

Health Care Associated Pneumonia

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Objectives

- Define the terms, pneumonia, community acquired pneumonia, health care associated pneumonia
- (HCAP) and ventilator associated pneumonia (VAP).
- Describe the pathogenesis of the health care associated pneumonia (hospital associated pneumonia) and VAP.
- Classify HCAP according to the time of onset.

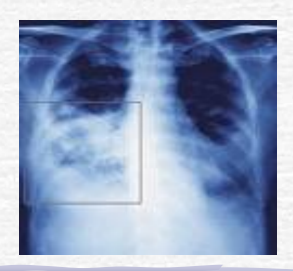
Objectives

- Name the different causative bacterial agents.
- Classify and describe types of VAP.
- Recognize the ways by which VAP is prevented.
- Describe the different chemotherapeutic anti microbial agents used for the treatment of health care associated pneumonia.
- Evaluate response to treatment and recognize reasons for failure of treatment.

Health Care Associated Pneumonia

Definition of Pneumonia:Infection of the pulmonary Parenchyma





PNEUMONIA can be:

- **A-Community acquired Pneumonia** acquired in the community, by community acquired organism, eg. *Streptococcus pneumoniae* usually susceptible to antibiotic.
- **B-Health care associated pneumonia** acquired **48-72 hours** after admission to health care institutions eg. pneumonia caused by organisms in hospital which are usually resistant to antibiotics-eg. *Pseudomonas aeruginosa*

Definition

- Nosocomial pneumonia: is defined as hospital associated pneumonia (HAP) or health care associated pneumonia (HCAP).
- Occurring at least 48 hours after admission and not incubating at the time of hospitalization.

Health care associated Pneumonia

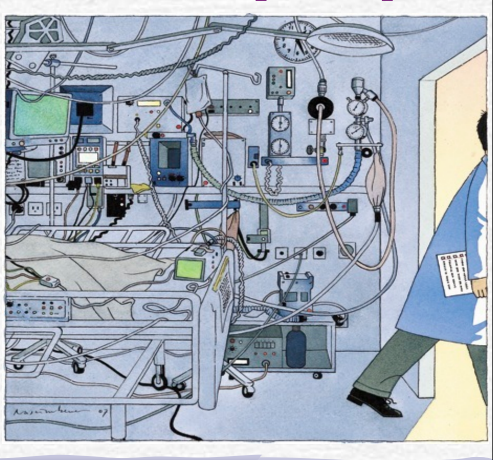
- A- Hospital Acquired Pneumonia(HAP)
- **B-** Ventilator Associated Pneumonia (**VAP**) in patients with assisted respiration for a period of 48 hours.

Introduction

- Nosocomial pneumonia is the 2nd most common hospital-acquired infections after urinary tract infection. Accounting for 31 % of all nosocomial infections
- Nosocomial pneumonia is the leading cause of death from hospital-acquired infections.
- The incidence of nosocomial pneumonia is highest in **ICU** (intensive care unit) patients.

Intensive Care Unit (ICU)





Introduction

- The incidence of nosocomial pneumonia in ventilated patients is 10-fold higher than non-ventilated patients
- The reported crude **mortality** for HAP is 30% to greater than 70%.

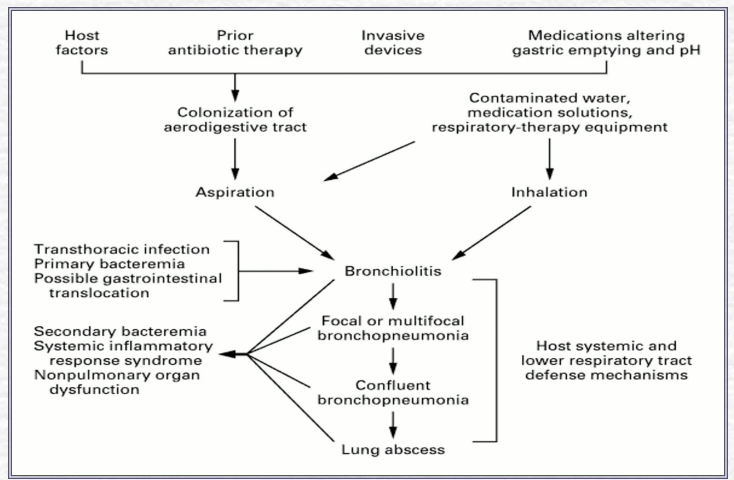
Pathogenesis of HAP



Pathogenesis

- For pneumonia to occur, at least one of the following **three conditions** must occur:
 - 1. Significant impairment of host defenses
 - 2. Introduction of a sufficient-size inoculum to overwhelm the host's lower respiratory tract defenses
 - 3. The introduction of highly virulent organisms into the lower respiratory tract
- Most common is microaspiration of oropharyngeal secretions colonized with pathogenic bacteria.

