

Acute Respiratory Failure

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Acute Respiratory Failure

Physiologic Classification

	Type 1 Hypoxemic	Type 2 Hypercarbic	Type 3 Post-op	Type 4 Shock
Mechanism	Shunt	$\downarrow V_a$	Atelectasis	\downarrow Cardiac Output
Etiology	Airspace Flooding	Increased Respiratory load, Decreased ventilatory drive	Decreased FRC and increased Closing Volume	Decreased FRC and increased Closing Volume
Clinical Setting	Water, Blood or Pus filling alveoli	CNS depression, Bronchospasm, Stiff respiratory system, respiratory muscle failure	Abdominal surgery, poor insp effort, obesity	Sepsis, MI, acute hemorrhage

Respiratory Failure

Physiologic Definition:

Inability of the lungs to meet the metabolic demands of the body

Can't take in enough O_2
or
Can't eliminate CO_2 fast enough to keep up with production

Ventilatory Failure



Inbalance between load on the lungs and the ability of bellows to compensate

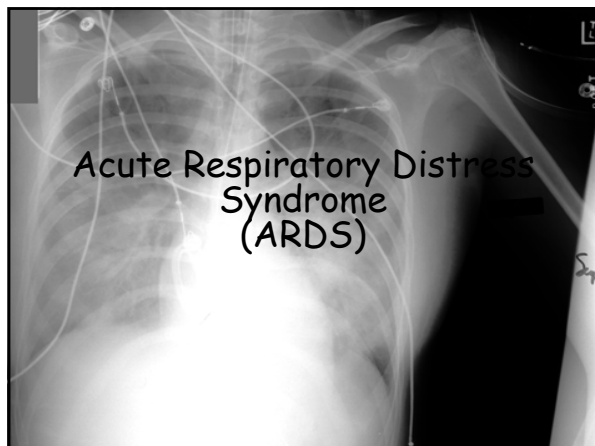
Respiratory Failure

- Failure of Oxygenation: $P_aO_2 < 60$ mmHg
- Failure of Ventilation*: $P_aCO_2 > 50$ mmHg

* P_aCO_2 is directly proportional to alveolar minute ventilation

Acute Hypoxemic Respiratory Failure

- Shunt disease - intracardiac or intrapulmonary
- Severe V/Q mismatch - asthma, PE
- Venous admixture due to low cardiac output states, severe anemia coupled with shunt and/or V/Q mismatch



Acute Respiratory Distress Syndrome

Each year in the United States

75,000-150,000 cases

Acute Respiratory Distress Syndrome (ARDS)

Leaky alveolar capillaries

Plasma fluid and leukocytes leak into the airspace

Shunt

Hypoxemia

Causes of ARDS

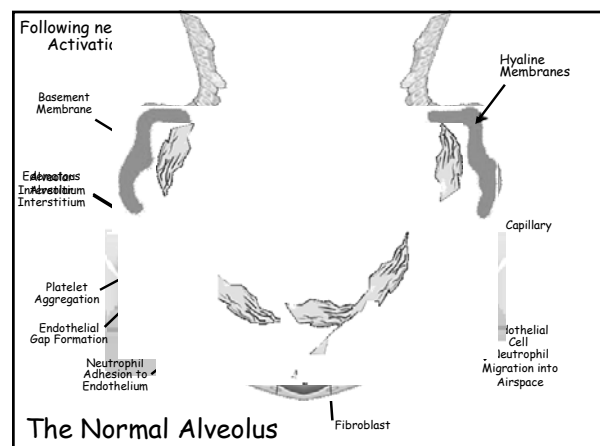
DIRECT LUNG INJURY	INDIRECT LUNG INJURY
Pneumonia	Non-pulmonary sepsis/SIRS
Aspiration of gastric contents	Severe trauma with shock
Pulmonary contusion	Cardiopulmonary bypass
Near-drowning	Drug overdose (Narcotics)
Inhalation injury (Cl ⁻ , smoke)	Acute pancreatitis
Reperfusion pulmonary edema after lung transplantation or pulmonary embolectomy	Transfusion (TRALI)
	Drug reaction (ARA-C, nitrofurantoin)
	fat/air/amniotic fluid embolism, bypass

