

# Infections of Nervous System

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## PATHOGENESIS

- CNS is normally sterile.
- parenchyma, coverings, and blood vessels of nervous system may be invaded by *virtually any pathogenic microorganism*.

### Principal routes of entry:

- hematogenous spread** (bacteria, viruses) via septicemia, septic emboli - most common!
  - ordinarily through *arterial* circulation, but retrograde *venous* spread can occur (e.g. via anastomotic connections between veins of face and cerebral circulation).
  - most common sources: pneumonia, bronchiectases, bacterial endocarditis.
- direct implantation** (bacteria) - invariably traumatic (rarely – iatrogenic\*); associated with congenital malformations (e.g. meningomyelocele).
 

\*esp. LP, ventriculo-peritoneal shunts
- local extension** (bacteria) from established infection – *paranasal sinus* (most often frontal), *middle ear, tooth, surgical site* in cranium or spine (osteomyelitis → bone erosion → propagation into CNS).
- retrograde transport through PNS** (certain viruses - rabies, herpes simplex, poliovirus).

### Infection becomes rapidly disseminated once organisms reach CSF.

- *CSF is area of impaired host defense* - lack of sufficient numbers of complement components and immunoglobulins for opsonization, contains no phagocytic cells; fluid medium impairs phagocytosis.

### Damage to nervous tissue:

- 1) direct invasion by infectious agent
- 2) microbial toxins
- 3) destructive inflammatory / immune-mediated response - recently recognized as very important (even in bacterial meningitis).

Inflammatory reaction in confined intracranial space can cause **ICP**↑

## CLASSIFICATION

- according to **major site of involvement**:

N.B. process frequently involves more than one of these structures (e.g. meningoencephalitis, encephalomyelitis)

1. **OSTEOMYELITIS** – inflammation of **bones**.
2. **MENINGITIS** – inflammation of **meninges**.
3. **ENCEPHALITIS** – **viral** invasion of **brain parenchyma**; often *diffuse*.
4. **CEREBRITIS** – *focal bacterial* invasion of **brain parenchyma**; no capsule or pus.
5. **MYELITIS** – inflammation of **spinal cord parenchyma**; no capsule or pus.
6. **ABSCESS** – *focal*, encapsulated, pus-containing cavity in **brain parenchyma** (rarely, in **spinal cord parenchyma**).
7. **EMPHYEMA** – abscess in enclosed or potential **space**:
  - a) subdural
  - b) extradural
8. **GRANULOMA** – *focal*, more or less encapsulated, chronic inflammatory lesion without pus (e.g. sarcoidosis, syphilis, tuberculosis, fungi, larvae of intestinal parasites).

### Infections of spine:

- 1) vertebral osteomyelitis/discitis
- 2) epidural abscess
- 3) subdural abscess\*
- 4) meningitis
- 5) spinal cord abscess\*

\*exceedingly rare.

## VIRUSES

**Neuroinvasive** - virus has ability to enter nervous system.

**Neurotropic** - virus infects nervous cells.

**Neurovirulent** - virus causes clinically recognizable neurologic symptoms.

### ACUTE viral infections:

- a) **viral (aseptic) meningitis**
- b) **encephalitis**
- c) **myelitis**

**DELAYED COMPLICATIONS of acute infection** - **postinfectious polyneuritis, acute disseminated encephalomyelitis (ADEM), acute cerebellar ataxia**.

**LATENT infections with recurrences** from time to time: **herpesviruses** (HSV, VZV).

### SLOWLY PROGRESSIVE disorders (slow viral infections):

- a) **CONVENTIONAL viruses**:
  - 1) **subacute sclerosing panencephalitis (SSPE)** (measles virus)
  - 2) **progressive rubella panencephalitis (PRP)** (rubella virus)
  - 3) **progressive multifocal leukoencephalopathy (PML)** (JC virus)
  - 4) **human T-lymphotrophic virus (HTLV)-associated myelopathy (HAM) / tropical spastic paraparesis (TSP)** (HTLV-I)
  - 5) **acquired immunodeficiency syndrome (AIDS)** (HIV)
- b) **UNCONVENTIONAL transmissible spongiform encephalopathy agents (prions)**.

