SIRS, Sepsis, and MODS

Amjad Bani Hani Asst. Prof of Cardiac Surgery and Intensive Care In 1992, the American College of Chest Physicians (ACCP) and the Society of Critical Care Medicine (SCCM) introduced definitions

Definitions

- Infection
- Systemic Inflammatory Response Syndrome (SIRS)
- Sepsis
- Severe Sepsis
- Septic Shock

Infection: Part of a bigger picture

· Infection:

- Presence of organisms in a closed space or location where not normally found

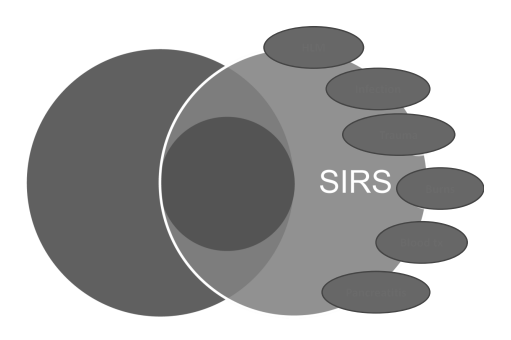
Infection

Definitions (ACCP/SCCM):

 Infection: A microbial phenomenon characterized by an inflammatory response to the presence of microorganisms or the invasion of normally sterile host tissue by those organisms.

SIRS

- self-defense mechanism.
- Inflammation is the body's response to nonspecific insults that arise from chemical, traumatic, or infectious stimuli.
- The inflammatory cascade is a complex process that involves humoral and cellular responses, complement, and cytokine cascades.



'Sepsis'

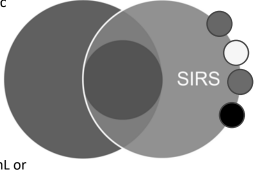
- Sepsis is the systemic response to infection
 - **SIRS** in the presence of proven or suspected infection

SIRS: Systemic Inflammatory Response Syndrome

 SIRS: A clinical response arising from a nonspecific insult manifested by
 ≥2 of the following:

Temperature≥38°C or ≤36°C

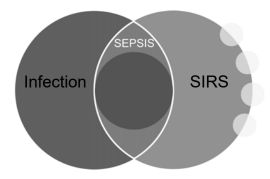
- HR ≥90 beats/min
- Respirations ≥20/min, Paco2 ≥ 32
- WBC count ≥12,000/mL or ≤4,000/mL or >10% immature neutrophils



Adapted from: Bone RC et al. *Chest*. 1992;101:1644-55. Opal SM et al. *Crit Care Med*. 2000;28:S81-2.

Sepsis: More Than Just Inflammation

- Sepsis:
 - SIRS criteria
 - Known or suspected infection



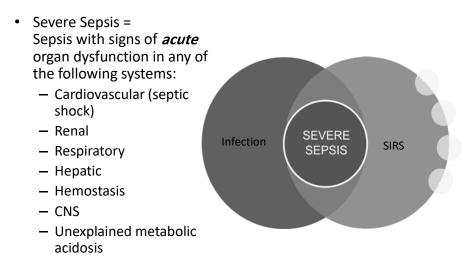
'Severe Sepsis'

- Sepsis associated with
- Organ-dysfunction
 - -Hypotension
 - -Hypoperfusion

'Septic Shock'

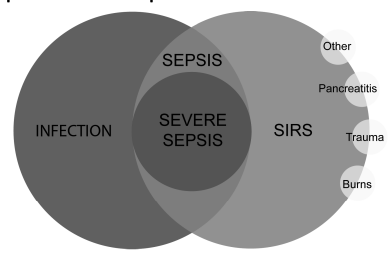
- Sepsis with hypotension despite adequate fluid resuscitation
- Include all patients on vasopressors or inotropic support

Severe Sepsis: Acute Organ Dysfunction



Adapted from: Bone RC et al. Chest. 1992;101:1644-55.

Sepsis: A Complex Disease

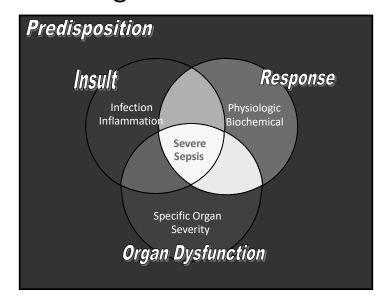


Adapted from: Bone RC et al. *Chest.* 1992;101:1644-55. Opal SM et al. *Crit Care Med.* 2000;28:S81-2.

