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Accounting for more than \$20 billion (5.2%) of total US hospital costs in 2011

The reported incidence of sepsis is increasing, likely reflecting aging populations with more comorbidities, greater recognition,

Sepsis is a leading cause of mortality and critical illness worldwide.

long-term physical, psychological, and cognitive disabilities with significant health care and social implications

Comparison With Other Major Diseases

Incidence of Severe Sepsis



Colon Breast CHF⁺ Severe

Cancer§

Mortality of Severe Sepsis

AIDS*

Breast

Cancer[§]

Severe

Sepsis[‡]



Sepsis[‡]

12/22/2023

<u> Cases/100,000</u>

AIDS*

Sepsis, Mortality Rates

- Overall = 30% 50%
- By syndrome definition:
 - **Sepsis = 16%**
 - -Septic shock = 46%

Sepsis is deadly



Sepsis is Common



Sepsis is increasing in incidence



Pathogenesis of SIRS/MODS



Pathogenesis of sepsis

An overview



Pathogenesis of sepsis

An overview



Inflammation

- Initial response to any pathogens is the release of pro-inflammatory mediators
 - to allow WBC to reach the infected area.
- Subsequently, an anti-inflammatory response
 - attempt to regain homeostasis and prevent "leaking capillary syndrome".
- The ability to activate and then eventually downregulate the inflammatory response to infection is a vital immune process and it is this ability that is <u>lost in sepsis</u> and severe sepsis.

Pathogenesis of sepsis

An overview



The role of the endothelium

- Release of mediators of vasodilatation and/or vasoconstriction
- Release of cytokines and inflammatory mediators
- Allows leucocytes to access infection sites
- Plays an important role in the coagulation cascade, maintaining the physiological equilibrium between coagulation and fibrinolysis



Tissue injury

Formation of fibrin clot

The role of the endothelium

- In sepsis, the regulatory function of the endothelium fails, leading to:
 - Excessive vasodilation and relative hypovolaemia
 - Leaking capillaries and generalised tissue damage
 - Tissue factor (TF) release initiates **procoagulant state**
 - Micro-thrombus formation compromising blood supply and leading to tissue necrosis
 - Inactivation of Protein C and suppression of fibrinolysis



Tissue injury

Formation of fibrin clot

Loss of homeostasis in sepsis



Pro-coagulant state

Disseminated Intravascular Coagulation (DIC)

DIC can cause:

- bleeding
- large vessel thrombosis
- haemorrhagic tissue necrosis
- microthrombi leading to organ failure

Widespread clotting causes consumption of:

- Low platelets
- clotting factors long clotting time
- fibrinogen

As a result, bleeding risk increases

Pathogenesis of sepsis

An overview



12/22/2023

Published in final edited form as: *Clin Chest Med.* 2008 December ; 29(4): 617–viii. doi:10.1016/j.ccm.2008.06.010.

The Compensatory Anti-inflammatory Response syndrome (CARS) in Critically ill patients

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Molecular Mediators in Pathophys

- Parallel to SIRS is CARS
 - Compensatory Anti-inflammatory Response System
 - Attempts to down regulate the SIRS response
 - IL-4, IL-10, transforming growth factor beta, CSF, soluble receptors to TNF, antagonists to TNF-alpha and IL-1
 - If CARS reaction is severe it will manifest as anergy and infection susceptibility



Figure Legend:

Fig. 2. Trauma-induced injury actives innate immune responses to produce pro- and antiinflammatory cytokines. Imbalance between the systemic inflammatory response syndrome and the compensatory antiinflammatory response (immunosupression) increases morbidity of trauma patients. In the first hours, the magnitude of the systemic inflammatory response syndrome is correlated with early multiple organ failure and infections. In the following days, immunosupression contributes to the increased incidence of nosocomial infections and late sepsis. CARS = compensatory anti-inflammatory response; MOF = multiple organ failure; SIRS = systemic inflammatory response syndrome.

