience, kindness, and compassion.
- K. PICC lines may be accessed for use by Paramedics with sterile techniques.
- L. Emergency Kits (eKit) may be given to patients by hospice to use at home for acute symptom exacerbation. These eKits are individualized and will be different for each patient, but typically contain medication that can address pain, nausea/vomiting, anxiety, and/or secretions. Not every hospice service utilizes an eKit. Family members are frequently trained in the use of the eKit
- M. In collaboration with hospice or palliative care provider, coordinate with guardian, power of attorney, or other accepted healthcare proxy if non-transport is considered.
- N. EMS providers cannot take medical orders from a hospice nurse, but their advice is often invaluable and may be followed with direction from OLMC. Providers can take orders from a hospice physician however, and this can include in the form of a written prescription which may be present in a patient's eKit.
- O. Consider OLMC if hospice or palliative care provider is not available or for on scene conflict.
- P. After medication administration, if no transport occurs, care may be transferred to the hospice nurse or palliative care provider.

TREATMENT TABLES

Acute Pain/Air Hunger (uncomfortable feeling of breathing difficulty)

Severity*	Medications		
	Fentanyl (IV/IN)	Hydromorphone (IV/IM)	Morphine (IV/IM)
Mild	25 mcg	0.5 mg	2 mg
Moderate	50 mcg	1 mg	4 mg
Severe	100 mcg	2 mg	8 mg

*Consider using moderate/severe dose in opiate tolerant patients. Opiate tolerant patients have a typical daily dose of narcotic that is equivalent to \geq 60 mg of oral morphine per day.

Anxiety/Agitation

Severity		Medications		
	Midazolam (IV/IM)	Droperidol (IV/IM)	Haloperidol (IV/IM)	
Mild/Moderate	1 mg	2.5 mg	2 mg	
Severe	2 mg	5 mg	4 mg	

May repeat dose in 15 minutes for IV administration, or 30 minutes for IM injections

Nausea/Vomiting:

Medications			
Ondansetron (PO)	Droperidol (IV/IM)	Haloperidol (IV/IM)	
8 mg	0.625 mg	1.25 mg	

Fire and EMS resources are frequently dispatched to provide lifting assistance. This assistance can vary but often involves an individual who has fallen or slipped and is now unable to get up or return to bed without assistance. In all calls from an individual or responsible party requesting lifting assistance, a medical evaluation must be completed looking for any injury, underlying medical process that contributed to this event, or for a deterioration in functional ability.

PROCEDURE:

- A. Initial evaluation should begin by assessing for any suspected medical cause or inability to mobilize (e.g. dizziness, lightheadedness, syncope, new weakness or balance problem, dehydration/poor oral intake, visual disturbance, recent illness or infection, etc.).
- B. Assess vital signs to include HR, RR, BP, SpO₂. In some instances, based on patient's past medical history or provider discretion, a temperature, EtCO₂, and blood glucose should also be checked.
- C. Determine if any acute injury or medical condition exists.
- D. Ascertain the duration of down time if found on the ground/floor. Consider hypothermia, compartment syndrome, or rhabdomyolysis.
- E. Determine if patient is on any oral anticoagulants which may increase risk level for unrecognized bleeding and may prompt the provider to recommend transport.

NOTES:

- A. Lift assist calls can be a sentinel event for someone that is developing a medical emergency or who has crossed the threshold from being able to live independently to someone who needs a little more help (assisted living, etc.).
- B. Anyone with impaired mobility that requires assistance to mobilize necessitates an assessment of their health status before deciding that the patient does not require further medical assessment.
- C. A PCR will be completed on all patient contacts in which a patient receives any assessment, assistance (i.e. lift assist), advice, or treatment by EMS. The PCR may be brief, but must include vital signs, any assessment/exam provided, and documentation of the lack of a medical complaint.
- D. Those who decline transport should be evaluated for medical decision-making capacity and the informed refusal process should be followed. Advise patient that they may call 911 if they develop any symptoms.
- E. If vitals are unable to be obtained, this must be documented on the PCR along with a reason.
- F. EMS/Fire agencies may (and are encouraged to) develop their own, more expansive and detailed documentation policies specific to their own operations.

To provide guidelines for emergency response personnel on scenes that involve multiple victims who have been exposed to a hazardous material or hazardous environment. This procedure would be used when MSDS and DOT information indicate that victims **may** suffer untoward effects from their exposure and need **short-term**, **continuing medical assessment**. It would also apply when victims are symptomatic and have been exposed to a hazardous environment that poses little risk of long-term effects, such as discharge of tear gas. <u>This protocol is NOT intended for use when there are symptomatic patients and the substance they were exposed to is unknown or when there is a potential for serious or long-term medical consequences.</u>

PROCEDURE:

- A. Triage determines that there are multiple victims who have been exposed to a hazardous material or environment, and that these victims are presently asymptomatic or have been exposed to an agent that has transient effects (e.g., tear gas).
- B. Triage will assist the Hazardous Materials (trained) Paramedic/EMT (HMP) in coordinating removal of the victims from the potentially hazardous environment, then isolate the victims as best as possible in a safe, well lit, and climate-controlled environment (Consider using a bus or a room in a nearby building). If clothing is contaminated, removal of contaminants and proper procedures will be employed prior to isolating victims.
- C. Access to and egress from the Triage and Treatment Area <u>must be strictly controlled at all times</u>. It is necessary to keep track of patients who are under the care of EMS providers, especially when the patient is a minor and his/her parent(s) are present. Patients should not be allowed to leave the treatment or triage area without Triage or Treatment's knowledge. It is recommended that a guard be posted at the entrance and exit to control patient movement.
- D. The HMP will attempt to determine the type and level of exposure. The HMP will then contact MRH with information on the type of chemical and level of exposure. MRH will consult with Poison Control to determine any symptoms that are to be expected, the approximate time line for onset of symptoms, and recommended treatment modalities. When possible, a three-way phone link among the scene, MRH, and Poison Center should be arranged. The HMP will report this information to Triage and to Medical.
- E. All potential patients entering the area will be triage tagged and baseline vitals will be obtained and recorded. It is recommended that the Triage consult with the Medical and assign one EMS provider for every 8 to 10 patients. If any exposure victim starts exhibiting symptoms, they will be immediately removed to the designated Treatment Area.
- F. In consultation with MRH, Triage and HMP will make a determination regarding how long the victims will be observed and the frequency of evaluating and taking vital signs of each patient. A log will be maintained of all patients treated and released. This log will include the patient's name, DOB, the date, symptoms (if any), and disposition.
 - 1. If the patients are asymptomatic after the designated observation time, they may be released. The HMP or Triage will individually brief the patients regarding the symptoms they should watch for and should recommend further medical evaluation by their own physician. Minor patients should only be released to their parent or guardian.
 - 2. Triage or the HMP will inform Medical of the number of patients being released.
- G. It is recommended that Medical proceed with initiating procedures normally undertaken during an MCI. Regional shall be notified that the all-call is precautionary.