Pathogenesis

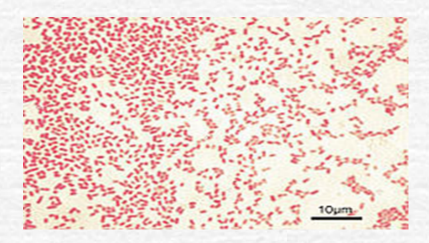


Classification

- Early-onset nosocomial pneumonia:
 - Occurs during the first 4 days of admission.
 - Usually is due to *S. pneumoniae*, MSSA (Methicillin sensitive *S.aureus*), *H. Influenza*.
- Late-onset nosocomial pneumonia:
 - OCCURS more than 4 days of admission.
 - More commonly by Gram negative organisms, especially: *P. aeruginosa, Acinetobacter, Enterobacteriaceae* (Klebsiella, Enterobacter, Serratia) or MRSA.

- Enteric Gram negative bacilli are isolated most frequently particularly in patients with late-onset disease and in patients with serious underlying disease often already on broad-spectrum antibiotics.
- Prior use of broad-spectrum antibiotics and an immunocompromised state make resistant Gram-negative organisms more likely.

• P. aeruginosa and Acinetobacter are common causes of late-onset pneumonia, particularly in ventilated patients.



- S. aureus is isolated in about 20~40% of cases and is particularly common in :
 - 1. Ventilated patients after head trauma, neurosurgery, and wound infection
 - 2. In patients who had received prior antibiotics or Prolonged care in ICU
- MRSA(methicillin resistant S.aureus) is seen more commonly in patients who:
 - Received corticosteroids
 Undergone mechanical ventilation >5 days
 Presented with chronic lung disease
 Had prior antibiotics therapy

- Anaerobes are common in patients predisposed to aspiration.
- Ventilator associated pneumonia (VAP) with anaerobes occurred more often with oropharyngeal intubation than nasopharyngeal intubation.

Ventilator-associated Pneumonia (VAP)





Ventilator-associated Pneumonia (VAP)

Definition:

Nosocomial pneumonia that has developed in patient who are receiving mechanical ventilation.

Classification:

Early-onset: within 48-72 hours after tracheal intubation, which complicates the intubation process