12/22/2023

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Response

- Physiology
 - Heart rate
 - Respiration
 - Fever
 - Blood pressure
 - Cardiac output
 - WBC
 - Hyperglycemia

- Markers of Inflammation
 - TNF
 - IL-1
 - IL-6
 - Procalcitonin
 - PAF

IDENTIFYING ACUTE ORGAN DYSFUNCTION AS A **MARKER OF SEVERE SEPSIS**



Organ Dysfunction

- Lungs
- Kidneys
- CVS
- CNS
- PNS
- Coagulation
- GI
- Liver
- Endocrine

- Adult Respiratory Distress Syndrome
- Acute Tubular Necrosis
- Shock
- Metabolic encephalopathy
- Critical Illness Polyneuropathy
- Disseminated Intravascular Coagulopathy
- Gastroparesis and ileus
- Cholestasis
- Adrenal insufficiency
- Skeletal Muscle > Rhabdomyolysis





accp/sccm consensus conference

Definitions for Sepsis and Organ Failure and Guidelines for the Use of Innovative Therapies in Sepsis

 THE ACCP/SCCM CONSENSUS CONFERENCE COMMITTEE:

 Roger C. Bone, M.D., F.C.C.P., Chairman
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Alan M. Fein, M.D., F.C.C.P. William A. Knaus, M.D. Roland M. H. Schein, M.D. William J. Sibbald, M.D., F.C.C.P.

2001 SCCM/ESICM/ACCP/ATS/SIS International Sepsis Definitions Conference

Mitchell M. Levy, MD, FCCP; Mitchell P. Fink, MD, FCCP; John C. Marshall, MD; Edward Abraham, MD; Derek Angus, MD, MPH, FCCP; Deborah Cook, MD, FCCP; Jonathan Cohen, MD; Steven M. Opal, MD; Jean-Louis Vincent, MD, FCCP, PhD; Graham Famsay, MD; For the International Sepsis Definitions Conference

Terminology



- Temp > 38 or < 36
- HR > 90
- RR > 20 or PaCO2 < 32
- WBC > 12 or < 4 or Bands > 10%

Sepsis

The systemic inflammatory response to infection.

Severe Sepsis

- Organ dysfunction secondary to Sepsis.
- e.g. hypoperfusion, hypotension, acute lung injury, encephalopathy, acute kidney injury, coagulopathy.

Septic Shock

Hypotension secondary to Sepsis that is resistant to adequate fluid administration and associated with hypoperfusion.

TWO out of four criteria acute change from baseline

Bone, R., Balk, R., Cerra, F., Dellinger, R., Fein, A., Knaus, W., Schein, R., et al. (1992). Definitions for sepsis and organ failure and guidelines for the use of innovative therapies in sepsis. The ACCP/SCCF/Gordenses 6002itee. American College of Chest Physicians/Society of Critical Care Medicine. *Chest*, 101(6), 1644–1655.



The Third International Consensus Definitions for Sepsis and Septic Shock (Sepsis-3)

The Sepsis Definitions Task Force

Special Communication | CARING FOR THE CRITICALLY ILL PATIENT

The Third International Consensus Definitions for Sepsis and Septic Shock (Sepsis-3)

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Editorial page 757

INFORTANCE Definitions of sepsis and septic shock were last revised in 2001. Considerable advances have since been made into the pathobiology (changes in organ function, morphology, cell biology, biochemistry, immunology, and circulation), management, and epidemiology of sepsis, suggesting the need for resumination. Author Video interview, Author Audio interview, and JAMA Report Video at Jama.com

DRJECTIVE. To evaluate and, as needed, update definitions for sepsis and septic shock.

CME Quiz at jamanetworkcime.com and CME Questions page 816

PRICESS A task force (in = 19) with expertise in sepsis pathobiology, clinical trials, and epidemiology was conversed by the Society of Critical Care Medicine and the European Society of Intensive Care Medicine. Definitions and clinical criteria were generated through meetings. Delphi processes, analysis of electronic health record databases, and voting, followed by circulation to international professional societies, requesting peer review and endorsement (by 3) societies listed in the Acknowledgment).