The purpose of this protocol is to describe who is in charge of patient care on the scene of medical emergencies and how to resolve disputes with other medical professionals in attendance. This protocol does not apply to MCI/MPS events where ICS is established.

PROCEDURE:

- A. EMS Providers On-Scene: The first arriving, highest certified EMS provider will be the Person-In-Charge (PIC) and will assume responsibility for directing overall patient care. The team approach to patient care assessment and treatment should be utilized by the PIC.
- B. When a higher-level EMS provider arrives, in an EMS role, that individual shall assume the role of PIC, after receiving verbal report from the initial PIC.
- C. The responsibilities of the PIC directing overall patient care include:
 - 1. Assuring that treatment, operations, and communications follow protocols.
 - 2. Coordinating patient care activities. This PIC must watch over the entire patient care scene activities and be sure that the patient care activities are being accomplished in a rapid, efficient, and appropriate manner.
 - 3. Directing other EMS providers to establish airway management, start IVs, etc.
 - 4. Establishing the appropriate time to be spent at the scene for doing patient care.
 - 5. Determining when transportation of the patient is to occur.
 - 6. Performing medical coordination with all agencies and personnel.
- D. The PIC directing overall patient care will be held responsible and accountable for patient care activities performed at the scene and be identified on all patient care reports.
- E. If a patient requires transport and the first arriving PIC is from a non-transporting agency, provision of patient care will be turned over to the transporting Paramedic or flight personnel when:
 - 1. The patient is placed on the transport unit's gurney, OR
 - 2. At a time agreed upon by both EMS providers, continued patient care will then become the responsibility of the transporting unit. There will be a verbal agreement anytime transfer of care from one EMS provider to another takes place.

Paramedic Direction On Scene:

EMS providers take medical direction from:

- Physician Supervisors.
- Regional Protocols.
- On-Line Medical Control (OLMC) as directed in protocols.

Physician On Scene Policy, (within office):

- A. When EMS is called to a physician's office, the EMS providers should receive information from the physician and attempt to provide the service requested by the physician.
- B. While in the physician's office, the physician shall remain in charge of the patient. The EMS providers may follow the direction of the physician if it is within the Scope of Practice and protocols of the PIC. Anytime there is a conflict between a physician's orders and the protocols, OLMC shall be contacted.

C. Once the patient is in the ambulance, unless the physician accompanies the patient, paramedics shall follow the protocols.

Physician On-Scene Policy, (outside office):

- A. Any physician (MD or DO) at the scene of an emergency may be qualified to provide assistance to EMS providers and shall be treated with professional courtesy.
- B. A licensed physician requesting control of patient care at the scene shall be:
 - 1. Thanked for the offer by the PIC.
 - 2. Advised that the EMS providers work under regional protocols and On-Line Medical Control.
 - 3. Advised that we are not permitted to relinquish medical control to a physician on the scene without agreement from On-Line Medical Control.
- C. If the physician requesting control is not the patient's "physician of record," EMS providers shall be authorized to proceed under the direction of the physician ONLY IF ALL THREE OF THE FOLLOWING PROVISIONS ARE MET:
 - 1. OLMC is contacted and authorizes transfer of patient care.
 - 2. The physician agrees to accompany the patient to the hospital in the ambulance.
 - 3. The physician agrees to complete and sign the appropriate patient care report.
- D. If communication with OLMC cannot be established, care may be provided only according to approved ALS protocols. No direction from an on-scene physician may be accepted.

Disputes On-Scene Between EMS providers or Other Medical Professionals:

- A. Disagreements about care should be handled in a professional manner and shall not detract from patient care.
- B. To the extent possible, the ALS and BLS protocols shall be followed and provide the basis for resolving disputes.
- C. If an unresolved dispute continues between EMS providers or other medical professionals concerning the care of a patient, **OLMC shall be contacted**.
- D. If a dispute arises which results in transfer of patient care from one PIC to another, the approximate time of the transfer shall be included on the patient care report.
- E. DISPUTES SHALL NOT APPEAR ON PATIENT CARE REPORTS. Written "Unusual Event Forms", or similar form should be completed pursuant to any dispute arising at the scene.

This protocol describes the steps an EMS provider should follow in contacting a receiving hospital for On-Line Medical Control (OLMC) and describes the contents of the various reports.

PROCEDURE:

- A. Calls to the Receiving Hospital: EMS Providers shall contact the Receiving Hospital by radio or telephone in the following situations:
 - 1. As required by the protocols.
 - 2. As required in approved studies.
 - 3. As required for trauma services.
 - 4. When On-Line Medical Control (OLMC) is needed.
- B. All scenes involving OLMC contact:
 - 1. One person at the scene must be designated as the contact person in charge of communications. The EMS provider designated as "in charge" of communications shall contact the Receiving Hospital by the time transport has begun, including all air ambulance transports.
 - 2. The receiving hospital should be contacted to provide patient status updates during transport for all patients except Trauma System entries.
 - 3. If BLS responders have initiated OLMC communications, ALS responders shall continue to use that medical direction source.
- C. When requesting OLMC, the following information must be relayed
 - 1. Unit number, identity and certification level of person making contact
 - 2. Location of the call, street address if appropriate
 - 3. Purpose of call (Identify the protocol being followed)
 - 4. Age and sex of patient
 - 5. Patient's chief complaint
 - 6. Brief history, prior medical history, medications, and allergies
 - 7. Vital signs
 - 8. Pertinent physical findings
 - 9. Treatment at scene
 - 10. Destination hospital and ETA, including loading time
- D. When contacting the receiving hospital for trauma system patients, the following information must be relayed:
 - 1. Unit number, identity, and certification level of person making contact
 - 2. Location of the incident, street address if appropriate
 - 3. Number of patients. Follow *Multi- Casualty Incident* protocol, if applicable
 - 4. Age and sex of the patients
 - 5. Trauma System entry criteria (be as specific as possible)
 - 6. Trauma Band number(s)
 - 7. Patient's vital signs. Specify if not taken or not present
 - 8. Approximate ETA of patient(s) to Trauma Center; include loading time if appropriate
 - 9. Unit number and mode of transport
 - 10. Patient destination based on incident location or request

- E. Consent and refusal guidelines for minors (reflecting Oregon Revised Statutes):
 - 1. A child under the age of 10 cannot be left alone even if he or she is not a patient. If no responsible adult is present and the child is not a patient, contact law enforcement.
 - 2. Minors who are ages 15 or older and less than 18 years can consent to treatment.
 - 3. If a minor age 15 or older and less than 18 years is refusing treatment/transport contact OLMC.
 - 4. If a minor age 15 or older and less than 18 years is not transported, attempt to contact parents to inform them of the EMS call.