**FUNGI**

- **opportunistic** organisms – infect only **immunosuppressed** individuals.  
(except few **pathogenic** fungi – *Histoplasma\**, *Blastomyces\**, *Coccidioides\**, *Paracoccidioides\*\** – may infect **normal** hosts).  
\*endemic to some areas of North America  
\*\*endemic to some areas of Central-South America
  - most fungi **invade brain** by **hematogenous dissemination** (but **direct extension** by *Mucor*).
  - lungs / skin / hair are usual **primary sites**.
- Cryptococcosis\* is most common mycotic CNS infection!
- \*may be primary infection and occur in **normal** individuals!
- meningitis
  - intraparenchymal abscess / granuloma
  - vasculitis → thrombosis → infarction (often strikingly hemorrhagic) - *Mucor*, *Aspergillus*

**PREDISPOSING FACTORS**

- Recent infection** that may progress to meningitis (e.g. upper respiratory infection, pneumonia, otitis media leading to pneumococcal meningitis; mumps, chickenpox).
- Exposure to others with infectious illness** (e.g. meningococcus or *Haemophilus influenzae*).
- Recent **travel** (e.g. mosquitoes → arbovirus encephalitis; Central America → cysticercosis).
- Occupation** (e.g. painter exposed to *Cryptococcus* in pigeon droppings)
- Underlying disease:**
  - lymphoma, leukemia, other malignancy
  - renal failure
  - AIDS and other immunodeficiency states
  - alcoholism
  - diabetes
- Drugs** (chemotherapy, immunosuppressant, steroids)
- Recent **head injury** (precedes 10% of pneumococcal meningitis), penetrating skull trauma.
- Recent **neurosurgical procedure**. see p. Op120 >>
- Recent **insect bite** (e.g. Lyme disease, rickettsial infection).
- History of **positive PPD**.

**DIAGNOSIS**

- **CT / MRI** is indicated in any patient with syndrome compatible with CNS infection!
- **CSF** is indicated in any patient (after exclusion of intracranial mass).
- **brain biopsy** (→ immunostaining techniques, electron microscopy, injection into susceptible animals and tissue culture cell lines) is still standard of diagnosis in some specific CNS infections.
- **CBC with differential** is nonspecific adjunct in diagnostic evaluation.  
CBC may be normal in elderly or immunosuppressed patients!
- 2-3 **blood cultures** should be obtained from all patients (even when antimicrobial therapy has already been administered).
- in suspected **any viral CNS infection**, draw **serum specimen** acutely and save to compare with convalescent sera (3-5 weeks after onset of illness) – ≥ 4-fold rise in **IgG titers**?
- **search of infection source** – chest X-ray (!), echocardiography, cultures of other body fluids, bone scans.
- **serum** electrolytes, glucose\*, urea nitrogen, creatinine.  
\*for interpretation of CSF glucose level.

**TREATMENT**

With exception of **viral meningitis**, all but **most chronic** CNS infections require **initial inpatient evaluation and treatment**:

- Bed rest
- Analgesics
- IV antimicrobials
- Fluid balance

**ANTIBIOTICS**

| DRUG IV                       | NEONATES<br>(0-7 days → 8-28 days)  | CHILDREN   | ADULTS  |
|-------------------------------|---|--|---|
| <b>PENICILLIN G</b>           | 100,000-150,000 U/kg/d<br>(divided every 12 hr) →<br>150,000-200,000 U/kg/d<br>(divided every 6-8 hr) | 250,000-400,000<br>U/kg/d (divided every<br>4 hr)  | 20-24 million U/d*<br>(divided every 4 hr)  |
| <b>AMPICILLIN</b>             | 50-75 mg/kg q12h →<br>50-100 mg/kg q6-8h  | 50-100 mg/kg q6h                                   | 2 g q4h   |
| <b>METHICILLIN</b>            |   | 50 mg/kg q6h                                       |   |
| <b>OXACILLIN</b>              | 50-75 mg/kg q12h →<br>50 mg/kg q6-8h  | 33 mg/kg q4h or 50<br>mg/kg q6h                    | 2 g q4h   |
| <b>nafcillin</b>              |   | 33 mg/kg q4h                                       | 1.5-2 g q4h   |
| <b>TICARCILLIN</b>            | 75-100 mg/kg q12h   | 75 mg/kg q6h                                       | 3 g q4h   |
| <b>GENTAMICIN</b>             | 2.5 mg/kg q12h → q8h  | 2.5 mg/kg q8h                                      | 1.66 mg/kg q8h  |
| <b>AMIKACIN</b>               | 7,5-10 mg/kg q12h → q8h   | 10 mg/kg q8-12h                                    | 7.5 mg/kg q12h  |
| <b>CEFOTAXIME</b>             | 50 mg/kg q12h →<br>50 mg/kg q6-8h   | 50 mg/kg q6h                                       | 1.5-2 g q4h   |
| <b>CEFTRIAZONE</b>            | - (displaces bilirubin from<br>albumin-binding sites)   | 40-50 mg/kg q12h                                   | 2-3 g q12h  |
| <b>CEFTAZIDIME</b>            | 30 mg/kg q12h → q8h   | 40-50 mg/kg q8h                                    | 2 g q8h   |
| <b>VANCOMYCIN<sup>R</sup></b> | 15 mg/kg q12h → q8h   | 10 mg/kg q6h                                       | Load 25-30 mg/kg<br>(max. 3000 mg)<br>↓<br>500-750 mg q6h<br>(check trough level<br>after 3 <sup>rd</sup> dose) |
| <b>CHLORAMPHENICOL</b>        | 25 mg/kg q24h → q12h  | 20-25 mg/kg q6h                                    | 1 g q6h   |
| <b>METRONIDAZOLE</b>          |   | 7.5 mg/kg q8h                                      | 500 mg q6h  |
| <b>SMX/TMP</b>                |   |  | 15-20 mg/kg/d<br>divided equally as<br>q6h or q8h doses   |
| oral <b>RIFAMPIN</b>          |   | > 1 yr.: 10 mg/kg<br>q12h<br>< 1 yr.: 5 mg/kg q12h | 600 mg q12h   |

\*may produce convulsions if large concentrations are introduced into CSF

<sup>R</sup> dose adjustment necessary for Creatinine Clearance < 60 mL/min

N.B. only **3<sup>rd</sup> generation cephalosporins** are used; CEFUROXIME enters CSF, but frequent treatment failures!

| Antibiotic   | Ratio CSF to serum |
|--------------|--------------------|
| PENICILLIN G | 2-5%               |
| AMPICILLIN   | 15-20%             |
| CEFOTAXIME   | 27-63%             |
| NAFCILLIN    | 10-15%             |
| VANCOMYCIN   | 10-15%             |

## ANTIVIRALS

- GANCICLOVIR** 5 mg/kg q12h IVI over 1 h for minimum 14-21 days → 6 mg/kg/d for indefinite period.
- FOSCARNET** 40-60 mg/kg q8h (or 100 mg/kg q12h) IVI over 1 h for 14-21 days → maintenance 60-120 mg/kg/d IV for indefinite period.
- ACYCLOVIR**
  - 10 mg/kg (or 500 mg/m<sup>2</sup>) IVI q8h over 60 min (to minimize risk of renal dysfunction).
    - dilute to concentration ≤ 7 mg/mL (e.g. 70-kg person - 700 mg is diluted in 100 mL).
    - extravasation → local inflammation and phlebitis.
    - excellent CSF penetration.
    - acyclovir-resistant strains are problem only in AIDS patients.
  - 800 mg orally ×5/d.
- VALACYCLOVIR** 1.0 g orally ×3/d.
- FAMCICLOVIR** 500-750 mg orally ×3/d.

