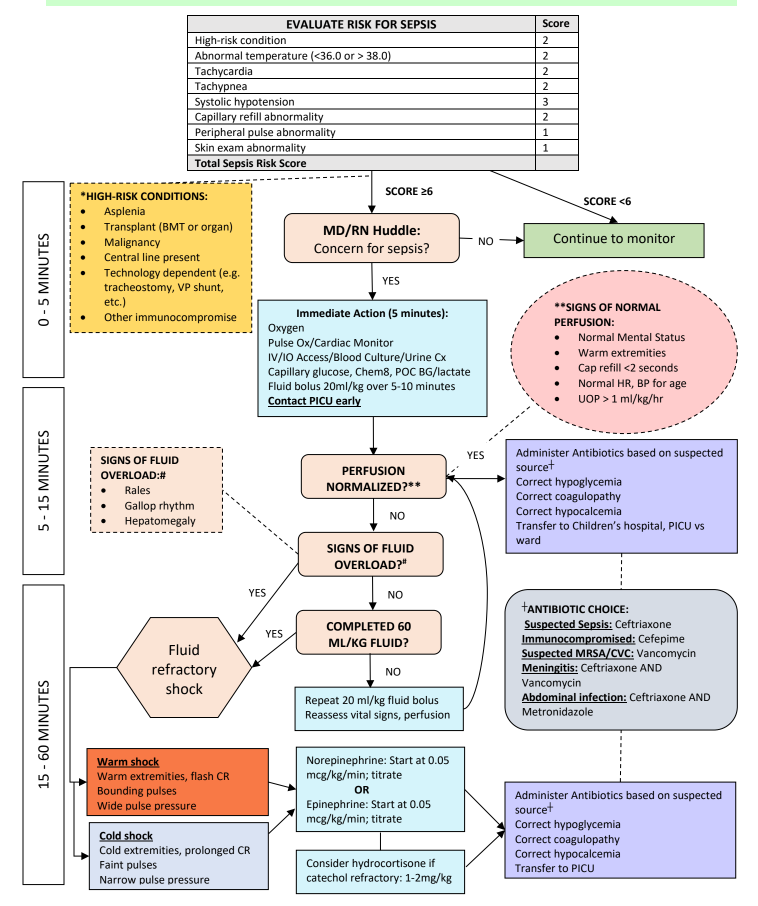
Pediatric Sepsis / Septic Shock in Patients > 60 days Clinical Pathway Mar 2022						
Outcomes/Goals	 Rapid identification and treatment of pediatric patients ≥ 60 days presenting in septic shock. Create a team-oriented approach to efficient and timely evaluation and work-up. Early and aggressive treatment to stabilize hemodynamic status and reverse shock. 					
NURSE	Chief complaint. Onset of symptoms. Presence of high-risk medical condition. Presence of					
Documentation	an indwelling catheter, CVC. Assessment including hemodynamic status (core temp, changes, cap refill, urine output, pulse quality, neuro status)					
Evaluation	Calculate risk score for sepsis (see algorithm)					
INTERVENTIONS	ESI Triage level II					
Initiate on arrival	Full set of vitals including core temperature					
	Apply cardiac monitor, continuous pulse oximetry					
	Establish IV (2 if possible, largest size appropriate)					
	Consider IO if cannot obtain IV in 3 attempts or 90 seconds					
	Bedside CBG					
	Oxygen					
	Initiate warming devices if applicable					
DIAGNOSTICS	POC blood gas/lactate, cap blood glucose					
	Consider POC Chem 8					
	Blood culture (if CVC present, from each lumen) — prior to antibiotics					
	CBC with differential					
	CMP, magnesium, phosphate, Ca					
	Coagulopathy panel, Type and Screen					
	Catheter specimen UA, microscopy, mandatory culture					
	Consider Chest x-ray (portable) +/- 2 view abdomen					
	Consider LP if hemodynamically stable (gram stain, culture, cell count, protein, glucose, hold					
	extra fluid)					
	Consider influenza, RSV, COVID-19					
PHYSICIAN (LIP)						
Fluids	Normal Saline bolus 20 ml/kg in the first 15 minutes; use 3-way stopcock if needed					
	-Reassess for normalization of perfusion (HR, BP, cap refill, pulses, mental status)					
	Repeat up to 60ml/kg/first hour until normalization of perfusion or signs of fluid overload					
Medication Antibiotics	***GOAL TO ADMINISTER WITHIN 1 HOUR OF ARRIVAL FOR SEVERE SEPSIS, 3 HOURS FOR SEPSIS***					
	Suspected Sepsis: Ceftriaxone 50mg/kg (max 2g) IV					
	Immunocompromised: Cefepime 50mg/kg (max 2g) IV					
	Suspected MRSA or Central Venous Catheter: Vancomycin 20mg/kg (max 2g) IV					
	Meningitis: Ceftriaxone 100mg/kg (max 2g) AND Vancomycin 20 mg/kg (max 2g)					
	Abdominal infection: Ceftriaxone 50mg/kg (max 2g) IV AND Metronidazole 10mg/kg (max					
	1.5g) IV					
Vasoactives	Either:					
	 Norepinephrine 0.05-1 mcg/kg/min 					
	Epinephrine 0.05-1 mcg/kg/min					
Calcium	Calcium gluconate 10% 50mg/kg IV over 5 minutes for iCa < 1.1					
Dextrose	D10 5ml/kg for CBG <60					
Antipyretics	Acetaminophen 12.5 mg/kg PO					
	Acetaminophen 15 mg/kg PR					
Corticosteroids	Hydrocortisone 1-2mg/kg (max 100mg) IV					
ADMISSION	Arrange transfer to Children's hospital, ward vs PICU. Fluid refractory shock should always b					
	admitted to PICU. Contact Peds ED or PICU early.					

