# ACNP Pulmonary Respiratory Failure Cheat Sheet by xkissmekatex (kissmekate) via cheatography.com/33594/cs/10535/

Hypoxemia	a
Workup	Get PaCO2 level, A-a gradient, and response to supplemental O2.
lf PaCO2 is elevated	<ul> <li>Hypoventilation is occurring.</li> <li>If the A-a gradient is normal, hypoventilation or low inspired PO2 is the only mechanism.</li> <li>If the A-a gradient is increased, hypoventilation is not the lone mechanism of hypoxemia.</li> </ul>
If PaCO2 is not elevated	<ul> <li>No hypoventilation.</li> <li>If A-a gradient is not increased, low inspired PaO2 is the cause of hypoxemia.</li> </ul>
If A-a gradient is increased	• With low to normal PaCO2. See if the lowered PaO2 is improved with supplemental O2. If so, there is V/Q mismatch. If it does not improve, shunting is present.

Hypoxemia	a (cont)
V/Q Mismatch	<ul> <li>Typically leads to hypoxia</li> <li>without hypercapnia (PaCO2 is usually low to normal).</li> <li>Most common mechanism of hypoxemia (esp. in chronic lung disease).</li> <li>Responsive to supplemental oxygen.</li> <li>Ventilation without perfusion.</li> </ul>
Intrapulm onary Shunting	<ul> <li>Little or no ventilation in perfused areas due to collapsed or fluid-filled alveoli.</li> <li>Venous blood is shunted into the arterial circulation without being oxygenated.</li> <li>Perfusion without ventilation.</li> <li>Causes include atelectasis or fluid buildup in alveoli (pneumonia or edema) or direct right to left intracardiac blood flow in congenital heart disease.</li> <li>No responsive to supplemental oxygen.</li> </ul>
Hypoventi ation	Leads to hypercapnia with secondary hypoxemia.

#### Hypoxemia (cont)

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Increased CO2	In sepsis, DKA, hyperthermia, etc. Leads to <b>hypercapnic</b>
production	respiratory failure.
Diffusion Impairment	In ILD causes hypoxemia without hypercapnia.

### Hypoxemic Failure

Нурохі	a PaO2<60mmHg and
	PaCO2>50mmHg.
	<ul> <li>Severe hypoxemia can result in</li> </ul>
	irreversible damage to CNS and
	CVS, and must be corrected rapidly.
	O2 saturation is <90% despite
	FiO2>0.6.
	<ul> <li>Causes include lung pathology like</li> </ul>
	ARDS, severe pneumonia, and
	pulmonary edema.
	<ul> <li>V/Q mismatch and</li> </ul>
	intrapulmonary shunting are the
	major pathophysiologic
	mechanisms.
Oxyge	n Delivery
Low	Nasal cannula has flow rate of 1-
Flow	6L/min, FiO2 up to 0.40, is easy to use
	and comfortable.
	• Simple face masks have a flow rate of
	1-101 /min_EiO2 of 0.40-0.60 and can

deliver higher flow rates than nasal cannula.

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Oxygen Delivery (cont)			Acute Respiratory Failure (cont)		
High • V Flow 10 me CC an • N rat 0.8	Venturi masks have a flow rate of 4- L/min, FiO2 up to 0.50 with precise easurements, and are preferred in D2 retainers due to higher precision d control of oxygenation. Nonrebreather masks have flow res up to 15L/min, FiO2 up to 0.70- 30, and can achieve higher FiO2 at		Signs	Inability to speak in complete sentences, use of accessor muscles tachypnea, tachycardia, cyanosis, impair mentation (due to fatigue or hypercapnia, of if cause of respiratory failure is CNS depression).	
NPPV • E ful im int • S <b>res</b> Pa aw pro	Ver flow rates. SIPAP or CPAP via nasal mask or I-face mask. Indicated in patients in pending respiratory failure to avoid ubation or mechanical ventilation. Success is highest in <b>hypercarbic</b> <b>spiratory failure (esp. COPD)</b> . tient must be neurologically intact, rake, cooperative, and able to patient their airway.		Causes	<ul> <li>CNS Causes: depression o insult from drug overdose, str or trauma.</li> <li>Neuromuscular Causes: myasthenia gravis, polio, Guillain-Barre syndrome, amyotrophic lateral sclerosis.</li> <li>Upper Airway Causes: obstruction due to stenoses, spasms, or paralysis</li> </ul>	
Acute Res	piratory Failure			Thorax and Pleural Causes mechanical restriction due to	
Pathophysi ology Symptoms	Pathophysi Results when there is plogy inadequate oxygenate of blood or inadequate ventilation or both. Symptoms dyspnea is the first. Cough may or may not be present, depending		kyphosco hemothro • CVS and valvular d anemia. • Lower A COPD, pi	kyphoscoliosis, flail chest or hemothroax. • CVS and Heme Causes: CH valvular diseases, PE, and anemia. • Lower Airway Causes: asth COPD, pneumonia, ARDS.	
	on the underlying cause.		Diagnosis	ABG, CXR, CT, CBC, BMP, a consider cardiac enzymes i pulmonary edema is suspect	

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### Hypercarbic Failure

ele	Hyperc	• PCO2>50mmHg.
y	apnia	Severe hypercapnia can lead to
		dyspnea and vasodilation of cerebral
red		vessels (with increased ICP and
		subsequent papilledema, HA,
		impaired consciousness, and finally
		coma).
		A failure of alveolar ventilation.
r		• Either a decrease in minute
roke.		ventilation or an increase in
, one,		physiologic dead space that leads
		to CO2 retention and eventually
		hypoxemia.
		May be caused by underlying lung
		disease (COPD, asthma, CF, severe
		bronchitis) or without lung disease
		(patients with impaired ventilation
		due to neuromuscular diseases, CNS
:		depression, mechanical restriction of
1		the chest, or respiratory fatigue). Also
		increased CO2 production or
		diffusion impairment.
HF,		<ul> <li>Hypercapnia can be caused by</li> </ul>
		hypoventilation (secondary to a
		variety of causes).
ima,		<ul> <li>Respiratory acidosis occurs when</li> </ul>
		hypercapnia is present.
and		<ul> <li>If chronic acidosis, renal</li> </ul>
if		compensation occurs and acidosis is
ed.		less severe.