The Multiple Organ Dysfunction Syndrome (MODS)

 The development of potentially reversible physiologic derangement involving two or more organ systems not involved in the disorder that resulted in ICU admission

Jargon 2002: PIRO



The Multiple Organ Failure Syndrome (MOFS)

 The development of potentially irreversible physiologic derangement involving two or more organ systems not involved in the disorder that resulted in ICU admission

Predisposition

- Pre-existing disease
 - Cardiac, Pulmonary, Renal
 - HIV
- Age (extremes of age)
- Gender (males)
- Genetics
 - TNF polymorphisms (TNF promoter high secretor genotype)

Response

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

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

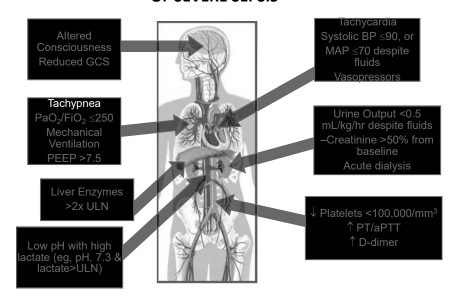
Organ Dysfunction

- Lungs > Adult Respiratory Distress Syndrome
- Kidneys
 Acute Tubular Necrosis
- CVS ➤ Shock
- CNS
 Metabolic encephalopathy
- Coagulation ➤ Disseminated Intravascular Coagulopathy
- Gl

 ➤ Gastroparesis and ileus
- Liver > Cholestasis
- Endocrine > Adrenal insufficiency
- Skeletal Muscle > Rhabdomyolysis

✓ Specific therapy exists

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



Magnitude of the Problem

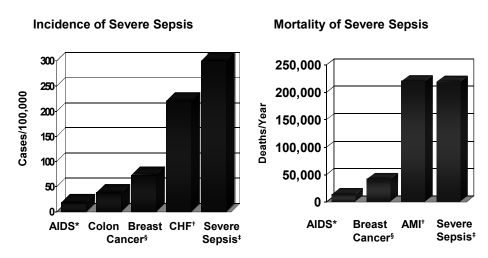
- Severe sepsis takes more lives than breast, colon/rectal, pancreatic, and prostate cancer combined.
- One of every three patients who develop severe sepsis will die within a month.

Source: Society of Critical Care

Sepsis, Mortality Rates

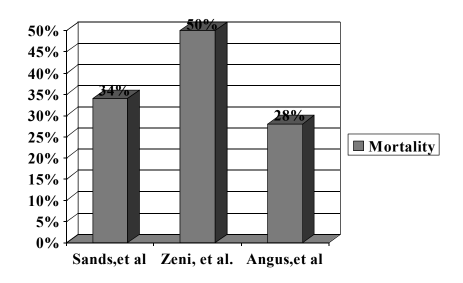
- Overall = 30% 50%
- By syndrome definition:
 - -SIRS = 4-7%
 - **-Sepsis = 16%**
 - -Severe sepsis = 20%
 - -Septic shock = 46%

Comparison With Other Major Diseases

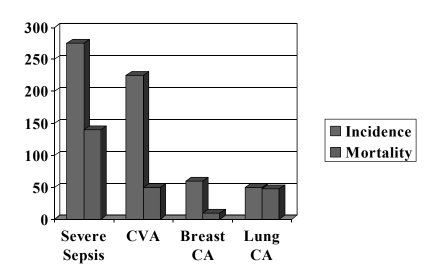


[†]National Center for Health Statistics, 2001. [§]American Cancer Society, 2001. ^{*}American Heart Association. 2000. [‡]Angus DC et al. *Crit Care Med.* 2001;29(7):1303-1310.

Severe Sepsis is deadly



Severe Sepsis is Common



Epidemiology of Sepsis The International Cohort Study

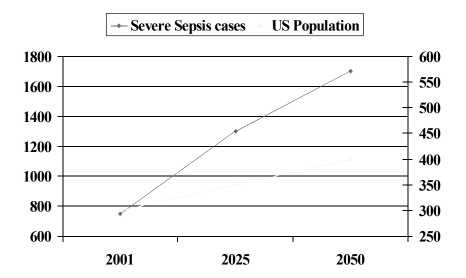
Infection Sepsis	Severe Sepsis	Septic Shock
------------------	------------------	-----------------

Percent of cases within each category

18	28	24	30
		35% m	ortality

8353 patients with LOS > 24h 4277 infections (2696 on admission)

Severe Sepsis is increasing in incidence



Sources of Sepsis
The International Cohort Study

	Severe Sepsis	Septic Shock
Respiratory	66	53
Abdomen	9	20
Bacteremia	14	16
Urinary	11	11
Multiple	-	-

Microbiology of Sepsis The International Cohort Study

	Severe Sepsis	Septic Shock
Gram-positive	44	40
Gram-negative	47	47
Fungal	9	13
Polymicrobial	-	-