F. High risk medical conditions where OLMC contact should be considered:

EMS providers are encouraged to contact OLMC for the following refusal situations:

- Suspected impaired decision-making capacity.
- Suspected high risk medical condition such as:
 - Age younger than 3 months.
 - Minor (age 17 or younger) without a patient or guardian who is refusing care.
 - Serious chief complaint (including but not limited to, chest pain/dysrhythmia, shortness of breath, BRUE, stroke-like symptoms, syncope, first time seizures, poison/overdose, suspected sepsis, or suspected cervical spine injury).
 - Significant MOI or suspicion of injury.
 - You believe a patient requires evaluation.
 - Conflict on scene regarding refusal of care.
 - Suspected abuse situation involving a minor, elderly, or a person with a disability.
 - Any unconscious or altered mental status (individual or parent/guardian for a minor).
 - Sustained abnormal vital signs:
 - Systolic BP less than 90
 - Respirations greater than 29 or less than 10
 - SpO₂ \leq 90%
 - EtCO₂ less than 25 mmHg or greater than 60 mmHg

DOCUMENTATION:

All instances of an identified patient, with or without impaired decision-making capacity, must be fully documented on a Patient Care Form. A signed refusal form must be obtained on all patients with decision making capacity who are refusing care and/or transport against medical advice. The following is considered minimum documentation criteria:

- General appearance and level of consciousness (mental status).
- History, vital signs, and physical exam.
- Presence of any intoxicants.
- Assessment of the person's decision-making capacity.

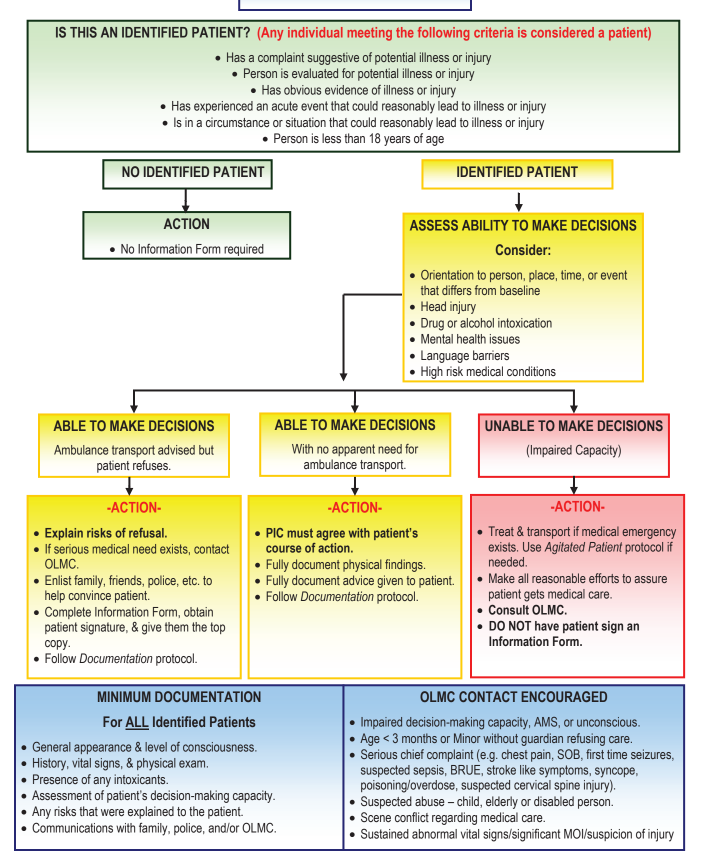
- Risks explained to patient.
- Communication with family, friends, police, and/or OLMC.

GUIDELINES & DEFINITIONS:

- A. Decision Making Capacity: The ability to make an informed decision about the need for medical care based on:
 - Accurate information given the patient regarding potential risks associated with refusing treatment and/or transport.
 - The persons perceived ability to understand and verbalize these risks.
 - The person's ability to make a decision that is consistent with his/her beliefs and life goals.
- B. Impaired Decision-Making Capacity: The inability to understand the nature of the illness or injuries, or the risks and consequences of refusing care.
- C. Emergency Rule: EMS providers may treat and/or transport under the doctrine of implied consent a person who requires immediate care to save a life or prevent further injury. Minors may be treated and transported without parental consent if a good faith effort has been made to contact the parents or guardians regarding care and transport to a hospital, and the patient, in the opinion of EMS provider, needs transport to a hospital. When in doubt, contact OLMC.

Refusal and Informed Consent – 50.117

ASSESS PATIENT'S MEDICAL NEED

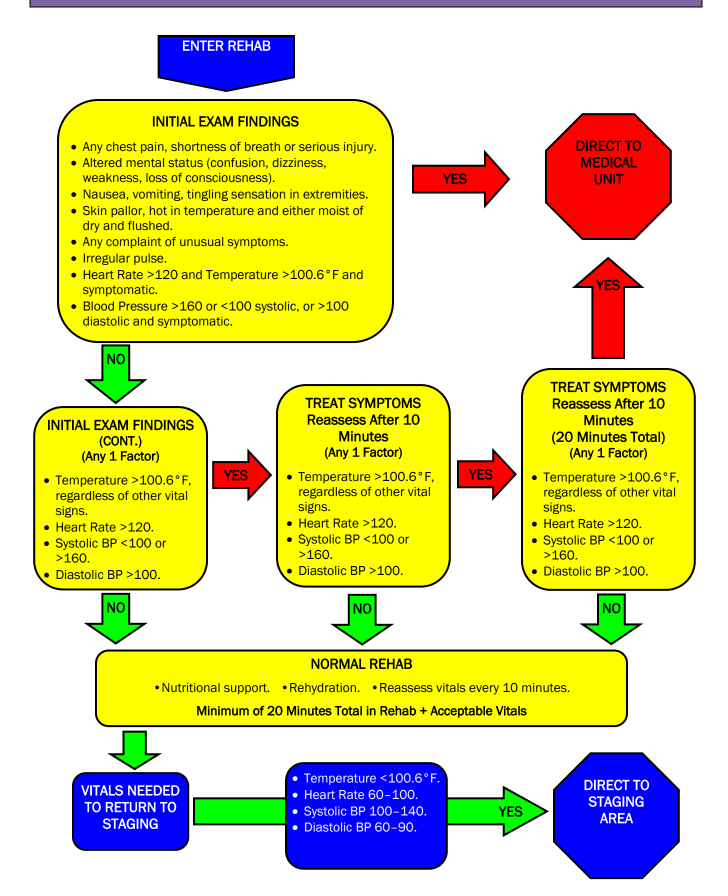


To establish guidelines for the evaluation and treatment of personnel in the Rehabilitation Group (Rehab).

