

# Pharmacology of Growth hormone and Pituitary Adenomas

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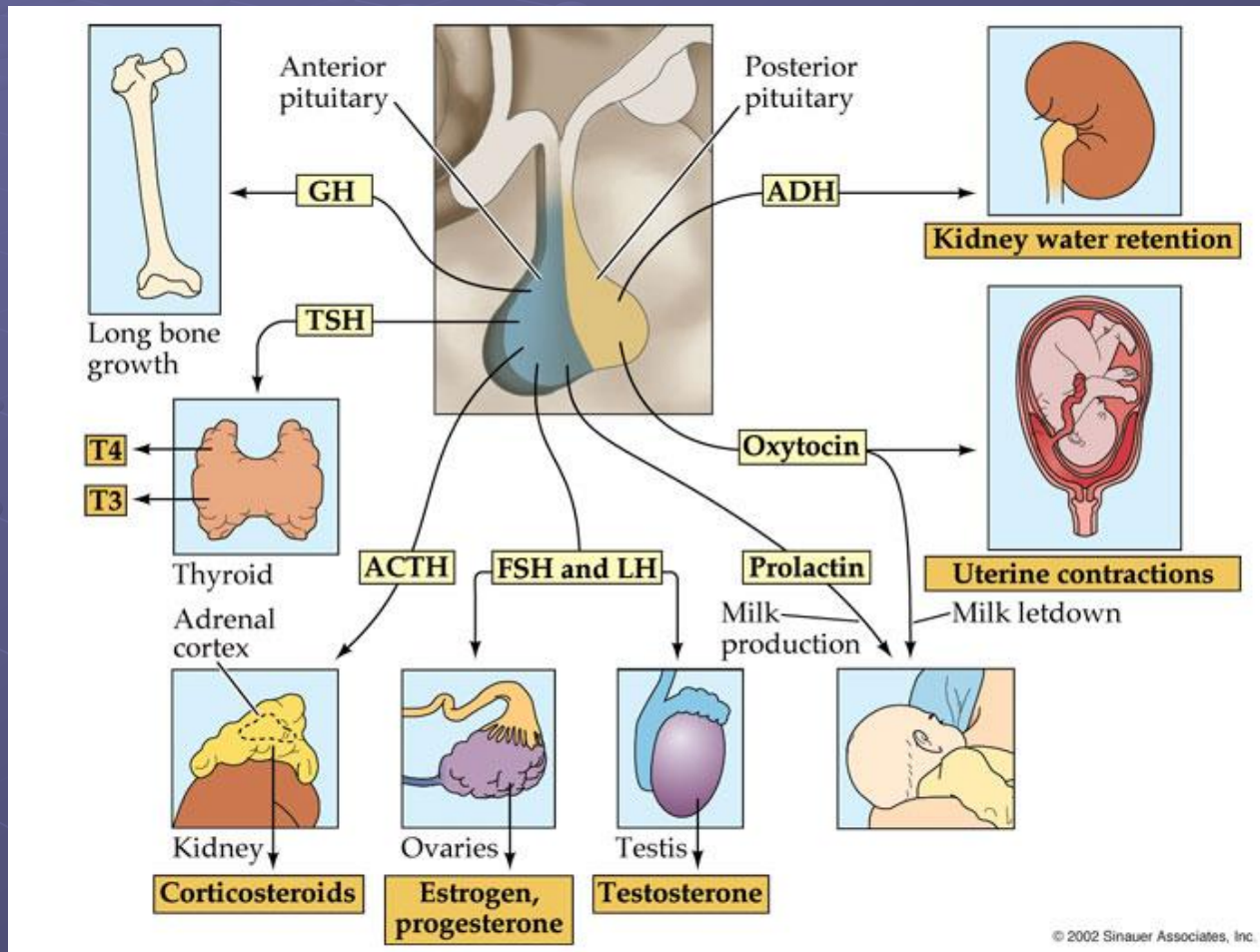
# Pituitary and Hypothalamus

**Pituitary and hypothalamus are the link between the nervous system and the endocrine system.**

Hypothalamus is also *major regulator of body homeostasis*

1. Homeostatic control includes regulating hunger, thirst, sex drive, sleep-wake cycles, body temperature, blood glucose.
2. Endocrine control via regulating the release of pituitary hormones.
3. Autonomic control via descending pathways to sympathetic and parasympathetic preganglionic neurons.
4. Limbic function via connections to limbic system regulating emotional behavior.

