Posterior Pituitary Hormones
ADH (Vasopressin) & Oxytocin

Nonapeptides (9 a.a)

Known as neurohormones

Synthesized in the hypothalamus

Stored in the posterior pituitary → release

? Role as neurotransmitters ($V_1$R’s in CNS)

Role of Oxytocin in man is unknown
**ADH (Vasopressin)**

Physiological and pharmacological actions:

- **Vasoconstriction** ($V_1$ receptors)
- **↑ reabsorption of $H_2O$ from collecting ducts** ($V_2$ receptors)
- **↑ synthesis of certain clotting factors** (VIII, Von Willebrand) ($V_2$ receptors)
- **↑ ACTH release** ($V_3$ receptors)
- **Oxytocin-like activity**
Factors/Drugs ↑ ADH release:
- Hypovolemia, hyperosmolarity, pain, stress, nausea, fever, hypoxia
- Angiotensin II
- Certain prostaglandins
- Nicotine, cholinergic agonists, β-adrenergics
- Tricyclic antidepressants
- Insulin, morphine, vincristine...
Factors/Drugs ↓ ADH release:
- Hypervolemia
- Hypoosmolality
- Alcohol
- Atrial natriuretic peptide
- Phenytoin
- Cortisol
- Anticholinergics, α-adrenergics, GABA...
Disorders affecting ADH release:

A. Excess production (inappropriate ADH secretion) → Dilutional hyponatremia

Causes:
- Head trauma, encephalitis
- Meningitis, oat cell carcinoma

Rx:
- Water restriction (Rx of choice)
- Hypertonic saline solution
- Fludrocortisone → ↑ Na⁺ blood level
- ? ADH antagonists
- **ADH antagonists**
  - Conivaptan, $V_1$ & $V_2$ R antagonist given IV
  - Tolvaptan; Lixivaptan & Satavaptan, orally effective selective $V_2$R antagonists

**Clinical uses:**
- Inappropriate ADH secretion
- CHF
B. Deficiency of ADH → Diabetes insipidus (DI) → polyuria

Causes:

- Idiopathic DI
- Congenital, Familial DI
- Hypothalamic surgery, head trauma, malignancies
- Gestational DI, overproduction or decreased clearance of vasopressinase

Rx:

ADH preparations (HRT)
**ADH preparations:**

- **Natural human ADH (Pitressin)**
  Given I.M, S.C, has short half-life (15 min)

- **Lypressin (synthetic, porcine source)**
  Given intranasally, I.V, I.M, has short DOA (4hrs)

- **Desmopressin (synthetic ADH-like drug)**
  Given intranasally, S.C
  Most widely used preparation, has long DOA (12 hrs)
- Felypressin (synthetic ADH-like drug)
  Has strong vasoconstrictor activity
  Mainly used in dentistry
  ■ Clinical uses to ADH:
  - DI
  - Nocturnal enuresis
  - Hemophilia
  - Bleeding esophageal varices
Side effects to ADH preparations:

- Allergy
- Pallor
- Headache, nausea, abdominal pain in ♀’s (oxytocin-like activity)
- Anginal pain (coronary artery vasospasm)
- H₂O intoxication (massive doses)
- Gangrene (rare particularly with desmopressin= has great affinity to V₂ receptors)
Drugs acting on the uterus
I. Uterine stimulants

1. Oxytocin: (nonapeptide=9 a.a peptide)
   - Contracts the myoepithelial cells of the breast → milk letdown; milk ejection
     Major stimuli, baby cry and suckling
   - Contracts the uterus → delivery

The uterus is insensitive to oxytocin in early pregnancy but its sensitivity increases with advanced pregnancy reaching maximum at time of delivery
   - Has slight ADH-like activity
**Oxytocin MOA:**

- Surface receptors $\rightarrow$ stimulation of voltage-sensitive Ca$^{++}$ channels $\rightarrow$ depolarization of uterine muscles $\rightarrow$ contractions
- ↑ intracellular Ca$^{++}$
- ↑ prostaglandin release
**Clinical uses to oxytocin:**

- Induction of labor
  Drug of choice given in units in an I.V infusion
- Postpartum hemorrhage, I.M. Ergot alkaloids are better (ergonovine, methylergonovine, syntometrine = oxytocin + ergometrine)
- Breast engorgement, intranasally
- Abortifacient, I.V infusion. ≥ 20 weeks of gestation, ineffective in early pregnancy
- **Side effects to oxytocin:**
  - Rupture of the uterus
  Major and most serious side effect
  - H$_2$O intoxication and hypertension
  Due to its ADH-like activity
- **Specific oxytocin antagonist**
  Atosiban (inhibitor to uterine contraction=tocolytic),
effective in the management of premature delivery,
given IV
2. Prostaglandins:

* Dinoprostone (PGE$_2$)
  Vaginal pessaries, inserts and gel, tab
  Abortifacient, induction of labor

* Dinoprost (PGF$_{2\alpha}$)
  I.V infusion and intramniotic
  Same uses as dinoprostone
* Carboprost (PGF$_{2\alpha}$)  
I.M and intramniotic  
Abortifacient and postpartum hemorrhage  
* Gemeprost (PGE$_1$)  
Vaginal pessaries  
Used to prime the cervix  

3. **Ergot alkaloids:**  
Ergonovine, Methylergonovine  
I.M, oral
Ergot alkaloids remain the drugs of choice to manage postpartum hemorrhage

As compared to oxytocin, ergot alkaloids are more potent, they produce more prolonged and sustained contractions of the uterus and they are less toxic

Ergot alkaloids are contraindicated to be used as inducers to delivery (associated with high incidence of fetal distress and mortality)
II. Uterine relaxants (Tocolytics)

Major clinical use: premature delivery (weeks 20-36) → improve the survival of the newborn

1. β-adrenergic agonists:
   ↑ cAMP → ↓ cytoplasmic Ca^{++}

* Ritodrine
  I.V infusion

Most widely used

* Terbutaline, Oral, S.C, I.V
Side Effects to $\beta$-adrenergics:
Sweating, tachycardia, chest pain...

2. Magnesium sulfate
I.V infusion

Activates adenylate cyclase and stimulates $\text{Ca}^{++}$ dependent ATPase

Uses: premature delivery and convulsions of pre-eclampsia
3. Progesterone
Oral, I.M
Dydrogesterone
4. Oxytocin competitive antagonists
Atosiban
5. Prostaglandin synthesis inhibitors
Indomethacin, Meloxicam
6. Nifedipine
** Major contraindication to tocolytics: fetal distress