Clinical Pathway Decision Making Process

Pediatric Sepsis / Septic Shock in Pts \geq 60 days

Mar 2022



Pediatric Sepsis / Septic Shock Rationale and Data

Goals of Clinical Pathway

- 1. Rapid identification and treatment of pediatric patients presenting in sepsis/septic shock.
- 2. Create a team-oriented approach to efficient and timely evaluation and work-up.
- 3. Early and aggressive treatment resulting in stabilization of hemodynamic status and reversal of shock.

Rapid Recognition

The 2005 International Pediatric Consensus Conference definition of sepsis remains the most widely used for children. According to this definition:

<u>Sepsis:</u> ≥2 Systemic Inflammatory Response Syndrome (SIRS) criteria (abnormal temperature, heart rate, respiratory rate, and white blood cell count) AND suspected or confirmed infection

<u>Severe Sepsis:</u> Sepsis criteria AND either cardiovascular dysfunction (hypotension or any two of the following: metabolic acidosis, elevated lactate, oliguria, or prolonged capillary refill), acute respiratory distress syndrome, or \geq 2 other criteria for end organ dysfunction due to sepsis

<u>Septic Shock</u>: Severe sepsis and hypotension or tissue hypoperfusion (cap refill <1 second (flash) or \geq 3 seconds, lactate \geq 4) Though these definitions are useful, sepsis presentations represent a continuum and it can be clinically difficult to identify transitions through stages. Also note that hypotension is a late sign of cardiovascular dysfunction and is not necessary for a diagnosis of shock.

Current guidelines recommend each institution develop a systematic screening tool to evaluate pediatric patients for possible sepsis.

Clinical Considerations Suggestive of Sepsis to Guide Initial Assessment

- 1. Temperature dysregulation: Fever or hypothermia
- 2. Mental status: Restless, agitated, anxious, progressive lethargy
- 3. Skin findings: petechial rash below nipple line, purpura, macular rash with mucosal changes
- 4. **Cardiovascular dysfunction**: Tachycardia (especially that does not resolve with normalization of temperature), abnormal pulses (diminished, weak, bounding), prolonged or flash capillary refill, hypotension (late finding)
- 5. **Respiratory**: Tachypnea, grunting (even in absence of pulmonary disease 2/2 metabolic acidosis)
- 6. **Presence of High-Risk medical conditions**: Patients with immunocompromise and other high risk medical conditions are especially susceptible to sepsis and may have more subtle physiologic derangements

Physical findings will vary according to the stage of shock. Frequent vital sign and physical reexamination is therefore <u>necessary</u>.

Resuscitation

Antimicrobials: Data suggests improved outcomes (including reduced hospital length of stay, shorter duration of organ dysfunction, and in some cases improved mortality) with decreasing time to antibiotic. Though the optimal time to antimicrobial is not clear, the Surviving Sepsis Guidelines for Children recommend administration as soon as possible within 1 hour for patients with septic shock and within 3 hours for those with organ dysfunction but without shock. For those without reason to suspect a specific source, combination therapy does not appear to be superior to extended-spectrum monotherapy.