# A 'global' view of hypothalamic pituitary functions



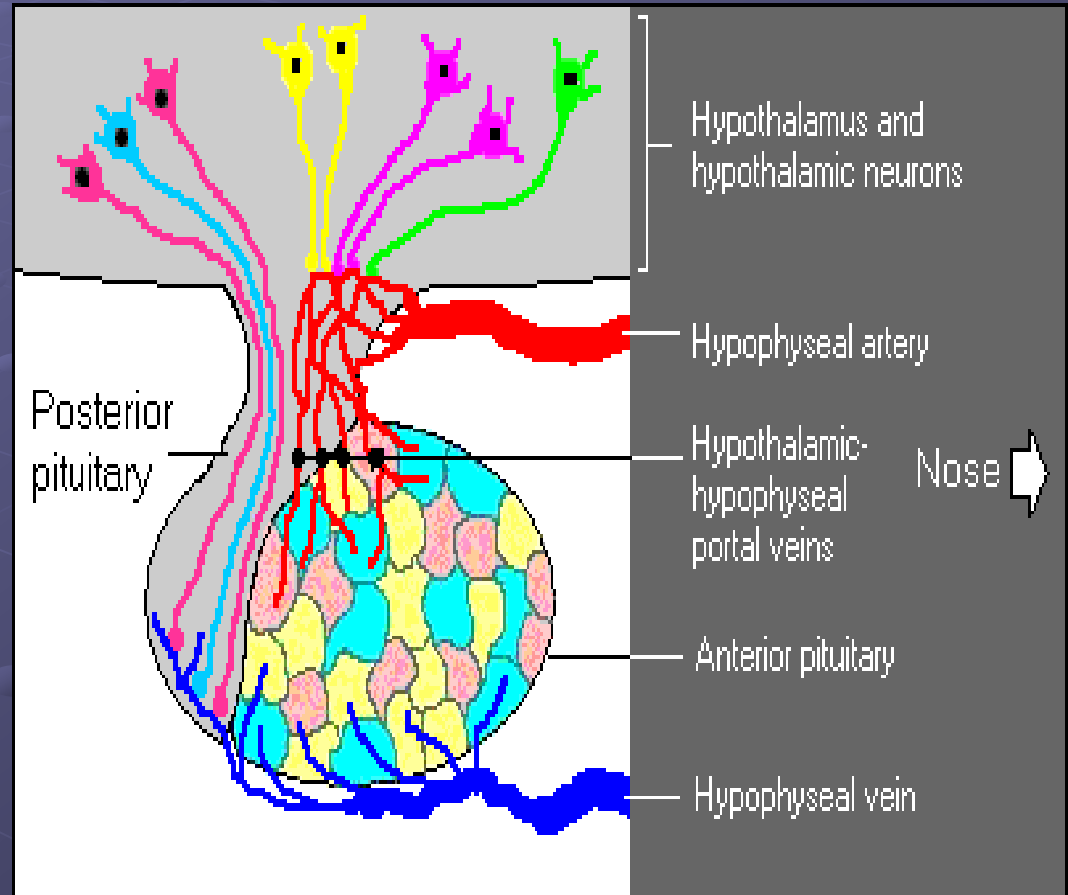
# Function

## ● Anterior