## ANTIFUNGALS

### CANDIDA

From 2016 guidelines:

*What Is the Treatment for Central Nervous System Infections in Neonates?*

#### Recommendations

- For initial treatment, AmB deoxycholate, 1 mg/kg intravenous daily, is recommended (*strong recommendation; low-quality evidence*).
- An alternative regimen is liposomal AmB, 5 mg/kg daily (*strong recommendation; low-quality evidence*).
- The addition of flucytosine, 25 mg/kg 4 times daily, may be considered as salvage therapy in patients who have not had a clinical response to initial AmB therapy, but adverse effects are frequent (*weak recommendation; low-quality evidence*).
- For step-down treatment after the patient has responded to initial treatment, fluconazole, 12 mg/kg daily, is recommended for isolates that are susceptible to fluconazole (*strong recommendation; low-quality evidence*).
- Therapy should continue until all signs, symptoms, and cerebrospinal fluid (CSF) and radiological abnormalities, if present, have resolved (*strong recommendation; low-quality evidence*).
- Infected central nervous system (CNS) devices, including ventriculostomy drains and shunts, should be removed if at all possible (*strong recommendation; low-quality evidence*).

### XIII. What Is the Treatment for Central Nervous System Candidiasis?

#### Recommendations

- For initial treatment, liposomal AmB, 5 mg/kg daily, with or without oral flucytosine, 25 mg/kg 4 times daily is recommended (*strong recommendation; low-quality evidence*).
- For step-down therapy after the patient has responded to initial treatment, fluconazole, 400–800 mg (6–12 mg/kg) daily, is recommended (*strong recommendation; low-quality evidence*).
- Therapy should continue until all signs and symptoms and CSF and radiological abnormalities have resolved (*strong recommendation; low-quality evidence*).
- Infected CNS devices, including ventriculostomy drains, shunts, stimulators, prosthetic reconstructive devices, and biopolymer wafers that deliver chemotherapy should be removed if possible (*strong recommendation; low-quality evidence*).
- For patients in whom a ventricular device cannot be removed, AmB deoxycholate could be administered through the device into the ventricle at a dosage ranging from 0.01 mg to 0.5 mg in 2 mL 5% dextrose in water (*weak recommendation; low-quality evidence*).

Related:

#### IX. Does the Isolation of *Candida* Species From the Respiratory Tract Require Antifungal Therapy?

##### Recommendation

- Growth of *Candida* from respiratory secretions usually indicates colonization and rarely requires treatment with antifungal therapy (*strong recommendation; moderate-quality evidence*).

#### **XIV. What Is the Treatment for Urinary Tract Infections Due to *Candida* Species?**

##### *What Is the Treatment for Asymptomatic Candiduria?*

###### **Recommendations**

97. Elimination of predisposing factors, such as indwelling bladder catheters, is recommended whenever feasible (*strong recommendation; low-quality evidence*).
98. Treatment with antifungal agents is NOT recommended unless the patient belongs to a group at high risk for dissemination; high-risk patients include neutropenic patients, very low-birth-weight infants (<1500 g), and patients who will undergo urologic manipulation (*strong recommendation; low-quality evidence*).
99. Neutropenic patients and very low-birth-weight infants should be treated as recommended for candidemia (see sections III and VII) (*strong recommendation; low-quality evidence*).
100. Patients undergoing urologic procedures should be treated with oral fluconazole, 400 mg (6 mg/kg) daily, OR AmB deoxycholate, 0.3–0.6 mg/kg daily, for several days before and after the procedure (*strong recommendation; low-quality evidence*).

##### *What Is the Treatment for Symptomatic Candida Cystitis?*

###### **Recommendations**

101. For fluconazole-susceptible organisms, oral fluconazole, 200 mg (3 mg/kg) daily for 2 weeks is recommended (*strong recommendation; moderate-quality evidence*).
102. For fluconazole-resistant *C. glabrata*, AmB deoxycholate, 0.3–0.6 mg/kg daily for 1–7 days OR oral flucytosine, 25 mg/kg 4 times daily for 7–10 days is recommended (*strong recommendation; low-quality evidence*).
103. For *C. krusei*, AmB deoxycholate, 0.3–0.6 mg/kg daily, for 1–7 days is recommended (*strong recommendation; low-quality evidence*).
104. Removal of an indwelling bladder catheter, if feasible, is strongly recommended (*strong recommendation; low-quality evidence*).
105. AmB deoxycholate bladder irrigation, 50 mg/L sterile water daily for 5 days, may be useful for treatment of cystitis due to fluconazole-resistant species, such as *C. glabrata* and *C. krusei* (*weak recommendation; low-quality evidence*).

BIBLIOGRAPHY for ch. "Infections of Nervous System" → follow this [LINK](#) >>