PROCEDURE:

- A. Personnel in Rehab will undergo an initial medical evaluation that will consist of a physical assessment including mental status and vital signs (blood pressure, pulse and temperature, pulse ox and CO monitoring [if available]). All medical evaluations will be recorded on the Medical Evaluation Form.
- B. Medical treatment or a resting period will be determined according to the following triage criteria based on entry findings:
 - 1. Findings mandating that the individual be transferred to the Medical Unit:
 - a. Any chest pain, shortness of breath or serious injury.
 - b. Altered mental status (confusion, dizziness, weakness, loss of consciousness).
 - c. Nausea, vomiting, or tingling sensation in extremities.
 - d. Skin pallor, hot in temperature and either moist of dry and flushed.
 - e. Any complaint of unusual symptoms.
 - f. Irregular pulse.
 - g. Heart Rate >120 and Temperature >100.6°F and symptomatic.
 - h. Blood Pressure >160 or <100 systolic, or >100 diastolic and symptomatic.
 - 2. If initial exam findings include any of the following the individual will require reassessment within 10 minutes:
 - a. Temperature >100.6°F, regardless of other vital signs.
 - b. Heart Rate >120.
 - c. Systolic BP <100 or >160.
 - d. Diastolic BP >100.
 - 3. If reassessment exam findings include any of the following, the individual will require an additional reassessment in 10 minutes:
 - a. Temperature >100.6°F, regardless of other vitals.
 - b. Heart Rate >120.
 - c. Systolic BP <100 or >160.
 - d. Diastolic BP >100.
 - 4. If, after an additional 10 minutes (20 minutes total in Rehab), reassessment exam findings include any of the following, the individual will be sent to the Medical Unit for further evaluation and/or treatment:
 - a. Temperature >100.6°F, regardless of other vitals.
 - b. Heart Rate >120.
 - c. Systolic BP <100 or >160.
 - d. Diastolic BP >100.
 - 5. Exam findings allowing an individual to enter Staging for reassignment include:
 - a. Temperature <100.6°F.
 - b. Heart Rate 60–100.
 - c. Systolic BP 100–140.
 - d. Diastolic BP 60–90.

Rehabilitation – 50.119



Operations - Revised 6/11

Reporting of Suspected Child Abuse – 50.120

PURPOSE: To establish guidelines for the reporting of suspected child abuse.

DEFINITIONS:

- A. Abuse: The non-accidental assault or physical injury to a child. This may include mental abuse, sexual abuse, neglect, etc.
- B. Child: An unmarried person under the age of 18.
- C. Public or Private Officials: physicians, including residents and interns, firefighter or EMT among others.

PROCEDURE:

It is the policy of the State of Oregon to require mandatory reporting of suspected child abuse.

A. DUTY TO REPORT CHILD ABUSE (ORS 419B.010)

All EMS PERSONNEL ARE REQUIRED to report child abuse REGARDLESS OF ON DUTY STATUS. ANY EMT WHO has reasonable cause to believe that a child has either been abused, or witnessed abuse of another child or adult, or who comes into contact with someone who has abused a child, **shall report** the contact to a law enforcement agency, i.e., any city or municipal police department, any county sheriff's office, or the Oregon State Police AS SOON AS POSSIBLE.

Passing the report only to a Nurse or Physician does not meet the requirement of the Statute or the Protocol.

B. CONTENT OF REPORT (ORS 419B.015)

Paramedic must file an Unusual Event Report with the EMS Office within 12 hours as outlined in the Documentation Protocol. If there is imminent danger to health or life, notify police, the Chief Officer and use your agency notification procedure. The report must contain, if known, the following information:

- The names and addresses of the child and parents/person responsible for the child's care.
- The child's age.
- The nature and extent of abuse (including any evidence of previous abuse).
- The explanation given for the abuse.
- Any information the official believes may be helpful in establishing the cause of abuse or the perpetrator's identity.

IMMUNITY OF PERSONS MAKING REPORTS (ORS 419.025):

Persons who acting in good faith and upon reasonable grounds, report child abuse are immune from civil and criminal liability.

To establish guidelines for the reporting of suspected elder abuse or abuse of a resident in a long term care facility.

PROCEDURE:

There are two separate elder abuse reporting requirements; a general reporting requirement which applies to patients outside long-term care facilities and a special reporting requirement, which applies to patients of long-term care facilities.

DEFINITIONS:

- A. Abuse: The non-accidental physical injury to an elderly person or patient of a long term cares facility. Abuse also includes:
 - 1. Outside long-term care facilities:
 - a. **Neglect** means the withholding of services necessary to maintain health and well being. Treatment solely by spiritual means is not neglect; however, the person must be voluntarily under the care of an accredited practitioner or in accordance with the practices of a recognized church or religion.
 - b. **Abandonment**, including desertion or willful forsaking of an elderly person or withdrawal or neglect of duties and obligations owed an elderly person by a caregiver.
 - c. Willful infliction of physical pain or injury.
 - 2. Inside long-term care facilities:
 - a. Illegal or improper use of the patient's financial resources for personal profit or gain.
 - b. Sexual contacts by force, threat, duress or coercion by an employee, agent or other resident.
 - c. Use of derogatory names, phrases, harassment, intimidation, punishment or involuntary seclusion.
- B. Elderly person Any person 65 years of age or over.
- C. Long-term care facility Any licensed skilled nursing facility or intermediate care facility.

PROCEDURE:

A. <u>DUTY TO REPORT ELDER ABUSE AND PATIENT ABUSE IN A LONG TERM CARE</u> <u>FACILITY</u>

An EMT who has reasonable cause to believe that an elderly person has been abused, or who comes into contact with someone who has abused an elderly person, shall file an Unusual Event Report with the EMS Office within 12 hours as outlined in the Documentation Protocol. If there is imminent danger to health or life, notify police, or DHS: 503-304-3400 or

1-800-232-3020. Passing the report only to a Nurse or Physician does not meet the requirement of the Statute or the Protocol.

B. CONTENT OF REPORT

The elder abuse report must contain, if known, the following information:

- The names & addresses of both the elderly person and anyone responsible for his/her care.
- The nature and extent of abuse including any evidence of previous abuse.
- The explanation given for the abuse.
- Any information the official believes may be helpful in establishing the cause of abuse.

Persons participating in good faith in making a report of elder abuse and who have reasonable grounds for making it are immune from civil and criminal liability including participation in any judicial proceeding resulting from their report. Persons making such a report of abuse of a patient in a long term care facility in addition have immunity from any criminal liability that might otherwise be incurred or imposed with respect to making such a report.