Acute Respiratory Distress Syndrome (ARDS)

American-European Consensus Definition:

- Refractory hypoxemia
 $P_{a}O_2/F_{i}O_2$ (P/F ratio)
 <300 for ALI
 <200 for ARDS
- A disease process likely to be associated with ARDS
- No evidence of elevated left atrial pressure elevation (by clinical exam, echo or PA catheter)
- Bilateral airspace filling disease on X-ray

Report of the American-European Consensus conference on acute respiratory distress syndrome: definitions, mechanisms, relevant outcomes, and clinical trial coordination. Consensus Committee.

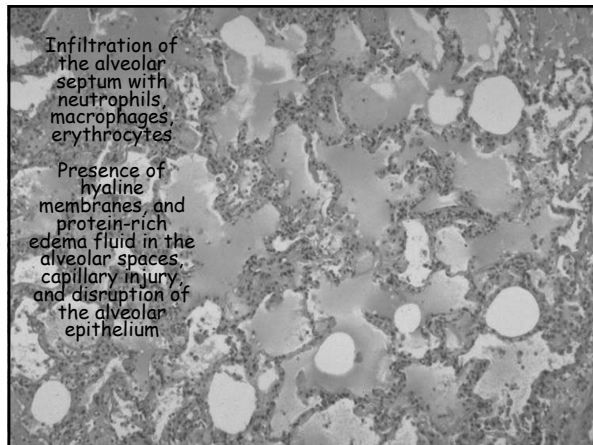
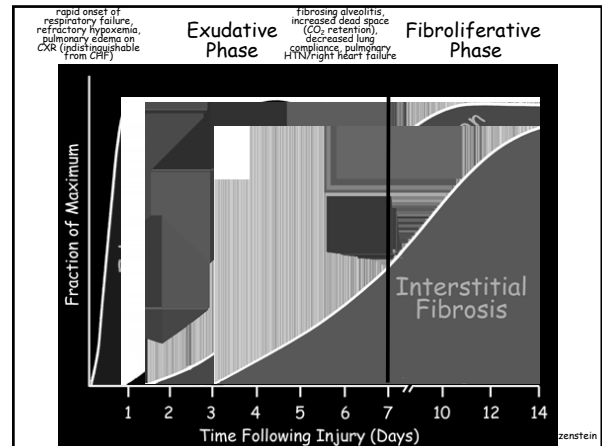


ARDS

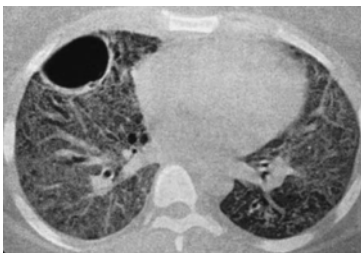
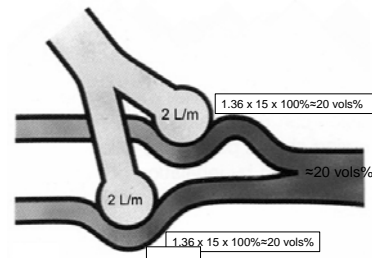
Fundamental Pathophysiology:

Increased alveolar permeability due to direct neutrophil-mediated injury to the alveolar epithelium

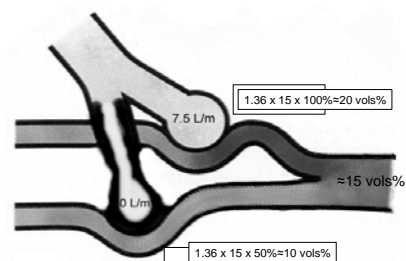
Not a distinct disease - rather a sequelae of activation of lung and systemic inflammatory pathways