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#### Treatment

### · Treat underlying disorder with bronchodilators, corticosteroids, antibiotics, etc.

• Provide supplemental oxygen if hypoxemic. In hypoxemic failure, use lowest concentration of oxygen that provides sufficient oxygenation to avoid oxygen toxicity (due to free radicals).

• In hypercarbic failure, traditionally do not give high concentrations of O2, especially in patients with COPD to not depress respiratory drive. Apply NPPV only for conscious patients. Intubation and mechanical ventilation

#### Acute Respiratory Distress Syndrome

Pathop

gy

 A diffuse inflammatory process hysiolo- involving both lungs. Neutrophils are activated in the systemic or pulmonary circulations. • Not a primary disease but a disorder that arises from other conditions that cause a widespread inflammatory process.

> There is massive intrapulmonary shunting, decreased pulmonary compliance (increased work), increased dead space (secondary to obstruction and destruction of pulmonary capillary bed), low VC, and low FRC

Acute F	Respiratory	/ Distress	Syndrome	(cont)
Addici	copilator	Disticus	oyna one	(Cont)

Intrapulm	<ul> <li>Severe hypoxemia with no</li> </ul>			
onary	significant improvement on			
Shunting	100% oxygen.			
	<ul> <li>Shunting secondary to</li> </ul>			
	widespread atelectasis, collapse			
	of alveoli, and surfactant			
	dysfunction.			
	<ul> <li>Interstitial edema and alveolar</li> </ul>			
	collapse are due to increased lung			
	fluid that leads to stiff lungs, an			
	increase in A-a gradient, and			
	ineffective gas exchange.			
	<ul> <li>Effects of the increase in</li> </ul>			
	pulmonary fluid are identical to			
	those in cardiogenic pulmonary			
	edema, but the cause is			
	increased alveolar capillary			
	permeability			

#### Acute Respiratory Distress Syndrome (cont)

Causes	• Sepsis is the most common (which can itself be secondary to pneumonia, urosepsis, wound infections, etc). Aspiration of gastric contents, severe trauma, fractures (femur, pelvis), <b>acute</b> <b>pancreatitis</b> , multiple or massive transfusions, near-drowning, drug overdose, toxic inhalation, intracranial HTN, and cardiopulmonary bypass.
Signs	<ul> <li>Dyspnea, tachypnea, tachycardia due to increased work of breathing.</li> <li>Progressive hypoxemia not responsive to supplemental oxygen (ratio of PaO2 to FiO2&lt;200).</li> <li>Patients are difficult to ventilate because of high peak airway pressures due to stiff, noncompliant lungs.</li> </ul>
Diagnosis	• Hypoxemia that is <b>refractory to</b> <b>oxygen therapy</b> (ratio of PaO2/FiO2<200-300), bilateral diffuse pulmonary infiltrates on CXR, no evidence of CHF (PCWP <18mmHg).
CXR	Shows diffuse bilateral pulmonary infiltrates. Improvement on CXR follows clinical improvement after 1-2 weeks

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Acute Respiratory Distress Syndrome (cont)			Acute Respiratory Distress Syndrome (cont)		
Arterial Blood Gas	<ul> <li>Hypoxemia (PaO2&lt;60mmHg).</li> <li>Initially, respiratory alkalosis</li> <li>(PaCo2&lt;40) is present, which</li> <li>gives way to respiratory acidosis</li> <li>as the work of breathing increases</li> <li>and PaCO2 increases.</li> <li>If patient is septic, metabolic</li> <li>acidosis may be present with or</li> <li>without respiratory</li> <li>compensation.</li> </ul>	Treatment		<ul> <li>Oxygenation &gt;90%.</li> <li>Mechanical ventilation with PEEP is usually required (increases lung volume by opening collapsed alveoli and decreasing shunting).</li> <li>Avoid volume overload and maintain low-normal intravascular volume (PCWP 12-15mmHg).</li> <li>Vasopressors may be needed to</li> </ul>	
Pulmonary Artery Catheter	<ul> <li>Enables a determination of PCWP, which reflects left heart filling pressures and is an indirect marker of intravascular volume status.</li> <li>The most useful parameter in differentiating ARDS from cardiogenic pulmonary edema.</li> <li>If PCWP is &lt;18mmHg, ARDS is more likely, whereas if PCWP is &gt;18mmHg, cardiogenic is more likely.</li> </ul>	Į	·	maintain BP. • Patients with sepsis have high fluid requirements, so this might be difficult. Treat underlying cause. Tube feedings are preferred over parenteral nutrition.	
Bronchosc opy	With bronchoalveolar lavage. May be considered if patient is acutely ill and infection is considered. Fluid collected can be cultured and analyzed for cell differential, cytology, Gram stain, and silver stain.	-			
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