NOV FINITIALS FINANCE SYNCHESES Limitations of previous definitions included an eccessive focus on inflammation, the misleading model that sepsis follows a continuum through severe sepsis to shock, and indequate specificity and simultivity of the systemic inflammatory response syndrome (SIRS) criteria. Multiple definitions and terminologies are currently in use for sepsis, septic shock, and organ dysfunction, leading to discrepancies in reported incidence and observed mortality. The task force concluded the term severe sepsis was redundant.

INCOMMENDATIONS Separa should be defined as life-threatening organ dysfunction caused by a dysregulated host response to infection. For clinical operationalization, organ dysfunction can be represented by an increase in the Sequential (Spssie-related) (Organ Failure Assessment (SOFA) score of 2 points or more, which is associated with an in-hospital mortality greater than 10%. Septic shock should be defined as a subset of sepsis in which particularly profound circulatory, cellular, and metabolic abnormalities are associated with a greater risk of mortality than with sepsis alone. Patterns with septic shock can be clinically identified by a vasopressor requirement to maintain a mean arterial pressure of 65 mm Hg or greater and serum lactate level greater than 2 mmol/L (>48 mg/dL) in the absence of hypovolemia. This combination is associated with hospital mortality rates greater than 40%. In our of hospital, emergency department, or general hospital ward setting, adult patients with suspected infection can be tapidly identified as being more likely to have poor outcomes typical of sepsis if they have at least 2 of the following clinical criteria that together constitute a new bedside clinical score termed quickSOFA (spOFA) respiratory rate of 22/min or greater, ahared mentation, or systic blood pressure of 100 mm Hg or less.

CONCLUSIONS AND RELEVANCE. These updated definitions and clinical criteria should replace previous definitions, offer greater consistency for epidemiologic studies and clinical trials, and facilitate earlier recognition and more timely management of patients with sepsis or at risk of developing sepsis.

JAMA. 20%-3/501-807-810. doi:10.2007/jama.20%.0287

The Document

Singer M, Deutschman CS, Seymour CW, Shankar-Hari M et al.

Third International Consensus Definitions for Sepsis and Septic Shock (Sepsis-3) JAMA 2016; 315: 801-10

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The Definition of Sepsis

Sepsis is life-threatening organ dysfunction caused by a dysregulated host response to infection The Definition of Septic Shock

□ What tangibly differentiates septic shock from sepsis ?

□ MORTALITY

Septic shock is "really bad" sepsis

Septic shock is a subset of sepsis in which profound circulatory, cellular and metabolic abnormalities are associated with a greater risk of mortality than with sepsis alone

Clinical criteria for sepsis

Sequential [Sepsis-Related] Organ Failure Assessment (SOFA) Score

System	0		2	3	4
Respiration PaO2/FiO2, mmHg (kPa)	≥400 (53.3)	<400 (53.3)	<300 (40)	<200 (26.7) with respiratory support	< 100 (13.3) with respiratory support
Coagulation Platelets, xl0³/uL	≥ 50	<150	<100	<50	<20
Liver Bilirubin, mg/dL (umol/L)	<i.2 (20)<="" td=""><td>1.2 - 1.9 (20 - 32)</td><td>2.0 - 5.9 (33 - IOI)</td><td>6.0 - II.9 (102 - 204)</td><td>>12.0 (204)</td></i.2>	1.2 - 1.9 (20 - 32)	2.0 - 5.9 (33 - IOI)	6.0 - II.9 (102 - 204)	>12.0 (204)
Cardiovascular	MAP ≥70mmHg	MAP <70mmHg	Dopamine <5 or Dobutamine (any dose)	Dopamine 5.1 - 15 or Epinephrine <0.1 or Norepinephrine <0.1	Dopamine >15 or Epinephrine >0.1 or Norepinephrine >0.1
CNS GCS Score	15	13 - 14	IO -I2	6 - 9	<6
Renal Creatinine, mg/dL (umol/L) Urine Output, mL/d	<1.2 (110)	I.2 - I.9 (110 - 170)	2.0 - 3.4 (171 - 299)	3.5 - 4.9 (300 - 440) <500	>5.0 (440) <200
*Catecholamine Doses = ug/kg/min for at least Ihr					
SOFA Score