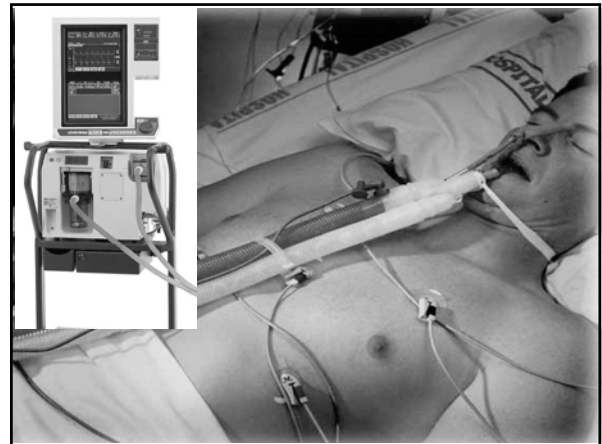
Optimal V/Q matching



Shunt



Severe Hypoxemia

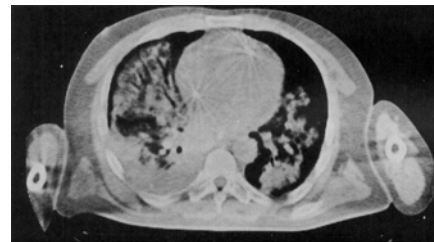


Therapeutic Goals

Maintain reasonable oxygen delivery

Find & fix the primary cause

"Baby Lungs"



FRC can be reduced by 80% or more in ARDS

Gattinoni, et. al. *Anesthesiology*, 74:15-23, 1991.



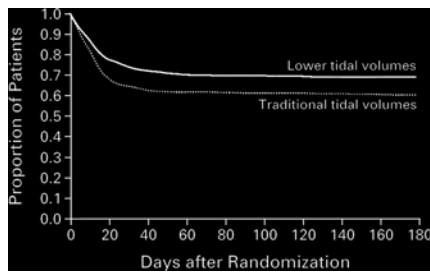
ARDS Network Trial

Day 1 Ventilatory Characteristics

	Low V_t Group n=432	Traditional V_t Group n=429
V_t :	6.2 ± 0.9	11.8 ± 0.8
PEEP:	9.4 ± 3.6	8.6 ± 3.6
F_{iO_2} :	0.56 ± 0.19	0.51 ± 0.17
P_{plat} :	25.7 ± 7	33 ± 9
P_{peak} :	32.8 ± 8	39 ± 10
P_aO_2 / F_{iO_2} :	158 ± 73	176 ± 76
P_aCO_2 :	40 ± 10	35 ± 8
pH:	7.38 ± 0.08	7.41 ± 0.07

NEJM 342:1301-1308, 2000

ARDS Network Trial



Mortality: 39.8 % in traditional tidal volume group, 31% in low tidal volume group ($P=0.007$)

Also: @ 28 days: more ventilator free days (12 vs. 10), more days without organ failure (15 vs 12), higher rate of liberation from ventilation rate (63.7% vs 55%)

NEJM 342:1301-1308, 2000

What happens to alveoli in ARDS?

Edema accumulates in alveoli

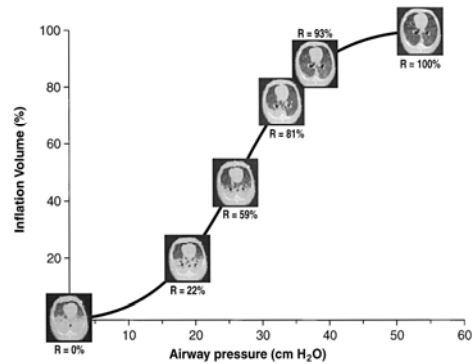
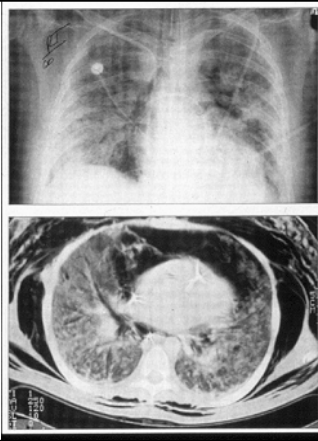
Diluting & disaggregating surfactant

Surface tension increases

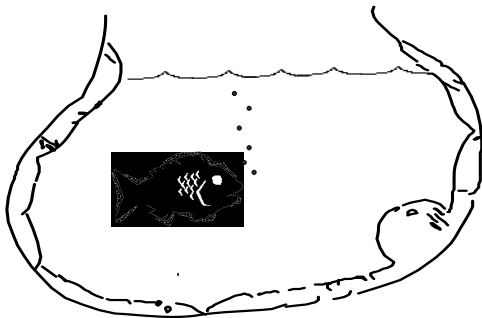
Alveoli collapse



Alveolar collapse decreases FRC and contributes to hypoxemia



What happens to alveoli in ARDS?