Etiology

- Infectouse
- Non-Infectouse

Infectouse

- Bacterial sepsis
- Burn wound infections
- Candidiasis
- Cellulitis
- Cholecystitis
- Community-acquired pneumonia^[3]
- Diabetic foot infection
- Erysipelas
- · Infective endocarditis
- Influenza

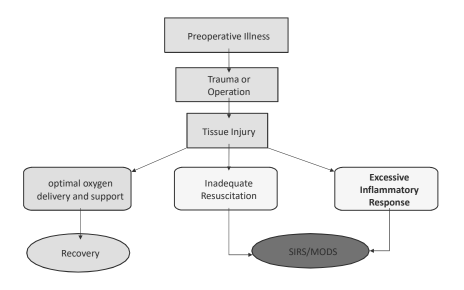
- Intraabdominal infections -Eg, diverticulitis, appendicitis
- Gas gangrene
- Meningitis
- · Nosocomial pneumonia
- Pseudomembranous colitis
- Pyelonephritis
- · Septic arthritis
- Toxic Schock Syndrom
- Urinary tract infections (male and female)

Noninfectious

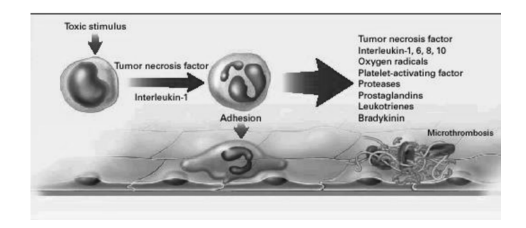
- Acute mesenteric ischemia
- Adrenal insufficiency
- Autoimmune disorders
- Burns
- Chemical aspiration
- Cirrhosis
- Cutaneous vasculitis
- Dehydration
- Drug reaction
- · Electrical injuries
- Erythema multiforme
- Hemorrhagic shock

- Hematologic malignancy
- Intestinal perforation
- Medication side effect Eg, from theophylline
- Myocardial infarction
- Pancreatitis^[4]
- Seizure
- Substance abuse Stimulants such as cocaine and amphetamines
- Surgical procedures
- Toxic epidermal necrolysis
- · Transfusion reactions
- Upper gastrointestinal bleeding
- Vasculitis

Pathogenesis of SIRS/MODS



Initiation of Inflammatory Response



From Wheeler & Bernard, NEJM 1999

Pro-inflammatory Mediators

- Bacterial Endotoxin
- TNF-α
- Interleukin-1
- Interleukin-6
- Interleukin-8
- Platelet Activating Factor (PAF)
- Interferon-Gamma
- Prostaglandins
- Leukotrienes
- Nitric Oxide

Anti-inflammatory Mediators

- Interleukin-10
- PGE2
- Protein C
- Interleukin-4
- Interleukin-12
- Lipoxins
- GM-CSF
- TGF
- IL-1RA

Pathophysiology of Sepsis

- In simple terms sepsis can be viewed as an imbalance of inflammation, coagulation, and fibrinolysis.
- In normal patients homeostasis is maintained when these are balanced.





Pathophysiology of Sepsis

During a normal response to bacteria in the blood the immune system releases inflammatory mediators to promote recovery of the tissue.

These mediators are known as:

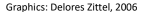
- Tumor Necrosis Factor (TNF)
- Interleukins (IL)
- Cytokines
- Prostaglandins
- Platelet Activating Factor

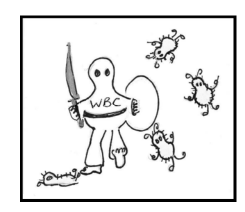
Source: New England Journal of Medicine, 2003



Pathophysiology of Sepsis

Once the bacteria or antigen is isolated, the pro-inflammatory mediators attract neutrophils or WBCs which attack the antigen and try to engulf it.





Pathophysiology of Sepsis

To prevent the response from damaging normal tissue, anti-inflammatory mediators are released including transforming growth factors and interleukins (IL-4). This balance of inflammatory and anti-inflammatory mediators restricts the inflammation response to the local site of infection.



upplement, 2004

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

Nicholas S. Ward, MDa,*, Brian Casserly, MDa, and Alfred Ayala, PhDb

^aDivision of Pulmonary, Critical Care, and Sleep Medicine, The Warren Alpert Medical School of Brown University, 593 Eddy Street, APC 707, Providence, RI 02912, USA

^bDivision of Surgical Research, Department of Surgery, The Warren Alpert Medical School of Brown University. Providence. RI 02912. USA

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

Cellular/molecular elements

Lymphocyte dysfunction (ie, reduced proliferative and/or type 1 helper T-cell [Th1] cytokine production in response-defined antigens or specific T-cell stimuli)

Lymphocyte Apoptosis

Down-regulation of monocyte HLA receptors Monocyte deactivation (ie, reduced Th1/proinflammatory cytokine production in response stimuli)