Lobe:

- FSH
- LH
- ACTH
- TSH
- Prolactin
- GH

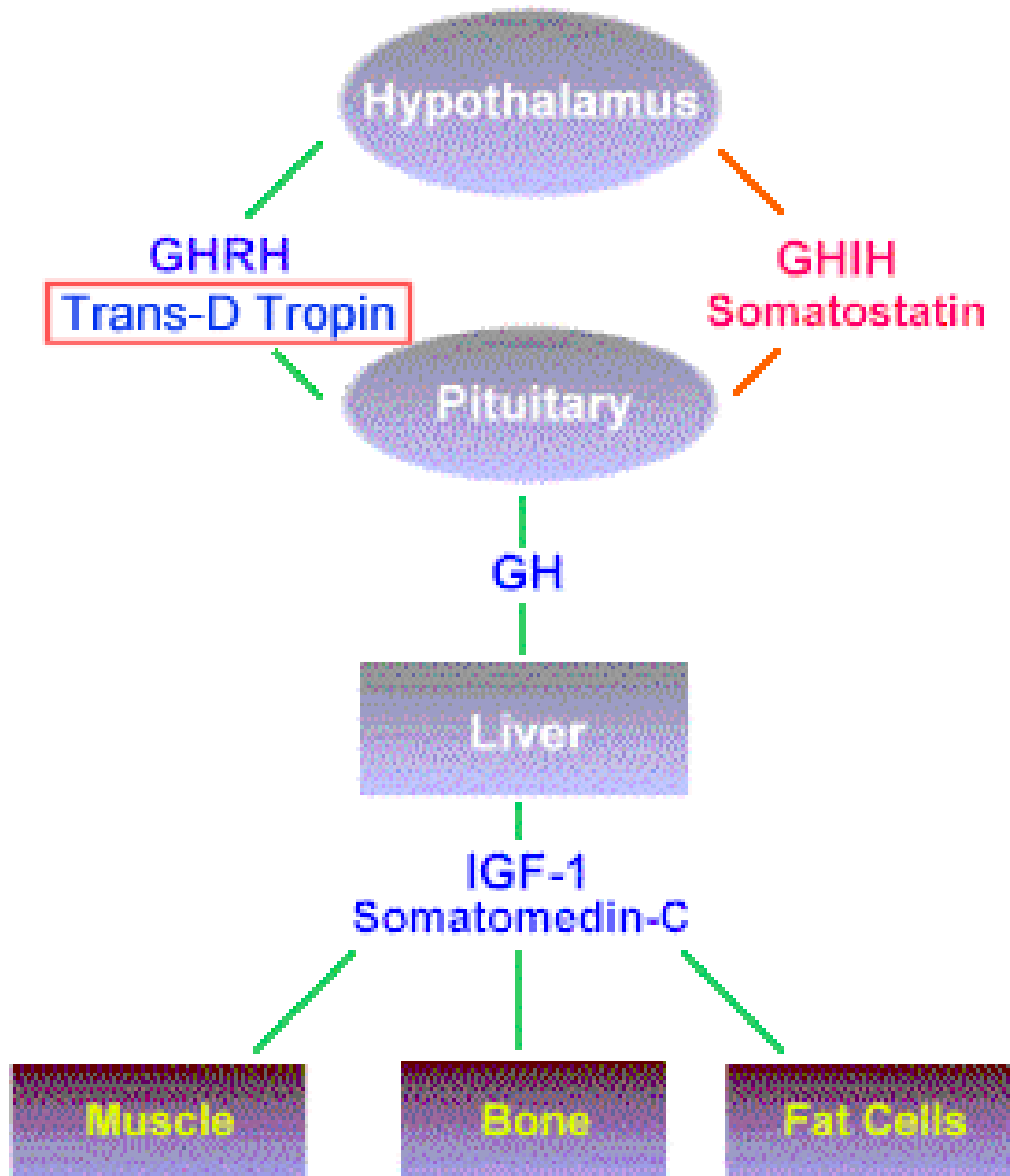
## ● Posterior Lobe:

- ADH
- Oxytocin



# Anterior Pituitary: Growth Hormone (GH)

- **Stimulates increase in size and mitotic rate of body cells, increases fat utilization**
- Hypothalamic growth hormone releasing hormone (**GHRH**) stimulates secretion of GH; **Somatostatin (SS)** inhibits secretion of GH
- **Enhances** amino acid movement through membranes and promotes **protein synthesis**
- **Promotes long bone growth**



Deficiency or absence of  
**somatotroph** cells



Underproduction of  
growth hormone  
(somatotrophin)



## **PITUITARY DWARF**

(Lorain Dwarf)

Delayed skeletal growth  
and retarded sexual  
development but alert,  
intelligent, well  
proportioned child.

Functional overactivity (or tumour) chiefly of the **SOMATOTROPH** cells of the anterior pituitary leads to **GIANTISM** in the CHILD: **ACROMEGALY** in the ADULT.

↓  
Overproduction of *growth Hormone*

↓  
↑ *IGF-1 (somatomedin C)*

↓  
Stimulates protein synthesis.  
Influences carbohydrate and fat metabolism and mitosis of **ALL CELLS** of the body

Overgrowth of all body tissues

Onset before bony epiphyses have closed at puberty

Onset after puberty



# Direct Effects of GH

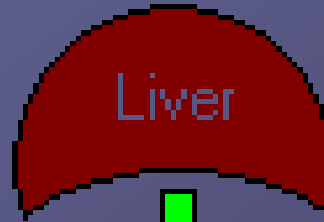
- Binds to adipocytes and causes them to break down triglycerides and prevents them from accumulating fat in the blood
- Releases insulin-like growth factor-1 (IGF-1) from the liver

# Indirect Effects of GH

## ● Stimulates:

- Cartilage cells (chondrocytes) growth
- Myoblasts growth and differentiation
- Amino Acid uptake
- Protein synthesis

**Growth  
Hormone**



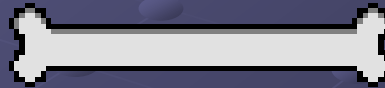
Liver



Fat



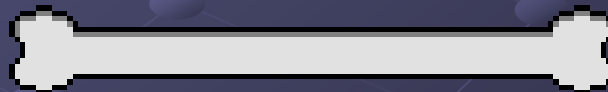
**IGF-1**



Direct  
effect



Indirect  
effect



## **Mechanism of Action:**

Binding of GH to its receptor activates the signaling cascade mediated by receptor associated to JAK tyrosine kinases

The effects of GH are primarily mediated by insulin-like growth factor 1. (IGF-1) released by liver in response to GH.

# Pituitary adenoma

Pituitary adenoma is a benign tumor of the anterior lobe of the pituitary that causes symptoms either by

**Underproduction:** growth hormone deficiency, major problem in children's growth, hypothyroidism,

or **overproduction** of the pituitary hormones

**Growth hormone excess resulting in acromegaly or gigantism .**

**Prolactin excess leads to galactorrhoea, menstrual abnormalities and infertility**

Cushing disease resulting from adrenocorticotrophic hormone (ACTH).

# Clinical Presentation

- Prolactin – Amenorrhea, galactorrhea, impotence
- Growth hormone – Gigantism and acromegaly
- Corticotropin – Cushing's disease,
- TSH - Hyperthyroidism

# Pharmacology of Growth Hormone Deficiency

Drugs Used:

Synthetic GHRH (Sermorelin)

Recombinant human growth hormone  
(Somatropin, Somatrem)

Recombinant IGF1 (Mecasermin)



**Sermorelin:** It is used if a patient possesses defective hypothalamic release of GHRH but normally functioning anterior pituitary somatotrophs

Treatment with Recombinant Human Growth Hormone (**Somatropin**, Somatrem)