Positive End-Expiratory Pressure (PEEP)

- Beneficial Effects
 - Increases FRC, CI, P_{aO_2}
 - Recruits Atelectatic Units
 - Decreases Q_s/Q_t
 - Allows Reduction in F_{iO_2}
- Detrimental Effects
 - Volutrauma
 - Alveolar Overdistention
 - Hemodynamic Derangements

PEEP

Oxygen is:

- A) good for you
- B) bad for you
- C) all of the above

$F_{I}O_2 > 0.6$ for 24 hours or more may cause lung injury

PEEP recruits collapsed alveoli,
improves FRC and
improves oxygenation

An essential therapy for patients with ARDS

Does Mechanical Ventilation Contribute to MSOF?

Ranieri, et al.*: randomized prospective study of the effects of mechanical ventilation on bronchoalveolar lavage fluid and plasma cytokines in patients with ARDS (primarily non-pulmonary causes).

Controls (n=19): Rate 10-15 bpm, V_T targeted to maintain $PaCO_2$ 35-40 mmHg (mean: 11 ml/kg), PEEP titrated to SpO_2 (mean: 6.5), P_{plat} maintained <35 cmH₂O

Lung protective ventilation (n=18): Rate 10-15 bpm, V_T targeted to keep P_{plat} less than upper inflexion point (mean: 7 ml/kg), PEEP 2-3 cmH₂O above LIP (mean: 14.8)

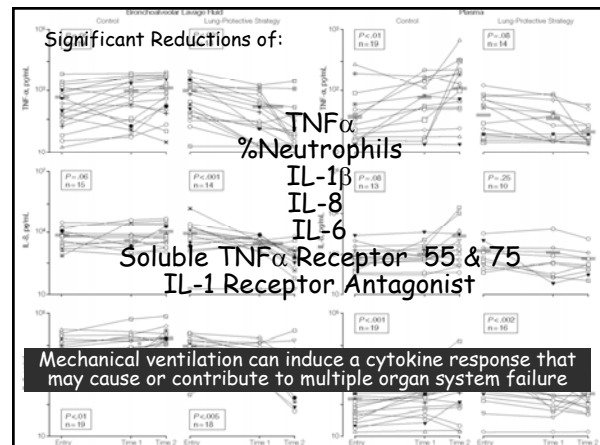
Plasma and BALF levels of IL-1 β , IL-6, IL-8, TNF α , TNF α -sr 55, TNF α -sr 75, IL-1ra, measured within 8 hrs of intubation and again @24-30 hours & 36-40 hours after entry

*Ranieri, et al. Effect of mechanical ventilation on inflammatory mediators in patients with acute respiratory distress syndrome: a randomized controlled trial. JAMA 282:54-61, 1999.

ARDS Network Trial

The standard of care

Assist Control
 V_T 6 cc/kg ideal body weight
PEEP of \approx 8-10



Cause of Death in ARDS Patients?

Generally not due to
respiratory failure

The lung is not just an innocent bystander - it functions as an immunomodulatory organ that may participate in the systemic inflammatory response that leads to multiple organ system dysfunction syndrome

Biotrauma

Goals for Management of ARDS

The American-European Consensus
Conference on ARDS, Part 2

- Ensure appropriate O_2 delivery to vital organs
- Minimize oxygen toxicity/tolerate mediocre ABG's
- Reduce edema accumulation
- Minimize airway pressures
- Prevent atelectasis/Recruit alveoli
- Use sedation and paralysis judiciously

Am J Resp Crit Care Med 157:1332-47, 1998.

Survival from "pure" ARDS

1979: 20-50%

2002: 50-90%