IL-10 production

Transforming growth factor-beta production Prostaglandin E2 production

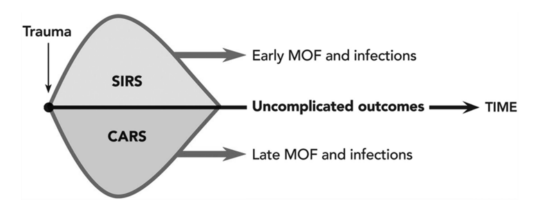


Figure Legend:

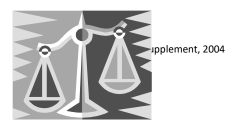
Fig. 2. Trauma-induced injury actives innate immune responses to produce pro- and antiinflammatory cytokines. Imbalance between the systemic inflammatory response syndrome and the compensatory antiinflammatory response (immunosupression) increases morbidity of trauma patients. In the first hours, the 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; MOE = multiple organ failure; SIRS = systemic inflammatory response syndrome.

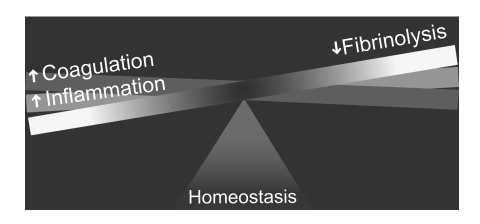
Pathophysiology of Sepsis

When the body is unable to maintain the appropriate balance, the immune response is no longer local but becomes systemic.

Inflammation and altered clotting quickly spread through the body.



Homeostasis Is Unbalanced in Severe Sepsis



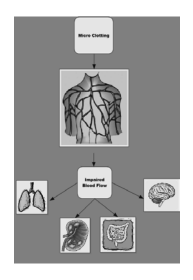
Pathophysiology of Sepsis

- The release of the inflammatory mediators starts the Coagulation Cascade leading to the development of a clot.
- To maintain this clot, inhibitors are released to suppress fibrinolysis or breakdown. This is necessary to have time for the body to destroy the bacteria before the clot is gone.

Source: Critical Care Nurse Supplement, 2004

Activation of Coagulation

The enhanced clotting continues making tiny clots or "microthrombi" in the vascular system which impairs blood flow and organ perfusion.



Activation of Fibrinolysis

Fibrinolysis, or the breakdown of clots, is the body's response to the increased clotting and inflammation.

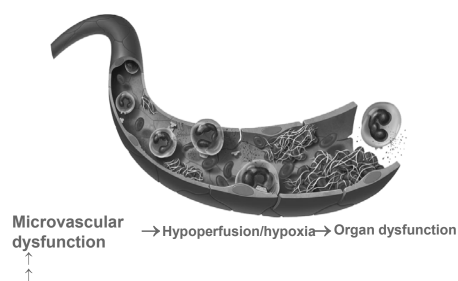
In sepsis this breakdown is inhibited or slowed through

- Plasminogen Activator Inhibitor-1 (PAI-1)
- Thrombin Activatable Fibrinolysis Inhibitor (TAFI)

Activation of Fibrinolysis

The increase levels of these two inhibitors,
Plasminogen Activator Inhibitor-1(PAI-1) and
Thrombin Activatable Fibrinolysis Inhibitor
(TAFI), suppress fibrinolysis even more creating a state of "coagulopathy".

SEVERE SEPSIS PATHOPHYSIOLOGY



Making Matters Worse The Role of Endothelium in Sepsis

Normal endothelium has anticoagulant abilities and plays a role in the body's homeostasis abilities including:

- Vasomotor tone
- Movement of cells and nutrients
- Maintaining blood fluidity

When activated, endothelium also plays a role in the inflammatory, coagulation, and fibrinolytic components of sepsis.

Making Matters Worse

- In sepsis the endothelium becomes damaged which makes the "inflammatory process" worse by releasing more cytokines (TNF-a and IL-1) causing neutrophils to stick to its' lining.
- The "activation" of the capillary endothelium leads to increased permeability causing fluid to "leak" out of the capillaries and into the extracellular spaces.

Source: http://www.xigris.com/Learning_Modules/course_01/module_02/index.htm

Organ Failure and Mortality

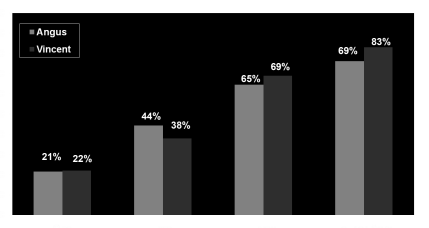
- •Knaus, et al. (1986):
 - Direct correlation between number of organ systems failed and mortality.
 - Mortality Data:

#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
					%	%	%

Question: Why do Septic Patients Die?