**Somatropin** (synthetic growth hormone), which is a 191-amino acid peptide, identical to the native form of hGH

## **GH Indications:**

Documented growth failure in pediatric patients associated with: GH deficiency and Turner syndrome (increase height in girls )

Idiopathic short stature

Wasting in patients with AIDS

Short bowel syndrome in patients who are also receiving specialized nutritional support

## Side Effects:

Leukemia,

rapid growth of melanocytic lesions

Hypothyroidism

Insulin resistance

Arthralgia

Increase in cytochrome P450 activity

# Treatment with Recombinant IGF1 (Mecasermin)

**Mecasermin** is used for children with severe IGF1 deficiency due to mutations in the GH receptor (Laron dwarfism) or development of neutralizing antibodies against GH. Its administered S.C, the common adverse effect is hypoglycemia, can be avoided by consumption of meal 20 min before or after the administration of drug.

## Features of Growth Hormone Excess

This usually results from benign tumor of the anterior pituitary.

(1) In children: **It causes gigantism**. occurs before the closure of epiphyses, because excess IGF1 causes excessive longitudinal bone growth

(2) In adults: It causes **acromegaly** (bones increase in size, including those of hands, feet and face).

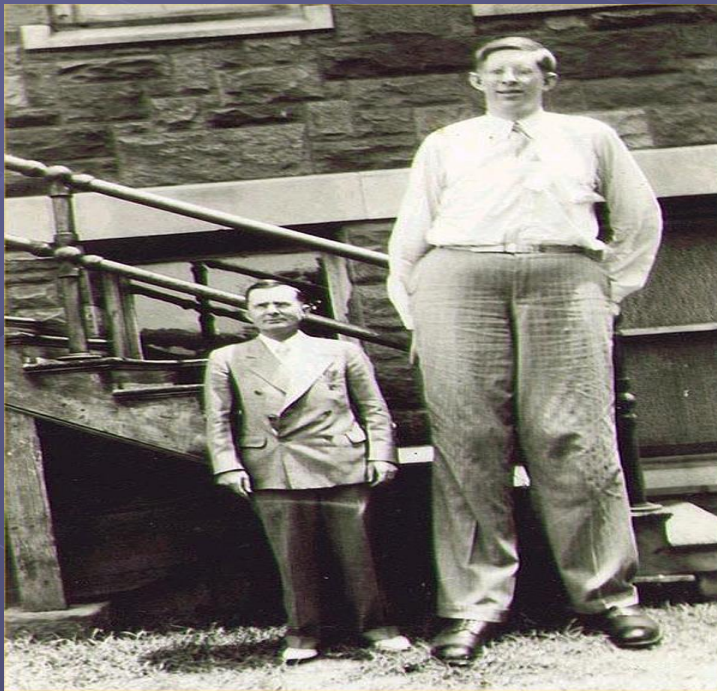
Growth retardation  
(dwarfism)



ADAM.



Hand comparison of twins - one on the left has acromegaly.  
R Gagel, IE McCutcheon. Pituitary Gigantism. NEJM  
1999;340:524. 1999.



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# Growth Hormone Antagonists

Drugs Used:

Somatostatin analogues (Octreotide S.C, IM, Lanreotide (I.M))

GH receptor antagonist (Pegvisomant)

Dopamine receptor agonist **only high doses** (Bromocriptine - described under hyperprolactinemia)



# Growth Hormone Antagonists

**Somatostatin analogues:** Somatostatin physiologically inhibits GH secretion, but is rarely used clinically, since it has a very short half-life ( a few minutes)

**Octreotide** is a synthetic long-lasting peptide analogue of somatostatin (45 times more potent)

Side effects : Octreotide and lanreotide cause significant gastrointestinal disturbances, gallstones, and cardiac conduction abnormalities