Trauma System

National Guideline for the Field Triage of Injured Patients

RED CRITERIA High Risk for Serious Injury

Injury Patterns

- Penetrating injuries to head, neck, torso, and proximal extremities
- Skull deformity, suspected skull fracture
- Suspected spinal injury with new motor or sensory loss
- Chest wall instability, deformity, or suspected flail chest
- Suspected pelvic fracture
- Suspected fracture of two or more proximal long bones
- Crushed, degloved, mangled, or pulseless extremity
- Amputation proximal to wrist or ankle
- Active bleeding requiring a tourniquet or wound packing with continuous pressure

Mental Status & Vital Signs

All Patients

- Unable to follow commands (motor GCS < 6)
- RR < 10 or > 29 breaths/min
- Respiratory distress or need for respiratory support
- Room-air pulse oximetry < 90%

Age 0-9 years

• SBP < 70mm Hg + (2 x age in years)

Age 10-64 years

- SBP < 90 mmHg or
- HR > SBP

• HR > SBP

Age \geq 65 years

• SBP < 110 mmHg or

Tx to Santiam unless in conjunction with other entry criteria

Patients meeting any one of the above RED criteria should be transported to the highest-level trauma center available within the geographic constraints of the regional trauma system

YELLOW CRITERIA Moderate Risk for Serious Injury

Mechanism of Injury	EMS Judgment
 High-Risk Auto Crash Partial or complete ejection Significant intrusion (including roof) >12 inches occupant site OR >18 inches any site OR Need for extrication for entrapped patient (rollover) Death in passenger compartment Child (age 0-9 years) unrestrained or in unsecured child safety seat Vehicle telemetry data consistent with severe injury Rider separated from transport vehicle with significant impact (eg, motorcycle, ATV, horse, etc.) Pedestrian/bicycle rider thrown, run over, or with significant impact Fall from height > 10 feet (all ages) 	 Consider risk factors, including: Low-level falls in young children (age ≤ 5 years) or older adults (age ≥ 65 years) with significant head impact Anticoagulant use Suspicion of child abuse Special, high-resource healthcare needs Pregnancy > 20 weeks Burns in conjunction with trauma Children should be triaged preferentially to pediatric capable centers If concerned, take to a trauma center

Patients meeting any one of the YELLOW CRITERIA WHO DO NOT MEET RED CRITERIA should be preferentially transported to a trauma center, as available within the geographic constraints of the regional trauma system (need not be the highest-level trauma center)

II. MEDICAL DIRECTION:

- A. Off-line medical direction for trauma patients is controlled by the Treatment Protocols.
- B. OLMC is provided by **Receiving Hospital**. OLMC may override off-line medical direction.

III. COMMUNICATIONS:

- A. Communications with OLMC:(Receiving Hospital)
 - The following information will be provided:
 - 1. Unit number.
 - 2. Age and sex of the patients.
 - 3. Trauma system entry criteria and vital signs.
 - 4. ETA to Trauma Center.

IV. TRAUMA CENTER DESTINATION:

- A. **Mode of Transportation:** An air ambulance should be used when it would reduce total prehospital time by 10 minutes or greater. This is usually achieved whenever the ground transport time will exceed 25 minutes (Scene is > 15 miles from Salem or other circumstances exist).
- B. **Patients or Guardians Request:** If the alert, competent and stable patient, or his/her competent guardian demands transport to a specific hospital, the EMT must honor that request.
- C. Multiple Patients: From the same scene, all patient destinations are to be that assigned by the above service areas unless the designated Trauma Center advises the that the facility cannot accept additional patients. In this instance, the OLMC will assist the paramedic in determining patient destinations. If there are more than two critical trauma patients (e.g., intubated, significant trauma) ready to be transported from the same scene, only the first two will be sent to the Level 1 facility designated by catchment area. Subsequent patients shall be directed to the next Level 1 center.
- D. **Diversion to Local Hospital:** If the paramedic is unable to establish an airway, the patient should be transported to the nearest acute care facility.

V. PATIENT EVALUATION PROTOCOL:

- A. Treatment priority should be approached in this order:
 - 1. Control of hemorrhage
 - 2. Airway (with control of the cervical spine). If unable to establish and maintain an adequate airway, the patient should be transported to the nearest acute care facility to obtain definitive airway control.
 - 3. Breathing
 - 4. Circulation
 - 5. Disability assessment (GCS, pupil size and reactivity, motor function)
 - 6. Exposure and temperature control
 - 7. Detailed head to toe exam
 - 8. Splinting of suspected fractures

Multi-Casualty Incidents

The National Incident Management System (NIMS) will be used to manage all incidents.

- 1. Incident Command (IC) is the responsibility of the agency having jurisdiction (AHJ).
- 2. Each assisting agency shall retain full authority to operate within the scope of its agency operational and administrative protocols and procedures.
- 3. Agencies that are assisting in the support of a single jurisdiction will function under the direction of that jurisdiction's designated Unified Incident Command.
- 4. Incident Command of a multi-discipline event should be predicated on the "Primary Hazard" of the event.
- 5. In a Unified Command, the "Lead Agency" may change as priorities change.

The **Mass Casualty Incident Protocol** is a tool that may be used in part or whole as determined by the on-scene Incident Commander in situations where the number of patients exceeds the resources of the on-scene responders. <u>There is no set number of patients that will automatically initiate this protocol</u>. If the Incident commander determines that additional resources or incident structure is needed to better manage due to the complexity of the incident, he/she shall announce to dispatch that an MCI is being declared. This may be done upon arrival or at any time during the incident.

- If the incident involves multiple asymptomatic patients (HazMat exposure) set up secure evaluation area. See *Multiple Toxic Exposure* protocol.
- During a declared MCI, the Trauma System is not in effect.
- "Licensed ambulances" are not needed for transport.
- If transport resources are limited, more than one critical patient may be placed in an ambulance.
- All MCI Forms have been moved to Patient Aids and in MCI Kit on Medic Units

Hazardous Materials

HAZMAT RESPONSE GUIDE SHEET

1. Awareness: WMD/Terrorism Indicators

Assess dispatch and size-up information for WMD/terrorism indicators:

- Responding to a target hazard / event or an existing threat of attack
- Explosion (single or multiple)
- Unexplained liquids, unusual odors, reports of a spray release
- Multiple victims (if non-traumatic, signs/symptoms may be similar)
- Injured responders, unexplained dead wildlife

One indicator: remain alert. If multiple indicators: possible WMD/terrorism incident. If this is so, use extreme caution, and communicate this information to Dispatch.

2. Arrival:

Cautious approach:

- Upwind / uphill / maintain safe distance / avoid building exhaust vents
- Park heading away from incident; consider escape routes
- Stay aware of surroundings (alert for secondary devices, other threats, etc)

3. Risk/Benefit Assessment:

Assess scene:

- Type of incident if known (explosive, radiological, chemical, biological, etc)
- Type of product if safe to ID; use Guide 111 if product is chemical and unknown
- Number of victims (ambulatory and non-ambulatory)
- Safety & security / secondary device awareness / hot zone size / threat assessment

Assess self-protection measures:

DECONTAMINATION ZONE

Note: All victims suspected of ingestion or significant exposure to **hydrogen cyanide** solution **require decontamination**. Others may be transferred immediately to the Support Zone.

- A. Decontamination
 - 1. Victims who are able and cooperative may assist with their own decontamination.
 - a. **Rapidly remove contaminated clothing** while flushing exposed skin and hair with plain water for 2 to 3 minutes.
 - b. Then wash twice with mild soap.
 - c. Rinse thoroughly with water.
 - d. Double bag contaminated clothing and personal belongings.
 - 2. Irrigate exposed or irritated eyes with plain water or saline for 5 minutes.
 - a. Continue eye irrigation during other basic care or transport.
 - b. Remove contact lenses if present and easily removable without additional trauma to the eye.
- B. Transfer to Support Zone as soon as decontamination is complete.