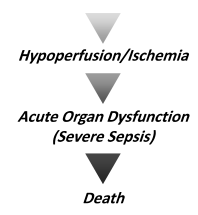
Answer: Organ Failure

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



Severe Sepsis: The Final Common Pathway

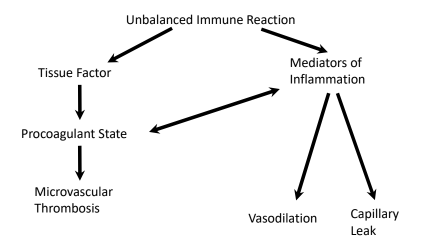
Endothelial Dysfunction and Microvascular
Thrombosis



How do you Quickly deliver complex care?

Mobilization and coordination of people and resources.

Sepsis Pathogenesis



System-based Approaches to sepsis

The New England Journal of Medicine

EARLY GOAL-DIRECTED THERAPY IN THE TREATMENT OF SEVERE SEPSIS AND SEPTIC SHOCK

EMANUEL RIVERS, M.D., M.P.H., BRYANT NGUYEN, M.D., SUZANNE HAVSTAD, M.A., JULIE RESSLER, B.S.,
ALEXANDRIA MUZZIN, B.S., BERNHARD KNOBLICH, M.D., EDWARD PETERSON, PH.D., AND MICHAEL TOMLANOVICH, M.D.,
FOR THE EARLY GOAL-DI

System-based Approaches to sepsis

■ Early-Goal Directed Therapy

■ INCLUSION = SEPSIS AND [BP < 90 after fluid OR Lactate > 4]

Control	Intervention	EGDT
CVP 8-12	Fluids	CVP 8-12
MAP > 65	Vasopressors	MAP > 65
	Transfusions Dobutamine	ScvO2 > 70%
49% mortality		33% mortality

Rivers, E., Nguyen, B., Havstad, S., Ressler, J., Muzzin, A., Knoblich, B., Peterson, E., et al. (2001). Early goal-directed therapy in the treatment of severe sepsis and septic shock. New England Journal of Medicine, 345(19), 1368–1377.

System-based Approaches to sepsis

Control	EGDT
49% mortality	33% mortality
Do whatever you normally do.	Use a rigid protocol with multiple dedicated team members

They did not control for the system of care.

System-based Approaches to sepsis

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FOR THE EARLY GOAL-DI

Used to promote:

- 1. CVP > 8 as an initial target
- 2. Use of Svo2 monitoring and use of blood/dobutamine

Rivers, E., Nguyen, B., Havstad, S., Ressler, J., Muzzin, A., Knoblich, B., Peterson, E., et al. (2001). Early goal-directed therapy in the treatment of severe sepsis and septic shock New England Journal of Medicine, 345(19), 1368–1377.

A Multidisciplinary Community Hospital Program for Early and Rapid Resuscitation of Shock in Nontrauma Patients

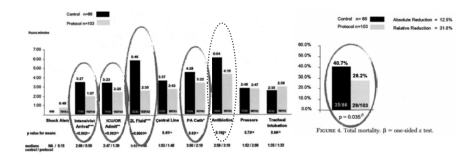
BEFORE (control)

AFTER (protocol)

Do what you normally do. We will be watching.

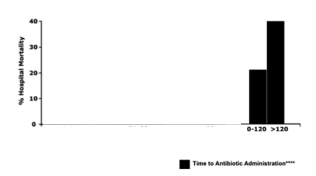
Screening Protocol, Educational Initiative, Shock Team, Treatment Protocols.

A Multidisciplinary Community Hospital Program for Early and Rapid Resuscitation of Shock in Nontrauma Patients

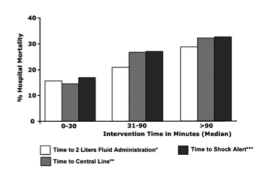


Sebat, F., Johnson, D., Musthafa, A. A., Watnik, M., Moore, S., Henry, K., & Saari, M. (2005). A multidisciplinary community hospital program for early and rapid resuscitation of shock in nontrauma patients. Chest, 1275), 1729—1743.

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Hospital-wide impact of a standardized order set for the management of bacteremic severe sepsis

BEFORE AFTER

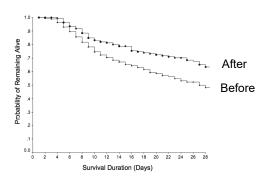
Do whatever it is that you normally do. We will be watching.