The European Society of Intensive Care Medicine

SOFA score	0 1		2	3	4	
Mortality	SOFA score		<300 142-220	<200 67-141	<100 <67	
<10%	0-6 7-9 10-12		Mortali	ty Sco	Score trend (First 48 hrs) Increasing	
15-20%				(Firs		
40-50%			>50%	Inc		
50-60%	13-14	4	27-359	6 Unc	Unchanged	
>80%	15		<27%	Dec	Decreasing	
>90%	15-24	4	2.0-3.4	3.5-4.9 or	5-4.9 or >5.0 or <20	
or unne output (mva)						

Clinical criteria for sepsis

□ Infection plus 2 or more SOFA points (above baseline)

Please visit www.qsofa.org

Clinical criteria for sepsis

□ Infection plus 2 or more SOFA points (above baseline)

Prompt outside the ICU to consider sepsis

Please visit www.qsofa.org



Clinical criteria for sepsis

□ Infection plus 2 or more SOFA points (above baseline)

Prompt outside the ICU to consider sepsis

□ Infection plus 2 or more qSOFA points

Please visit www.qsofa.org

- Outside the ICU, patients with suspected or presumed infection who are highly likely to have poor outcomes can be clinically identified using qSOFA
 SBP < 100mm Hg
 - □ RR > 22 breath/min
 - □ Altered mental status
- In the ICU, patients with suspected or presumed infection who are highly likely to have poor outcomes can be clinically identified by the presence of 2 or more SOFA points

Patients with a SOFA score of 2 or more had an overall mortality risk of approximately 10% in a general hospital population with presumed infection.

2016 Septic Shock Criteria





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Mortality of Septic shock exceeds 40 %



SOFA Score	0	1	2	3	4
paO ₂ /FiO ₂ (mmHg)	> 400	≤ 400	≤ 300	≤ 200 with respiratory support	≤ 100 with respiratory support
Platelets x10 ³ /mm ³	> 150	≤ 150	≤ 100	≤ 50	≤ 20
Bilirubin (mg/dL)	< 1.2	1.2 - 1.9	2.0 - 5.9	6.0-11.9	≥ 12.0
Hypotension	No hypotension	MAP < 70 mmHg	Dopamine ≤ 5 <i>or</i> Dobutamine (any dose)	Dopamine > 5 or Epinephrine $\leq 0.1 \text{ or}$ Norepinephrine ≤ 0.1	Dopamine > 15 or Epinephrine > 0.1 or Norepinephrine > 0.1
Glasgow Coma Score	15	13 - 14	10-12	6 - 9	< 6
Creatinine (mg/dL) or Urine output	< 1.2	1.2 - 1.9	2.0 - 3.4	3.5 – 4.9 <i>or</i> < 500 mL/d	> 5.0 <i>or</i> < 200 mL/d

Why do Septic Patients Die?

Organ Failure



Organ Failure and Mortality

•Knaus, et al. (1986):

•Direct correlation between number of organ systems failed and mortality.