IDENTIFICATION CAS 74-90-8 UN 1051

Synonyms include formic anammonide and formonitrile. Aqueous solutions are referred to as hydrocyanic acid and prussic acid.

Hydrogen cyanide is very volatile, producing potentially lethal concentrations at room temperature. At temperature below 78°F, hydrogen cyanide is colorless or pale blue liquid (hydrocyanic acid); at higher temperatures, it is a colorless gas. It has a faint bitter almond odor and a bitter burning taste. It is soluble in water. Hydrogen cyanide is lighter than air.

PRECAUTIONS

- A. Persons whose clothing or skin is contaminated with cyanide containing solutions can secondarily contaminate personnel by direct contact or through off-gassing vapor.
 - 1. Avoid dermal contact with cyanide-contaminated victims and their bodily fluids.
 - 2. Take special care with victims who may have ingested cyanide, as cyanide salts dissolve in the stomach and react with hydrochloric acid to produce hydrogen cyanide gas. Transport patients in vehicles with windows opened and/or good ventilation. These patients who meet *Death in the Field* criteria should be considered a Hot Zone.
 - 3. Victims exposed only to hydrogen cyanide gas do not pose contamination risks to rescuers.
- B. Hydrogen cyanide is a volatile flammable liquid at room temperature; as a gas, it is flammable and potentially explosive.
- C. Hydrogen cyanide is absorbed well by inhalation and can produce death within minutes.
 - 1. Substantial absorption can occur through intact skin if vapor concentration is high.
 - 2. Exposure by any route may cause systemic effects.

HEALTH EFFECTS

HCN is classified a systemic (chemical) asphyxiant. Cyanides interfere with the intracellular utilization of oxygen resulting in cellular dysfunction and cell death. Effects are most profound and first evidenced in the CNS and cardiovascular system. Initial symptoms may include CNS excitation and cardiovascular compensation followed by depression/collapse of both systems.

ROUTES OF EXPOSURE

A. Inhalation

- 1. Hydrogen cyanide is readily absorbed from the lungs; symptoms of poisoning begin within seconds to minutes.
- 2. The odor of cyanide does not provide adequate warning of hazardous concentrations. Perception of the odor is a genetic trait (20% to 40% of the general population cannot detect hydrogen cyanide); also, rapid olfactory fatigue can occur.
- B. Skin/Eye Contact: Exposure to hydrogen cyanide can cause skin and eye irritation and can contribute to systemic poisoning with delayed symptoms.
- C. Ingestion of hydrogen cyanide solutions or cyanide salts can be rapidly fatal

SIGNS AND SYMPTOMS

- A. Signs and symptoms usually develop rapidly. Initial symptoms are nonspecific and include excitement, dizziness, n/v, HA and weakness.
- B. Progressive signs and symptoms may include: Drowsiness, tetanic spasm, convulsions, hallucinations and loss of consciousness.
- C. Cardiovascular Can cause various life-threatening dysrhythmias.
- D. Respiratory
 - 1. Victims may complain of shortness of breath and chest tightness.
 - 2. Pulmonary findings may include rapid breathing and increased depth of respiration.
 - 3. As poisoning progresses, respirations become slow and gasping; cyanosis may be present, and pulmonary edema may develop.

RESCUER PROTECTION

- A. Respiratory protection: Pressure demand self-contained breathing apparatus (SCBA) is recommended in response situations that involve exposure to potentially unsafe levels of hydrogen cyanide.
- B. Skin protection: Chemical protective clothing is recommended because both hydrogen cyanide vapor and liquid can be absorbed through the skin to produce systemic toxicity.

DECONTAMINATION ZONE

- A. Refer to Decontamination page.
- B. Transfer to Support Zone as soon as decontamination is complete.

SUPPORT ZONE

- A. Be certain that victims have been decontaminated properly. Additional decontamination may be required for exposed skin and eyes.
- B. Decontaminated victims or those exposed only to vapor, pose no serious risks of secondary contamination to rescuers. In these cases, Support Zone personnel require no specialized protective gear.

TREATMENT

Patients who rapidly regain consciousness and who have no other signs or symptoms may not require antidote treatment. Patients who remain comatose or develop shock should be treated promptly with the antidotes per OLMC direction. In cases of ingestion—emesis and activated charcoal are *contraindicated*.

- A. High flow oxygen, establish IV access, apply cardiac monitor and secure protected airway following Airway Management protocol.
- B. If Cyanide Toxicity is suspected based on findings (soot in mouth, nose or oropharynx, know exposure) and patient is comatose, in cardiac or respiratory arrest, or has persistent hypotension despite fluid resuscitation:
 - 1. Administer Hydroxocobalamin (CYANOKIT[®]) 5 g IV or IO over 15 minutes. Repeat once if needed. For cardiac arrest, hydroxocobalamin should be administered as a rapid fluid bolus.
 - 2. If Hydroxocobalamin (CYANOKIT[®]) is not available, then administer Sodium Thiosulfate 50 ml of 25% solution over 10-20 minutes. Pediatric dose is 1.65 ml/kg.

- 3. Do NOT administer Hydroxocobalamin (CYANOKIT[®]) and Sodium Thiosulfate to the same patient.
- 4. Treat other presenting symptoms per appropriate protocol.
- 5. Initiate emergent transport to appropriate facility.
- 6. Patients in shock or having seizures should be treated according to existing protocols. These patients may be seriously acidotic; consider giving sodium bicarbonate 50 mEq, with OLMC direction.
- C. **MULTI-CASUALTY TRIAGE -** Patients who have only brief inhalation exposure and mild or transient symptoms may be discharged.

IDENTIFICATION CAS 7664-39-3 UN 1052 (Anhydrous) UN 1790 (Solution)

Synonyms include fluoric acid, hydro fluoride, hydrofluoric acid, and fluorine monohydride.

Hydrogen fluoride is a colorless, corrosive fuming liquid or gas (boiling temperature 67°F) with a strong irritating odor. It is usually shipped in cylinders as a compressed gas. Hydrogen fluoride readily dissolves in water to form colorless hydrofluoric acid solutions. Dilute solutions are indistinguishable from water. It is present in a variety of over-the-counter products at concentrations of 6% to 12%.

PRECAUTIONS

- A. Victims whose clothing or skin is contaminated with HF liquid, solution or condensed vapor, can secondarily contaminate response personnel by direct contact or through off-gassing vapor.
- B. Inhalation hazards result not only from HF gas but also from fumes arising from concentrated hydrogen fluoride liquid <u>or from the patient's bodily fluids.</u>
- C. Rapid flushing of exposed areas with water is critical. HF is water-soluble.

HEALTH EFFECTS

The toxic effects of hydrogen fluoride are due primarily to the fluoride ion. The fluoride ion combines with endogenous calcium and magnesium to form insoluble calcium fluoride and magnesium fluoride.

- A. This results in cell destruction and local bone demineralization.
- B. Life threatening hypocalcemia, hypomagnesemia, and hyperkalemia can occur.
- C. The adverse action of the fluoride ion may progress for several days.