All physicians, nurses, and patient care technicians in the emergency department and intensive care units received formal order set clinical education. Additionally, all hospital floor clinical nurse specialists and advance practice nurses, along with the house staff physicians in these areas, were in-serviced on the order sets...These educational endeavors included training in sepsis pathophysiology, monitoring of central venous pressures, assessment of central venous blood oxygen saturation, and the pharmacotherapy of sepsis

1. EDUCATION

2. ORDER SET with recommendations and goals for sepsis treatment.

Hospital-wide impact of a standardized order set for the management of bacteremic severe sepsis



Thiel, S. W., Asghar, M. F., Micek, S. T., Reichley, R. M., Doherty, J. A., & Kollef, M. H. (2009). Hospital-wide impact of a standardized order set for the management of bacteremic severe sepsis*. Critical Care Medicine. 37(3), 819–824. doi:10.1097/CCM.0b013e318196206b

Acute Phase

- Identify Sepsis as early as possible
- Broad Spectrum antibiotics ASAP and Identify source(s) of infection
- Identify severity: Vitals, mental status, UOP, LACTATE, other labs.
- Volume and physiologic resuscitation ASAP with GOALS.
- Tweak your system so these things happen FAST

Summary of Trials

	Rivers 2001 RCT	Sebat 2005 Before-After	Nguyen 2007 Complete or Not	Thiel 2009 Before-After	Levy 2011 Before-After
Goals	CVP >8 MAP > 65 ScVO2 >70% HCT >30	MAP > 70 SaO2 > 92 UOP > 30ml/h SvO2 > 60 Cl > 2.5	ABX in 4 h CVP > 8, MAP > 65, ScVO2 > 70%, HCT > 30 Check Lactate Steroids	Appropriate ABX in 4 h, CVP > 8, MAP > 65, ScVO2 > 70%	Early ABX, Blood Cultures, Appropriate ABX, CVP > 8, MAP > 65, SvO2 > 70%
Specific Interventions	Fluids, Blood, Pressors	ABX, Fluids Pressors	ABX, Fluids, Blood, Pressors	ABX, Fluids, Pressors, Steroids, Xigris, Other Supportive Care	ABX, Fluids, Pressors, Steroids, Xigris, Other Supportive Care
System Interventions	ED-based Sepsis Team	Screening, Education, Shock Team, Protocols	Education, Inservices, Protocols	Education, Inservices, Order Set, Protocols	Screening, Education, Order Sets
Absolute Change in Mortality	-16%	-12%	-19%	-16%	-7%

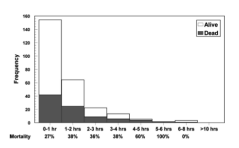
Sepsis Identification

- Train all providers
- Vital sign/Laboratory alerting systems
- ?Biomarkers

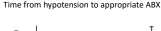
Antibiotics

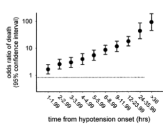
No randomized-controlled data

Time from EDGT qualification to ABX



Gaieski DF, Mikkelsen ME, Band RA, et al. Impact of time to antibiotics on survival in patients with severe sepsis or septic shock in whom early goal-directed therapy was initiated in the emergency department*. Critical Care Medicine 2010:38(4):1045-53.





Kumar A. Roberts D. Wood KE, et al. Duration of hypotension before initiation of effective antimicrobial therapy is the critical determinant of survival in human septic shock*. Critical Care Medicine 2006;34(6):1589-96.

Defining the severity of sepsis

- Importance of looking for organ failure is self evident.
- Identification of "shock" dramatically alters the treatment and mortality.
 - Blood Pressure, Response to Fluid, LACTATE

Source Control

- Don't be satisfied with a diagnosis of sepsis and no source.
- If a source exists and is potentially removable, get the ball rolling.

Goals in resuscitation

Early, quantitative resuscitation goals vs. standard care have resulted in improved mortality

The effect of a quantitative resuscitation strategy on mortality in patients with sepsis: A meta-analysis *. Jones, Alan E. MD; Brown, Michael D. MD, MSc; Trzeciak, Stephen MD, MPH; Shapiro, Nathan I. MD, MPH; Garrett, John S. MD; Heffner, Alan C. MD; Kline, Jeffrey A. MD; on behalf of the Emergency Medicine Shock Research Network investigators Critical Care Medicine. 36(10):2734-2739, October 2008.

Surviving Sepsis targets of fluid resuscitation

What are they?

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

Resuscitation

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

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

Chronic Phase

- Monitor for and prevent recurrence of sepsis
 - VAP, CLABSI, UTI. Infection Control Practices.
- Lung Protective Ventilator Strategies
- Protocolized Sedation, Daily Awakenings
- Nutritional Support
- Early Mobilization
- Success with these measures is most likely with a multi-disciplinary approach.

Summary

- System-based strategies are effective for improving sepsis care
- Processes should aim to:
 - Identify patients early and identify the severity of sepsis
 - Quickly administer appropriate antibiotics and source control
 - Establish institutional goals for physiologic resuscitation
 - Multidisciplinary chronic phase of care to ensure compliance

Oxygen delivery

What does it mean?