•Mortality Data:

#OSF	D1	D2	D3	D4	D5	D6	D7
1	22%	31%	34%	35%	40%	42%	41%
2	52%	67%	66%	62%	56%	64%	68%
3	80%	95%	93%	96%	100	100%	100%
			Fourth Yea	r Lectures 2021	%		

SEVERE SEPSIS-ASSOCIATED MORTALITY INCREASES WITH THE NUMBER OF ORGAN DYSFUNCTIONS



Organ Dysfunctions

Angus DC, et al. Crit Care Med. 2001;29:1303-1310. Vincent JL, et al. Crit Care Med. 1998;21:1793-1800. Fourth Year Lectures 2021

Evolution of Sepsis care

Established Core Rx: Source Control Antibiotics Resuscitation Supportive Care

Steroids

Established Core Rx: Source Control More Antibiotics Faster Resuscitation Better Supportive Care

In general the process of care has improved

No Steroids Endotoxin Antagonist igris LPS/LPS receptor a **highorfully compile Cioints** of anti-TNF NSAIDs Nitric Oxide Synthase Inhibitors Tissue Factor Pathway Inhibitors anti-TLR4

Mortality

Loosen Bilydder Bikieg Giddist Pol

Surviving Sepsis Campaign: International Guidelines for Management of Severe Sepsis and Septic Shock: 2012

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Dellinger RP, Levy MM, Rhodes A, et al. Surviving Sepsis Campaign: International Guidelines for Management of Severe Sepsis and Septic Shock: 2012. Critical Care Medicine 2013;41(2):580–637.

How do we manage sepsis and septic shock?

- 1) Investigate and treat sepsis
 - Try and find and treat source
 - Early blood cultures
 - Start antibiotics asap ideally within 1 hour and after cultures taken
- 2) Assess extent of end organ hypoperfusion and improve oxygen delivery

2005

6-hour Resuscitation Bundle

- Measure serum lactate
- Obtain blood cultures prior to antibiotics
- Administer broad spectrum antibiotics within 3 hours of ED or 1 hour non-ED admission
- With hypotension &/or serum lactate > 4 mmol/L:
 - Crystalloid 20ml/Kg
 - Vasopressors if unresponsive
- Persistent hypotension &/or lactate > 4 mmol/L achieve:
 - CVP ≥ 8 mm Hg
 - ScvO2 ≥ 70 % or SvO2 ≥ 65%

24-hour Management Bundle

- Low dose steroids
- Human activated protein C (rhAPC)
- Maintain glucose 70 -150 mg/dL
- Maintain median inspiratory plateau pressure < 30 cm H2O in mechanical ventilation

2013

3-hour Bundle

- Measure serum lactate
- Obtain blood cultures prior to antibiotics
- Administer broad spectrum antibiotics
- With hypotension &/or serum lactate > 4 mmol/L:
 - Crystalloid 30ml/Kg

6-hour Bundle

- Vasopressors for hypotension after fluid
- For persistent arterial hypotension after fluid or with lactate > 4 mmol/L;
 - Measure CVP
 - Measure ScvO2

24-hour Bundle no longer recommended

2018

1-hour Bundle

- Measure serum lactate. Remeasure if initial > 2 mmol/L
- Obtain blood cultures prior to antibiotics
- Administer broad spectrum antibiotics
- Begin rapid crystalloid 30 ml/kg
- Apply vasopressors if hypotension remains after fluid resuscitation to MAP <u>></u> 65 mm Hg

1 Hour Bundle



WWW.SEPSISTRUST.ORG Design by hugo beaumont

CX **1.GIVE O2 TO KEEP SATS ABOVE 94% 2.TAKE BLOOD CULTURES 3.GIVE IV ANTIBIOTICS 4.GIVE A FLUID CHALLENGE 5.MEASURE LACTATE**

6.MEASURE URINE OUTPUT

SEPSIS TRUST

THE SEPSIS

Surviving Sepsis targets of fluid resuscitation

What are they?

- SBP
- MAP
- CVP
- U/o
- Lactate
- ScvO2
- HCt

Surviving Sepsis targets of fluid resuscitation

What are they?