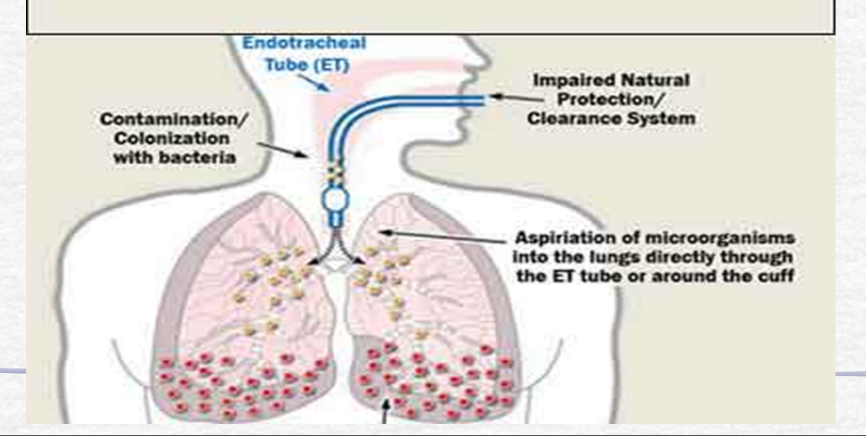
Late-onset: after 72 hours

Pathogenesis

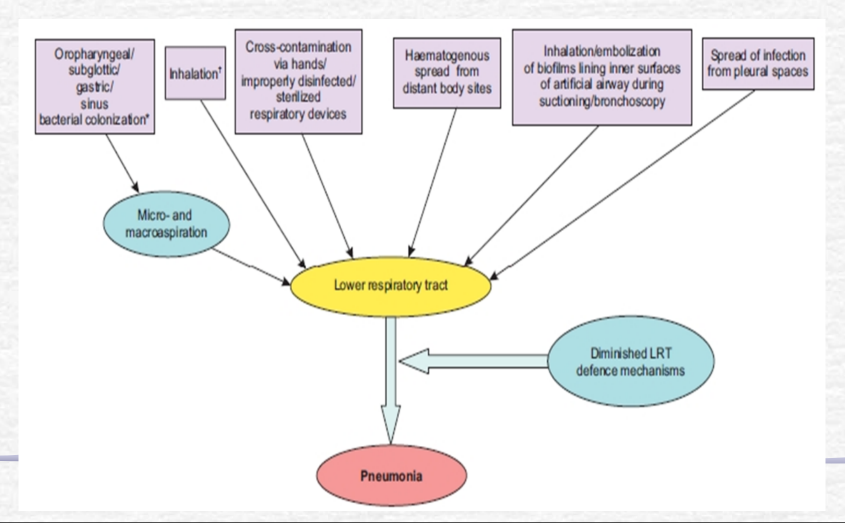
- Requires 2 important processes:
 - 1. Bacterial colonization of the aerodigestive tract
 - 2. Aspiration of contaminated secretion into the Lower airway
- Mechanical ventilation prevents mechanical clearance by cough and the mucociliary escalator.
- Sources of infection: endogenous or exogenous.