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 (early goal directed therapy)

Oxygen delivery

What does it mean?

Delivery (DO2) = O2 content x cardiac output

 $= ([Hb] \times SpO2 \times 1.34) \times (HR \times SV)$

Oxygen content = $[Hb] \times SpO2 \times 1.34$

Cardiac output = HR x SV

Fluid Challenge

What is the difference between an infusion and a challenge?

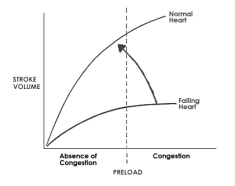
Fluid Challenge

What is the difference between an infusion and a challenge?

250 to 500 ml colloid (or blood products)
500 to 1000ml Hartmann's
[NOT 5% dextrose]
As fast a possible (with pressure bag)
You at the bedside

Fluid Challenge

Aim is to improve SV (and hence CO) by increasing preload Frank-Starling mechanism



Markers of perfusion

What are they?

Markers of perfusion

CVP

What are they?

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

What does it mean?

CVP

Lactate

What does it mean?

Starling's Law

Estimate of LVEDV (i.e. preload)

Not always a good correlation with volumeresponsiveness

However if low strongly suggestive of hypovolaemia

What does it mean?

Lactate ScvO2

What does it mean?

- 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

ScvO2

What does it mean?

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

• Target > 70%

What does it mean?

ScvO2

What can I do if it's low?

ScvO2 ScvO2

What can I do if it's low?

Delivery = [Hb] \times SpO2 \times 1.34 \times HR \times SV

Surviving Sepsis Campaign: International guidelines for management of severe sepsis and septic shock: 2012

R. Phillip Dellinger, Mitchell M. Levy, Andrew Rhodes, Djillali Annane, Herwig Gerlach, Steven M. Opal, Jonathan E. Sevransky, Charles L. Sprung, Ivor S. Douglas, Roman Jaeschke, Tiffany M. Osborn, Mark E. Nunnally, Sean R. Townsend, Konrad Reinhart, Ruth M. Kleinpell, Derek C. Angus, Clifford S. Deutschman, Flavia R. Machado, Gordon D. Rubenfeld, Steven A. Webb, Richard J. Beale, Jean-Louis Vincent, Rui Moreno, and the Surviving Sepsis Campaign Guidelines Committee including the Pediatric Subgroup.

Crit Care Med. 2013; 41:580-637

Intensive Care Medicine 2013; ..

What can I do if it's low?

Delivery = $[Hb] \times SpO2 \times 1.34 \times HR \times SV$

Fluid optimise

Transfuse packet cells

HCt > 30%

Inotropes

Current Surviving Sepsis Campaign Guideline Sponsors

- American Association of Critical-Care Nurses
- American College of Chest Physicians
- American College of Emergency Physicians
- Australian and New Zealand Intensive Care Society
- Asia Pacific Association of Critical Care Medicine
- · American Thoracic Society
- Brazilian Society of Critical Care(AIMB)
- Canadian Critical Care Society
- · Chinese Society of Critical Care Medicine
- **Emirates Intensive Care Society**
- European Respiratory Society
- European Society of Clinical Microbiology and Infectious Diseases
- European Society of Intensive Care Medicine
- European Society of Pediatric and Neonatal Intensive Care
- Infectious Diseases Society of America

- · Indian Society of Critical Care Medicine
- International Pan Arab Critical Care Medicine Society
- Japanese Association for Acute Medicine
- Japanese Society of Intensive Care Medicine
- Pediatric Acute Lung Injury and Sepsis Investigators
- Society Academic Emergency Medicine
- Society of Critical Care Medicine
- · Society of Hospital Medicine
- · Surgical Infection Society
- · World Federation of Critical Care Nurses
- World Federation of Pediatric Intensive and Critical Care Societies
- World Federation of Societies of Intensive and Critical Care Medicine

Participation and endorsement:

German Sepsis Society

Latin American Sepsis Institute

"Time Zero"

- Time Zero = time of presentation
 - -ED, Medical Floors, ICU
- Both bundles time based
- Most important time based elements:
 - Antibiotic timing
 - Resuscitation timing (EGDT)

Hospital Mortality by Time to Antibiotics

Time to ABX ¹ , hrs	OR ²	959	% CI	p-value	Probability of mortality ³	959	% 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

¹Time to ABX is based on 15,948 observations that are greater than or equal to zero

Antibiotic therapy

1. We recommend that intravenous antimicrobial therapy be started as early as possible and within the first hour of recognition of septic shock (1B) and severe sepsis without septic shock (grade1C).