ACUTE EXPOSURE

- A. <u>Respiratory</u>—Due to HF's water solubility, effects of exposure generally occur in the upper airway including the glottis. However, people incapacitated in large clouds of HF can have severe deep lung injury.
 - 1. **Mild effects** mucous membrane irritation, cough and narrowing of the bronchi.
 - 2. Severe effects:
 - a. Almost immediate narrowing and swelling of the throat, causing upper airway obstruction.
 - b. Lung injury may evolve rapidly or may be delayed in onset for 12 to 36 hours.
 - c. Pulmonary edema and constriction of the bronchi. Partial or complete lung collapse can occur.
 - d. Pulmonary effects can result even from splashes on the skin.
- B. <u>Dermal</u>—Depending on the concentration and duration of exposure, skin contact may produce pain, redness of the skin, and deep, slow healing burns with symptoms delayed up to 24 hours. HF can penetrate tissues deeply, causing both local cellular destruction and systemic toxicity.

C. Ocular

- 1. **Mild effects** Rapid onset of eye irritation.
- 2. **More severe effects** May result from even minor hydrofluoric acid splash include, sloughing of the surface of the eye, swelling of the structures of the eye, and cell death due to lack of blood supply. Potentially permanent clouding of the eye surface may develop immediately or after several days.

D. Gastrointestinal

- 1. A small amount of ingested HF is likely to produce systemic effects including acid-base imbalance and may be fatal.
- 2. Ingestion of hydrofluoric acid may cause corrosive injury to the mouth, throat and esophagus as well as inflammation and bleeding of the stomach.
- 3. Nausea, vomiting, diarrhea, and abdominal pain may occur.
- E. **Electrolyte disturbances**—Exposure by any route may result in systemic effects: Hypocalcemia and/or hypomagnesemia and/or hyperkalemia.

PREHOSPITAL MANAGEMENT

HOT ZONE

Rescuer Protection

- A. SCBA is recommended in response situations that involve exposure to potentially unsafe levels of hydrogen fluoride.
- B. Skin protection: Chemical protective clothing, i.e. level A or level B, is recommended because skin exposure to either vapor or liquid may cause severe consequences.

DECONTAMINATION ZONE

- A. Victims exposed only to hydrogen fluoride gas or vapor who have no skin or eye irritation do not need decontamination, they may be transferred immediately to the Treatment Area.
- B. Rescuer Protection: If exposure levels are determined to be safe, personnel wearing a lower level of protection than that worn in the Hot Zone may conduct decontamination.
- C. ABC Reminders:
 - 1. Quickly ensure a patent airway—anticipate airway edema.
 - 2. Stabilize the cervical spine with a c-collar and a backboard if trauma is suspected.
 - 3. Administer supplemental O₂.
 - 4. Assist ventilation with a bag-valve-mask device if necessary.
- D. Basic decontamination:
 - 1. Victims who are able and cooperative may assist with their own decontamination
 - a. **RAPIDLY REMOVE CONTAMINATED CLOTHING** while flushing exposed skin and hair with plain water for 15 minutes.
 - b. If treatment recommended below is available, water flushing may be reduced to 5 minutes and the treatment should be started immediately.
 - Calcium gluconate 3 g mixed with 5 oz water soluble lubricant and applied to burn.
 - c. Double bag contaminated clothing and personal belongings.
 - 2. Irrigate exposed or irritated eyes with plain water or saline or 5 minutes.
 - a. Continue eye irrigation during other basic care or transport.

b. Remove contact lenses if present and easily removable without additional trauma to the eye.

- 3. In case of ingestion, do not induce emesis or administer activated charcoal.
 - a. Victims who are conscious and able to swallow should be given 4 to 8 ounces of water or milk.
 - b. If available, also give 2 to 4 ounces of an antacid containing magnesium (e.g., Maalox, Milk of Magnesia) or calcium (e.g., TUMS).
- 4. As soon as basic decontamination is complete, move the victim to the Treatment Area.

TREATMENT

Be certain that victims have been decontaminated properly. Treatment Area personnel require no specialized protective gear if victims have undergone decontamination.

- A. ABCs, C-spine (prn), Pulse Oximetry, and ECG to obtain baseline QT interval (may be of benefit for this).
- B. Treat patients who are symptomatic per existing protocols.
- C. Observe for signs of hypocalcemia and contact OLMC regarding treatment with Calcium Gluconate.
 - 1. ECG—prolonged Q-T interval or QRS or ventricular dysrhythmias.
 - 2. Other-Muscular tetany. This is probable after ingestion of even small amounts of HF.
- D. For inhalation victims.
 - 1. Administer 2.5% calcium gluconate by nebulizer. Mix 1cc of 10% Calcium Gluconate with 3ccs of Normal Saline into the nebulizer.
 - 2. If wheezes are present, consider use of Albuterol per Respiratory Distress protocol.

E. Minor Burns.

- 1. Initially, the health care provider should wear rubber or latex gloves to prevent secondary contamination.
- 2. Calcium gluconate 3 g mixed with 5 oz water soluble lubricant and applied to burn.
- 3. Continue this procedure until pain is relieved or more definitive care is rendered.

F. Hand Exposure

- 1. Subungual (under the nail) burns often do not respond to immersion
 - treatment. The treatment for hand burns requires expert assistance; consult with OLMC.
- 2. Treatment of hand exposures can be accomplished by placing calcium gluconate gel into an exam glove and placing the glove on the affected hand.
- G. **Optical Exposure**—Irrigate exposed eyes with a 1% aqueous solution of calcium gluconate (10 ml of 10% solution in 90 ml of sterile saline in Buretrol) using a nasal cannula.
 - 1. Up to 500 ml over 1 to 2 hours may be used.
 - 2. If calcium gluconate is not available, use normal saline for irrigation.

MULTI-CASUALTY TRIAGE

Consult with the OLMC for advice regarding triage of multiple victims. Persons who have had only minor or brief exposure to hydrogen fluoride gas or vapor and are initially asymptomatic are not likely to develop complications. See Multiple Toxic Exposure Protocol.

IDENTIFICATION CAS 56-38-2 UN 2783

Synonyms include Alkron, Alleron, Danthion, DNTP, DPP, Ethyl Parathion, Etilon, E-605, Stathion, Sulphos, and Thiophos.

The term organophosphate (OP) is generally understood to mean an organic derivative of phosphoric or similar acids. There are many different OPs and they differ to some extent in their properties. Many OPs inhibit an enzyme known as acetylcholinesterase. This is a class effect of OPs, but not all OPs (e.g. glyphosate) demonstrate this effect. Inhibitors of acetylcholinesterase affect certain nerve junctions in animals, as well as parasympathetic effector sites (the heart, lungs, stomach, intestines, urinary bladder, prostate, eyes and salivary glands). By inhibiting the enzyme acetylcholinesterase, OPs prevent the nerve junction from functioning properly.