Fluids: Administer crystalloid in 10-20ml/kg boluses up to 40-60ml/kg over the first hour until perfusion normalizes or signs of fluid overload develop. *Clinical reassessment should occur after each fluid bolus.* Though no high-quality RCTs exist to support this practice, many observational studies have shown improved patient outcomes with routine aggressive fluid resuscitation. Lactated ringers are preferable though NS is acceptable.

Blood culture: Obtain prior to antibiotics if possible to guide antimicrobial therapy. If blood culture is difficult, do not delay administration of antibiotics.

Lactate: Lactate is a specific but not sensitive marker for CV dysfunction in sepsis. Thresholds of 2 and 4 mmol/L have been used in children.

Vasoactives: Limited evidence from pediatric studies suggests epinephrine is superior to dopamine for fluid refractory shock, while extrapolation from adult studies also suggests norepinephrine is superior to dopamine.

Corticosteroids: No high quality studies exist demonstrating benefit for catecholamine-refractory shock in children, though it *should* be used in those who are known or suspected to be adrenally insufficient.

Intubation and Induction agents: Intubation should be considered in those with fluid-refractory, catecholamine resistant septic shock, though no RCTs exist to support this practice. Avoid etomidate as it is known to cause adrenal insufficiency. Ketamine is the preferred agent for sepsis.

Pediatric Sepsis / Septic Shock Rationale and Data

Bacterial Pathogen Consideration

Suspected Source of Sepsis								
	Lungs		Abdomen	Skin/Soft Tissue	Urinary Tract	CNS		
Major Community	Streptococcus	Esche	erichia coli	Streptococcus	Escherichia coli	Streptococcus		
Acquired	pneumoniae	Bacte	eroides	pyogenes	Klebsiella sp.	pneumoniae		
Pathogens			is	Staphylococcus	Enterobacter sp.	Neiserria		
	influenzae			aureus	Proteus sp.	meningitides		
	Legionella sp.			Clostridium sp.		Listeria		
				Pseudomonas		monocyogenes		
				aeruginosa		Escherichia coli		
						Haemophilus		
						influenzae		
Major Nosocomial	Aerobic gram	Aoro	hicaram	Stanbulacacque	Acrobic gram	Pseudomonas		
	negative bacilli		bic gram tive bacilli	Staphylococcus	Aerobic gram negative bacilli	aeruginosa		
Pathogens	negative bacili	-	robes	aureus Aerobic gram	Enterococcus sp.	Escherichia coli		
			ida sp.	negative bacilli	Enterococcus sp.	Klebsiella sp.		
		Canu	iua sp.	negative bacili		Staphylococcus sp.		
			Antihioti	C Selection				
Antibiotic therapy sho	uld be directed at th	e most		f infection. For patien	ts without a clear sou	urce of infection or		
				ter increasing risk for :				
				strated to be inferior t				
following represent re						leroblar therapy. The		
Diagnosis/Suspected S		5	Preferred Regimen		Alternative Reg	Alternative Regimen/Other		
			0		considerations			
Sepsis (respiratory, genitourinary, or no clear			Ceftriaxone 50mg/kg IV		If toxic: conside	er Vancomycin		
source)					15mg/kg IV	15mg/kg IV		
· · ·				/1 <i>.</i>				
mmunocompromised			Cefepime 50mg/kg IV		If toxic: add Va	If toxic: add Vancomycin 15mg/kg		
Central Venous Catheter, Skin/Soft Tissue			Vancomycin 15mg/kg IV			If concern for toxic shock syndrome:		
nfection, or other concern for MRSA					add Clindamyc	n 10mg/kg IV		
Meningitis			Ceftriaxone 100mg/kg IV					
			Vancomycin					
Abdominal infection			Ceftriaxone		Piperacillin-taz	obactam 100mg/kg IV		
			Metronidazo	ole 10mg/kg IV				

References:

Goldstein B, Giroir B, Randolph A, International Consensus Conference on Pediatric Sepsis. International pediatric sepsis consensus conference: definitions for sepsis and organ dysfunction in pediatrics. Pediatr Crit Care Med 2005; 6:2.

Davis AL, Carcillo JA, Aneja RK, et al. American College of Critical Care Medicine Clinical Practice Parameters for Hemodynamic Support of Pediatric and Neonatal Septic Shock. Crit Care Med 2017; 45:1061.

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