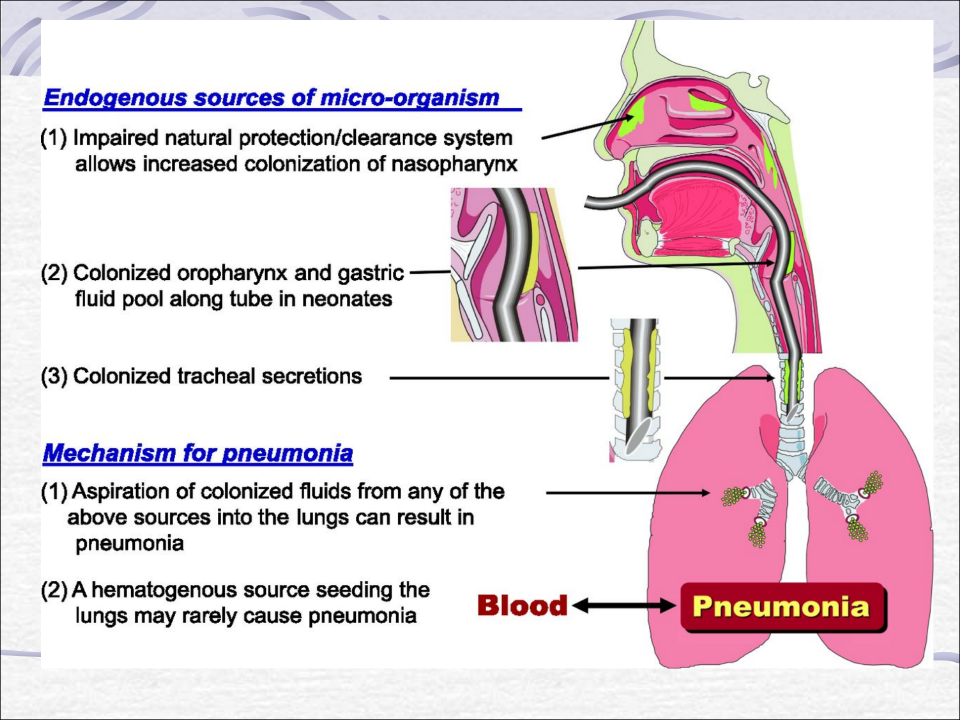
Pathogenesis of VAP

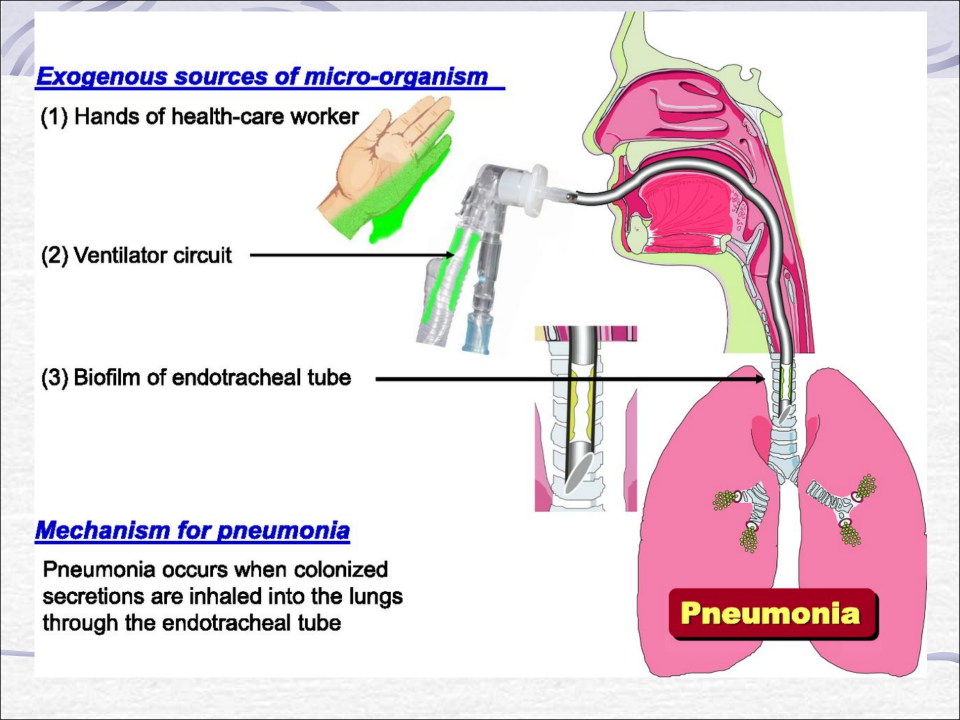
Ventilator Associated Pneumonia



Pathogenesis of VAP







Prevention for VAP

The oral regimen (topical Gentamicin, Colistin, Vancomycin cream given every 6h for 3 weeks) treating oropharyngeal colonization could prevent VAP.

--- Prevention of VAP by oral decontamination

American journal of respiratory critical care medicine 2001 164:382-8

Preventions for VAP

Non-pharmacologic strategies

- Effective hand washing and use of protective gowns and gloves
- Semirecumbent positioning
- Avoidance of large gastric volume
- Oral (non-nasal) intubation
- Continuous subglottic suctioning
- Humidification with heat and moisture exchanger
- Posture change

Preventions for VAP

Pharmacologic strategies

- Avoiding stress-ulcer prophylaxis
- Prophylactic antibiotic therapy
- Chlorhexidine oral rinse
- Prophylactic treatment of neutropenic patients
- Vaccines

Most initial therapy is empiric because no pathogen is identified or results are not available when antimicrobial decisions are made in most patients.

- Initially be treated with a broadspectrum antibiotic regimen aimed at covering all likely bacterial pathogens
- This regimen should subsequently be narrowed, according to the result of culture.

- The pathogen may be influenced by coexisting illnesses, prior treatment, and length of hospitalization.
- The frequency of ICU-acquired *P. aeruginosa* carriage or colonization/infection was 23.4% at 7 days and 57.8% at 14 days.

---- Current opinion in infectious disease 2002, 15:387-94, copyright LWW

The mortality can be reduced with early appropriate empiric therapy.

(Form 30 % with appropriate therapy to more than 90 % with inappropriate therapy).

- Cefepime or
- Piperacillin-tazobactam or
- Meropenem or
- Levofloxacin or

If risk for MRSA or more sever disease add vancomycin

Response to Therapy

If no clinical response is noted or deterioration occurs, we need to consider:

1. Infectious causes:

Resistant pathogen

Superinfection

Unusual pathogens

Lung abscess

Extrapulmonary infection

2. Noninfectious events:

Heart: congestive heart failure (CHF)

Lung: fibroproliferative acute respiratory distress syndrome (ARDS), pulmonary emboli, Atelectesis.

Reference book

Ryan, Kenneth J. Sherris Medical Microbiology. Letest edition.

Mc Graw -Hill eduction