# Somatostatin analogues:

- **Octreotide (very expensive)** :45 times more potent.
  - half-life in plasma being 113 min
  - suppress GH levels for 6–12 h
  - Given every 4 weeks
  - Mechanism of action
    - Inhibit GH secretion
    - partially inhibits GH-induced IGF-1 generation
    - reduce GHRH release

	Octreotide (S/C) 100 to 500 mic.gm TDS	Octreotide (I/M) at 28 days interval	Lanreotide (I/M) every 7-14 days	Pegvisomant
GH REDUCTION	47%	56%	50%	Not useful
IGF1 REDUCTION	46%	66%	48%	97%

Freda PU:clinical review 150:somatostatin analogs in acromegaly.j clin endocrinol metab 87:3013-3018,2002

# Dopamine agonists :

- Used both as primary and adjuvant treatment
  - Bromocriptine up to 20 mg/day
  - Cabergoline 1–2 mg/week
- Response rate low

# Dopamine agonists :

	Bromocriptine	Cabergoline
GH REDUCTION	20%	44%
IGF1 REDUCTION	10%	35%

Freda PU:clinical review 150:somatostatin analogs in acromegaly.j clin endocrinol metab 87:3013-3018,2002

# GH-Receptor Antagonist :

- Pegvisomant given s.c:
- Check IGF 1 level every 4-6 weeks
- Monitoring GH not useful
- Dose 10-40 mg/d

# Growth Hormone Antagonists

## Pegvisomant

Pegvisomant is a GH receptor antagonist approved for treatment of acromegaly.

Normally, GH, which has 2 distinct receptor binding sites, initiates cellular signaling cascades by dimerizing 2 GH receptors.

Pegvisomant is a long-acting derivative of a mutant GH that is able to cross-link GH receptors but is incapable of inducing the conformational changes required for receptor activation.

## Dopamine D<sub>2</sub> Receptor Agonists

Dopamine D<sub>2</sub> receptor agonists such as **bromocriptine** are more effective at inhibiting prolactin release than inhibiting GH release. However, high doses of D<sub>2</sub> receptor agonists have some efficacy in the treatment of small GH-secreting tumors.



# Prolactinoma (pituitary adenoma with excess release of prolactin)

- Initial therapy is generally **dopamine agonists**. **Bromocriptine**, a dopamine agonist, is generally given orally, ergot derivatives.
- **Cabergoline** is given once or twice weekly. Better tolerated and more effective than bromocriptine for tumor shrinkage **but more expensive**. Side effects include **orthostatic hypotension, nausea, and dizziness**; avoided by beginning with **low-dose therapy**.
- Other compounds include **pergolide mesylate**, a **long-acting ergot derivative** with dopaminergic properties **but strong vasospasm and uterotonic**.

## Dopamine agonists:

- Bromocriptine
- Cabergoline.
- Pergolide mesylate

Side effects– GI intolerance, postural hypotension, constipation, **nasal stuffiness**

# Mechanism of action of Dopamine agonist

Selective activation of D2 receptors located on lactotroph cell surface



Decrease adenylate cyclase activity



Decrease in C-AMP level



Inhibition of PRL synthesis and release.

# Bromocriptine:

- (Purpose was inhibiting prolactin secretion without the uterotonic, vasospastic properties of **other ergots**)
- **Bromocriptine is safer in pregnancy**

# Bromocriptine:

- The absorption rate from the GI tract is 25-30%.
- Very high first-pass effect, with 93.6% of a dose being metabolized and only 6.5% of an absorbed dose reaching the systemic circulation unchanged
- Excreted via the biliary route into the feces
- start low dose at 2.5 mg day at night before increasing to 2.5 – 10 mg per day in divided doses
- **Take with food** to reduce side effects

## Cabergoline (Ergot drug) :

- more effective
- Well tolerated but not safer in pregnancy
- **more expensive**
- given once or twice a week with a starting dose of 0.25 mg 2 x week

Titrate these based on prolactin levels and tolerability



THANK YOU