Fluid therapy

4. We recommend that initial fluid challenge in patients with sepsis-induced tissue hypoperfusion with suspicion of hypovolemnic be started with ≥ 1000 mL of crystalloids (to achieve a minimum of 30ml/kg of crystalloids in the first 4 to 6 hours).
(Grade 1B).

²Hospital mortality odds ratio referent group is 0 hours for the time to ABX and is adjusted by the number of baseline organ failures, infection type (community vs. nosocomial), and geographic region (Europe, North America, and South America)

Logistic Regression Model

Compliance indicator	Hospital mortality odds ratio ¹	95% CI	<i>p</i> -value
Serum lactate within 6 hours	0.71	0.67 - 0.75	< 0.001
2. Blood cultures before antibiotics	0.81	0.76 - 0.86	< 0.001
3. Broad spectrum antibiotics	0.83	0.79 - 0.88	< 0.001
4. Fluids and vasopressors	0.57	0.54 - 0.61	< 0.001
CVP ≥ 8 mm Hg within 6 hours	0.74	0.69 - 0.79	< 0.001
6. ScvO ₂ ≥ 70% within 6 hours	0.73	0.67 - 0.78	< 0.001
7. Resuscitation bundle	0.77	0.72 - 0.83	< 0.001
8. Low-dose steroids policy	0.82	0.770.88	< 0.001
9. Drotrecogin alfa policy	0.93	0.88 – 0.98	0.008
10. Glucose control maintained	0.70	0.69 - 0.74	< 0.001
11. IPP < 30 cm H ₂ 0	0.78	0.71 - 0.86	< 0.001
12. Management bundle	0.72	0.68 - 0.77	< 0.001
High resuscitation performance	0.79	0.75 - 0.83	< 0.001
High management performance	0.84	0.80 - 0.88	< 0.001

SSC Bundle:

TO BE COMPLETED WITHIN 6 HOURS OF TIME OF PRESENTATION:

- Apply vasopressors (for hypotension that does not respond to initial fluid resuscitation to maintain a mean arterial pressure (MAP) ≥65mmHg)
- 6. In the event of persistent arterial hypotension despite volume resuscitation (septic shock) or initial lactate ≥4 mmol/L (36mg/dl):
 - Measure central venous pressure (CVP)*
 - Measure central venous oxygen saturation (ScvO2)*
- 7. Remeasure lactate*
- * Targets for quantitative resuscitation included in the guidelines are CVP of ≥8 mm Hg, ScvO2 of ≥70% and lactate normalization.

SSC Bundle:

TO BE COMPLETED WITHIN 3 HOURS OF TIME OF PRESENTATION :

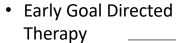
- Measure lactate level
- 2. Obtain blood cultures prior to administration of antibiotics
- 3. Administer broad spectrum antibiotics
- 4. Administer 30ml/kg crystalloid for hypotension or lactate ≥4mmol/L
- "time of presentation" is defined as the time of triage in the Emergency Department or, if presenting from another care venue, from the earliest chart annotation consistent with all elements severe sepsis or septic shock ascertained through chart review.

The Importance of Early Detection

- Efforts to just treat recognized sepsis alone are incomplete
- A critical aspect of mortality reduction in the Campaign has been pushing practitioners to identify sepsis early.
 - Levy MM, Dellinger RP, Townsend SR, et al. The Surviving Sepsis Campaign: Results Of An International Guideline-Based Performance Improvement Program Targeting Severe Sepsis. Crit Care Med. 2010 Feb;38(2):367-74.
- It may well be that earlier recognition accounts for much of the signal in mortality reduction and partially explains sharply increasing incidence.
 - Gaieski DF, Edwards JM, Kallan MJ, et al. Benchmarking the Incidence and Mortality of Severe Sepsis in the United States. Crit Care Med. 2013 Feb 25. [Epub ahead of print]
- Without recognition that the clock is ticking, there is simply no incentive to recognize a challenging diagnosis early.

6 Hour Resuscitation Bundle

- Early Identification
- Early Antibiotics and Cultures





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VOLUME 345	NOVEMBER 8, 2001	NUMBER 19
	The New England Journal of Medicine	

EARLY GOAL-DIRECTED THERAPY IN THE TREATMENT OF SEVERE SEPSIS AND SEPTIC SHOCK

EMANUEL RIVERS, M.D., M.P.H., BRYANT NGUVEN, M.D., SUZAINE HAVSTAD, M.A., JULIE RESSLER, B.S., ALEXANDRIA MUZZIN, B.S., BERNHARD KROBLICH, M.D., EDWARD PETERSON, PH.D., AND MICHAEL TOMLANOVICH, M.D., FOR THE EARLY GOAL-D

Thank You

Rhode Island Hospital EGDT Data