PRECAUTIONS

- A. Organophosphates are highly contaminating.
- B. Victims whose skin or clothing is contaminated with liquid or powdered organophosphate can secondarily contaminate response personnel by direct contact or off gassing of solvent vapor.
- C. Clothing and leather goods (e.g., belts or shoes) cannot be reliably decontaminated; they should be incinerated.
- A. Special care should be taken to avoid contact with the vomitus of a patient who has ingested organophosphate.

PHYSICAL PROPERTIES

- A. At room temperature, organophosphate are powders or combustible liquids.
- B. Organophosphates are almost insoluble in water, slightly soluble in petroleum oils, and miscible with many organic solvents. Accordingly, most commercial products contain hydrocarbon solvents.
- C. Organophosphates have low vapor pressures; thus significant inhalation is unlikely at normal temperatures (Exception: Dichlorvos (a.k.a. DDVP and Vapona) when in a poorly ventilated confined space). However, the hydrocarbon solvents remain volatile and flammable, as well as possessing toxic properties.

ROUTES OF EXPOSURE

A. Inhalation:

- 1. Toxic inhalation of organophosphate vapor is unlikely at ordinary temperatures because of its low volatility, but toxic effects can occur after inhalation of organophosphate sprays or dusts.
- 2. The hydrocarbon solvents (most commonly toluene and xylene) used to dissolve organophosphate are more volatile than organophosphate itself, and toxicity can result from inhalation of solvent vapor as well.
- B. Skin/Eye Contact—Organophosphates are rapidly absorbed through intact skin or eyes, contributing to systemic toxicity.
- C. Ingestion—Acute toxic effects. May be rapidly fatal.

HEALTH EFFECTS

- A. Introduction:
 - Organophosphates are known as cholinesterase inhibitors. Normally, the neurotransmitter acetylcholine (ACh) is broken down by acetylcholinesterase (AChE). Organophosphates inhibit the activity of AChE and thus ACh is not broken down. The resulting accumulation of ACh overstimulates ACh receptors (aka cholinergic receptors) within the central and peripheral nervous systems. The toxic effects of organophosphates result from this overstimulation of ACh receptors. There are two types of ACh receptors, muscarinic and nicotinic.
 - 2. Signs and symptoms of poisoning vary according to age, dose, and concentration:
 - a. **CNS effects**—Irritability, nervousness, giddiness, fatigue, lethargy, impairment of memory, confusion, slurred speech, visual disturbance, depression, impaired gait, convulsions, loss of consciousness, coma, and respiratory depression. CNS effects can be some of the earliest symptoms.
 - b. **PNS Effects**—Nicotinic and muscarinic stimulation can provide opposing effects. In general, nicotinic signs and symptoms predominate early in organophosphate poisoning, while muscarinic signs and symptoms predominate later.
 - Muscarinic effects— SLUDGE (Salivation, Lacrimation, Urination, Defecation, Gastroenteritis, Emesis), or DUMBELS (Diarrhea, Urination, Miosis, Bradycardia, Bronchorrhea, Bronchospasm, Emesis, Lacrimation, Salivation, Secretion, Sweating).
 - **ii. Nicotinic effects**—**MTWHF** (Mydriasis, Tachycardia, Weakness, Hypertension, Hyperglycemia, Fasciculations, Flaccidity).

PREHOSPITAL MANAGEMENT

HOT ZONE

- A. Respiratory Protection: SCBA is recommended in response situations that involve exposure to potentially unsafe levels of organophosphates.
- B. Skin Protection: Chemical-protective clothing is recommended because organophosphates are rapidly absorbed through the skin and may cause systemic poisoning.

DECONTAMINATION ZONE

All victims suspected of organophosphate ingestion, or substantial exposure to aerosolized organophosphates, or who have skin or eye exposure to liquid or powdered organophosphates require thorough decontamination.

BASIC DECONTAMINATION

Follow Decontamination General Guidelines. Then, move the victim to the Treatment Area upon completion.

SIGNS AND SYMPTOMS

- A. Mild poisoning HA, n/v, abdominal cramps, and diarrhea.
- B. Moderate poisoning: Generalized muscle weakness and twitching, slurred speech, pinpoint pupils, excessive secretions, and shortness of breath.

C. Severe poisoning: Seizures, skeletal-muscle paralysis, respiratory failure, and coma.

TREATMENT

- A. Secure protected airway in cases of respiratory compromise per Airway Management protocol.
- B. There is no contra-indication to the use of paralytic agents is in this setting, however both *succinylcholine and vecuronium will have a significantly sustained duration of paralysis in the presence of organophosphates*.
- C. The initial intravenous dose of atropine in adults should be determined by the severity of symptoms. In seriously poisoned patients, very large doses may be required. Alterations of pulse rate and pupillary size are unreliable indicators of treatment adequacy. **Atropine works** *only* to correct muscarinic effects.
 - 1. In mild to moderate poisonings (e.g. headache, mild bronchorrhea, nausea, vomiting, diarrhea but normal mentation), administer atropine 1-2 mg IV/IO/IM every 3-5 minutes until symptoms improve.
 - For severe poisoning (e.g. altered mental status, unconsciousness, seizures), administer atropine 3-5 mg IV/IO/IM every 3-5 minutes until symptoms begin to improve.
 - 3. Treat seizures per seizure protocol.
- D. Administer pralidoxime (2-PAM), if profound weakness or paralysis present.
 - 1. Moderate symptoms—1,200 mg (two Mark 1 injectors or one Duodote).
 - 2. Severe symptoms—1,800 mg (three Mark 1 injectors or three Duodote injectors).

CAUTION: When administering 2-PAM intravenously, administer at rate of less than 200 mg/minute, (4 mg/minute for children).

Note: The Mark 1 auto-Injector atropine is 2 mg. The 2-Pam auto-injector is 600 mg pralidoxime. The Duodote Auto-Injector is atropine 2.1 mg/0.7 mL and pralidoxime chloride 600 mg/2 mL.

E. Patients who are comatose, hypotensive, have seizures or cardiac dysrhythmias should be treated according to ALS protocols.

TRANSPORT TO MEDICAL FACILITY

- A. Report to OLMC, and the receiving medical facility, the condition *of* the patient, treatment given, and estimated time of arrival at the medical facility.
- B. If organophosphate has been ingested:
 - 1. Prepare the ambulance in case the victim vomits toxic material.
 - 2. Prepare several towels (or other absorbent material) and open plastic bags to quickly clean up and isolate vomitus.

MULTI-CASUALTY TRIAGE

Patients who have histories or evidence suggesting substantial exposure and all persons who have ingested organophosphate should be transported to a medical facility for evaluation.

Organophosphates – 70.040

- A. Others may be discharged from the scene after their names, addresses, and telephone numbers are recorded.
- B. They should be advised to seek medical care promptly if symptoms develop or recur.

PEDIATRIC PATIENTS:

Atropine: In children, dose is 0.05 mg/kg IV/IO. Pralidoxime: Pediatric dose: 25 to 50 mg/kg and must be given slowly via IV (4 mg/min.)