

# Cerebral Edema

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## ETIOPATHOPHYSIOLOGY

**BRAIN EDEMA** - brain volume↑ due to increase in *extravascular* brain water.  
 – it is general reaction to insults.

N.B. differentiate from **BRAIN ENGORGEMENT** - brain volume↑ due to increase in *intravascular* volume (e.g. obstruction of cerebral veins, arterial vasodilatation).

	Cerebral Edema		
	Vasogenic	Cytotoxic	Hydrocephalic (s. interstitial)
<b>Pathogenesis</b>	Capillary permeability↑	Cellular swelling	Intraventricular fluid↑
<b>Location</b>	White matter	Gray & white matter	Ventricular, white matter (periventricular)
<b>Edema fluid</b>	Plasma filtrate	Intracellular H <sub>2</sub> O & Na	CSF
<b>Extracellular fluid volume</b>	↑	↓	↑
<b>Contrast enhancement</b>	+	-	-

**VASOGENIC EDEMA** - *increased capillary permeability* to macromolecules = *BBB disruption* (widening of tight junctions + increase in pinocytotic vesicles).

- etiology (most common type of edema!):
  - 1) **tumor**
  - 2) **abscess**, meningitis, encephalitis
  - 3) **stroke** (ischemia, infarction, hemorrhage)
  - 4) **trauma** (diffuse BBB disruption up to several hours after trauma - window of opportunity to administer cerebral protective drugs that would not penetrate intact BBB)
  - 5) **lead encephalopathy**.
- accumulates preferentially in **white matter** and can become very widespread.
  - exception is corpus callosum (so tightly bundled that there is little extracellular space - edema does not spread readily).
- paucity of brain lymphatics impairs resorption of excess fluid.
- eventually resolves (edema fluid is reabsorbed into vascular space or ventricular system).
- BBB disruption causes **CT/MRI contrast enhancement**, **CSF protein**↑.

**CYTOTOXIC EDEMA** - *swelling of cells* (neurons, glia, endothelial) due to *membrane pump failure*.

- etiology:

- a) **decreased energy supply** to brain cells (e.g. ischemia, hypoxia, trauma) → increased intracellular osmoles (Na<sup>+</sup>, lactate, H<sup>+</sup>) → rapid water entry into cells.
  - even after short ischemia, brain may respond to reperfusion with severe brain edema.
- b) **plasma osmolality** ↓:
  - 1) **osmotic disequilibrium syndromes** (in hemodialysis, diabetic ketoacidosis) - excessive *intracellular solutes* (*organic acids* in uremia; *glucose & ketone bodies* in diabetic ketoacidosis) result in excessive cellular hydration when plasma osmolality is rapidly reduced with therapy.
  - 2) acute dilutional hyponatremia
  - 3) inappropriate secretion of ADH
- c) acute **hepatic encephalopathy**, **Reye's syndrome**.
  - accumulates in **white & grey matter**.
  - extracellular fluid volume is compensatory reduced!

Conditions associated with generalized edema have elements of both **vasogenic** and **cytotoxic** edema.

- both *vasogenic* and *cytotoxic* edema occur in setting of **trauma**!
- acute **hypoxia** causes *cytotoxic* edema, which is followed by *vasogenic* edema as infarction develops.

**INTERSTITIAL (s. HYDROCEPHALIC) EDEMA** (best characterized in **obstructive hydrocephalus**) - **CSF movement across ventricular walls**.

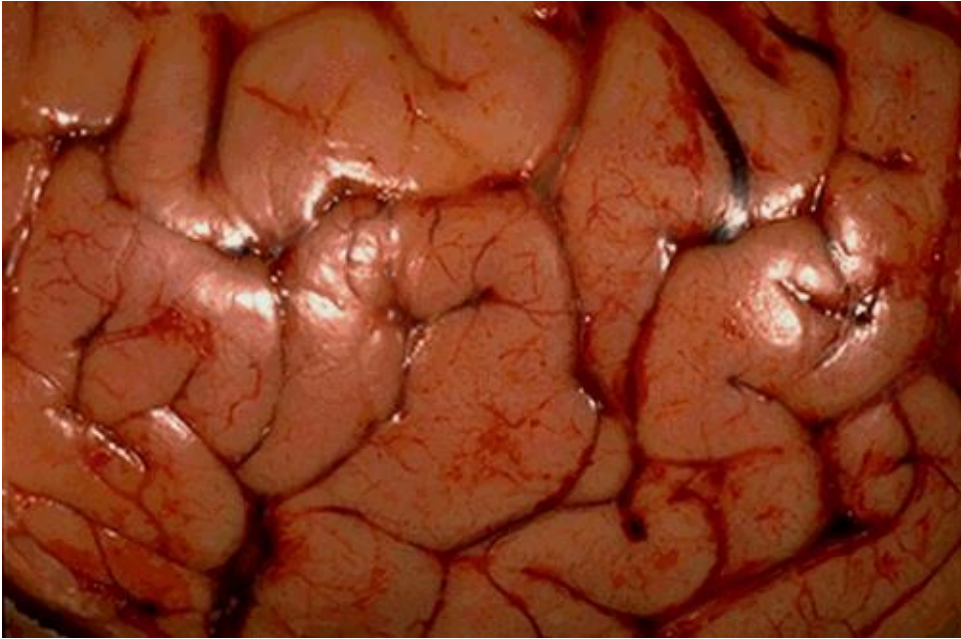
- accumulates in **periventricular white matter** (esp. at angles of lateral ventricles).
- volume of periventricular white matter is reduced! (after successful CSF shunting, edema is reduced, and thickness of mantle is restored)