- SBP > 90
- MAP > 65
- CVP 8 12
- U/o > 0.5 ml/kg/hr
- Lactate < 1
- ScvO2 >70
- HCt > 30

• 30 mL/kg of IV crystalloid fluid be given within the first 3 h

 additional fluids be guided by frequent reassessment of hemodynamic status (BPS)

- Crystalloids are favored as the initial fluid
- Hydroxyethyl starches are likely harmful
- Albumin may have a role, particularly if alot of fluid is given

Markers of perfusion

What are they?

- Clinical signs
 - Warm skin, conscious level, u/o
- Haemodynamic variables
 - -CVP
- Bloods
 - Serum Lactate
 - ScvO2

CVP

CVP

What does it mean? Starling's Law Estimate of LVEDV (i.e. preload) Not always a good correlation with volume-responsiveness However if low strongly suggestive of hypovolaemia

Lactate

Lactate

- Increased production (anaerobic glycolysis)
 - Tissue hypoperfusion
 - Tissue dysoxia
- Reduced metabolism
 - Hepatic
 - Renal
- <1 is normal, 1-2 is a concern, >2 is bad,
 >4 is very bad

What does it mean?

- Balance between oxygen delivery and consumption (VO2)
- ScvO2 = SaO2 <u>VO2</u>

CO

• Target > 70%

What can I do if it's low?

What can I do if it's low?

Delivery = [Hb] x SpO2 x 1.34 x HR x SV

What can I do if it's low? Delivery = $[Hh] \times SnO2$

Delivery = [Hb] x SpO2 x 1.34 x HR x SV

Fluid optimise

Transfuse packet cells

HCt > 30%

Inotropes

"Time Zero"

- Time Zero = time of presentation
 - -ED, Medical Floors, ICU
- 1 Hour Bundle

microbiologic cultures (including blood) be obtained before starting antimicrobial therapy in patients with suspected sepsis or septic shock if doing so results in no substantial delay in the start of antimicrobials.
Antibiotic therapy

- intravenous antimicrobial therapy as early as possible and within the first hour of recognition
- empiric broad-spectrum therapy with one or more antimicrobials to cover all likely pathogens (including bacterial and potentially fungal or viral coverage)
- antimicrobial therapy to be narrowed once pathogen identification and sensitivities are established and/or adequate clinical improvement is noted.

Hospital Mortality by Time to Antibiotics

Time to ABX ¹ , hrs	OR ²	95% CI		<i>p</i> -value	Probability of mortality ³	95% CI	
0 (ref)	1.00				18.7	17.5	19.9
1	1.05	1.02	1.07	< 0.001	19.3	18.3	20.4
2	1.09	1.04	1.15	< 0.001	20.0	19.1	21.0
3	1.14	1.06	1.23	< 0.001	20.8	19.7	21.8
4	1.19	1.08	1.32	< 0.001	21.5	20.3	22.8
5	1.25	1.11	1.41	< 0.001	22.3	20.7	23.9
6	1.31	1.13	1.51	< 0.001	23.1	21.2	25.1
¹ Time to ABX i ² Hospital mort	s based or tality odds	n 15,948 ob ratio refer	servation ent group	s that are gre is 0 hours fo	eater than or equal t r the time to ABX ar	o zero nd is adjust	ed by the

(Europe, North America, and South America)

Septic Shock: Timing of Antibiotics



12/22/2023

Kumar Crit Care Med 2006

Source Control

a specific anatomic diagnosis of infection requiring emergent source control to be identified or excluded as rapidly as possible and that any required source control intervention be implemented as soon as medically and logistically practical after the diagnosis is made.

Vasoactive agents

• Norepinephrine is the first choice vasopressor

CORTICOSTEROIDS

intravenous hydrocortisone to treat septic shock patients if adequate fluid resuscitation and vasopressor therapy are UNABLE to restore hemodynamic stability.

GLUCOSE CONTROL

We recommend a protocolized approach to blood glucose management in ICU patients This approach should target an upper blood glucose level ≤180 mg/dL

- Hit fast and hit Hard
- IV fluids
- Antibiotics
- Source control

Thank You