**MECHANISMS** by which edema alters neuronal function:

1. ICP ↑
2. Increased distances for nutrient diffusion (e.g. O<sub>2</sub>).
3. Lipid peroxidation in membranes

## **PATHOLOGY**

- edematous brain is softer and appears to "overflow" cranial vault.
- **gyri** are flattened, intervening **sulci** are narrowed.
- **ventricular cavities** are compressed.
- as brain expands, **herniation** may occur. see S54 p.



Source of picture: "WebPath - The Internet Pathology Laboratory for Medical Education" (by Edward C. Klatt, MD) >>

Multiple small metastases causing cerebral edema seen at right which obscures structures:



Source of picture: "WebPath - The Internet Pathology Laboratory for Medical Education" (by Edward C. Klatt, MD) >>

## CLINICAL FEATURES

### - **intracranial hypertension:**

1. **Generalized brain dysfunction** (disturbances of consciousness, etc) – due to DIFFUSE edema.
2. **Focal neurologic deficits** – due to FOCAL edema, brain herniation.

N.B. in brain tumor, clinical signs are often caused more by surrounding edema than by tumor mass itself!

- *rate of edema formation* is directly proportional to *severity of neurologic deficits*.
- in chronic hydrocephalus **interstitial edema** manifestations are usually minor (in advanced cases - **dementia** and **gait disorder** become prominent) - CSF accumulation in extracellular space is much better tolerated than is presence of plasma in extracellular space (as in vasogenic edema).

## DIAGNOSIS

**Neuroimaging** (MRI better than CT):

- 1) **decreased density** of brain parenchyma (water content↑) – T1-MRI and CT signal↓, T2-MRI signal↑.
    - blurring or loss of visible distinction between gray matter and white matter.
  - 2) **mass effect**:
    - a) **diffuse edema**: loss of definition of cortical sulci → bilateral compromise of ventricles → effacement of basal cisterns.
    - b) **focal edema** - focal mass effect.
- **MRI / CT with contrast** → brain parenchymal **enhancement** in **vasogenic edema** (BBB disruption!); no enhancement in cytotoxic edema.

**CSE** - **protein**↑ in **vasogenic edema** (BBB disruption!); normal in cytotoxic edema.

**EEG**:

- a) **vasogenic edema** – slowing.
- b) **interstitial edema** – normal.

## TREATMENT

1. **Intensive care** - patent airway, avoidance of hypoxia, maintenance of BP.  
N.B. avoid salt-free fluids IV!
2. **Surgery** - excision / decompression of intracranial mass lesions, shunting procedures.
3. **Pathogenetic treatment**:

Citotoksinė edema - didink CPP + didink intravazalinį osmozinį spaudimą.  
Vazogeninė edema - mažink hidrostatinį spaudimą kapiliaruose + mažink BBB pralaidumą.

- 1) **GLUCOCORTICOIDS** (**DEXAMETHASONE** 10 mg IV or IM loading → 4 mg q6h maintenance; pediatric dose 1-2 mg/kg loading → 0.25 mg/kg qid maintenance) - direct effect on endothelial cell function – decreased BBB permeability – for **vasogenic edema** (around tumor, abscess, radiotherapy field).

Glucocorticoids dramatically and rapidly (in hours) reduce focal and general signs of brain edema around tumors!

- usual **complications** of steroid therapy are expected (esp. **gastric hemorrhage** - all patients receiving steroids for more than few hours should receive RANTIDINE and oral ANTACIDS!).
  - not useful in cytotoxic edema (e.g. questionable efficacy in head trauma, hypoxia).
  - conflicting reports about efficacy in acute bacterial or tuberculous meningitis (e.g. steroids reduce deafness in infants with bacterial meningitis).
- 2) **OSMOTHERAPY** (**MANNITOL**) – for **cytotoxic edema**.
    - effect is **short-lived** (solute reaches equilibrium concentration in brain after delay of only few hours).
    - parts of brain most likely to “shrink” are normal areas (e.g. regions of vasogenic edema with increased capillary permeability do not shrink\*).
      - \*even develop rebound edema following mannitol use because solute accumulates in edematous tissue.

- no rationale for long-term use - brain adapts to hyperosmolality with increase in intracellular osmolality.
- 3) **DRUGS THAT REDUCE CSF FORMATION** (ACETAZOLAMIDE, FUROSEMIDE) – for interstitial edema.

BIBLIOGRAPHY see p. S50