8. ENDOCRINE PHYSIOLOGY

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PRINCIPLES OF ENDOCRINOLOGY

tenanc preadstropin

1. What is a hormone? What is a hormone is classically considered a substance produced in small amounts, re-as endocrine hormone is classically considered as a substance produced in small amounts, re-As endocrine hormone as consequently consequent a substance produced in small amounts, re-as a more blood where it is transported to a distant organ to exert its specific action on anget seed into the blood with a receptor for the hormone. A hormone can also act on periodopaid into the blood where the formone. A hormone can also act on neighboring tissue size of support with a receptor for the hormone. A hormone can also act on neighboring tissue size of support of the formone and can act on the tissue that produced it (annually size of sums gland (paracrine) and can act on the tissue that produced it (annually size of sums gland (paracrine)). is a supple with a respective to an action the tissue that produced it (autocrine). However, and is seen a gland (paracrine) and can act on the tissue that produced it (autocrine). However, a glas be synthesized and released into the bloodstream by nerves (neuronal and also be synthesized). 1800 - Sume gland (put not 100.) and cent act on the fissue that produced it (unterine) with the sume gland (put not 100.) and the bloodstream by nerves (neurocrine), most can also be synthesized and released into the bloodstream by nerves (neurocrine), most can also be synthesized.

- 1 List general functions that hormones regulate. List general to menstrual cycle, ovulation, spermatogenesis, pregnancy, lactation agreement secured differentiation Reproduction pregnancy, inclusion from an development — sexual differentiation, secondary sex characteristics, growth
 - vehicity

 Maintenance of the internal environment—extracellular fluid volume, blood pressure,
 Maintenance of the internal environment of playme for such as artistically Maintenance, and regulation of plasma ions such as calcium and sodium
 - Energy flux storage, distribution, and consumption of calories; heat production Behavior - food and water intake, sexual behavior, mood

t What is the chemical nature of classic hormones, and where are they produced? The chemical nature of hormones is determined by the site of synthesis and, in turn, deterde of transport in blood, the mechanism of action, and rate of metabolism.

	SITE OF RELEASE	HORMONE
es and tyrosine	Adrenal medulla	Catecholamines (epinephrine, norepinephrine, dopamine)
detratives	Thyroid gland	Triiodothyronine (T3), thyroxine (T4)
å	Gonads and placenta Adrenal cortex Diet/skin/liver/kidney	Testosterone, estrogens, progesterone Cortisol, aldosterone, adrenal androgens Secosteroids (vitamin D and its metabolites)
optides proteins	Posterior pituitary Hypothalamus Amerior pituitary Gastrointestinal Endocrine pancreas Placenta Parathyroid/thyroid Many others	Oxytocin, vasopressin TRH, somatotstatin, GnRH, CRH, GHRH MSH, ACTH, prolactin, GH Gastrin, somatotstatin, cholocystokinin, secretin Insulin, gluczgon, pancreatic polypeptide Chorionic somatomamnotropin PTH, calciformin Examples: heart (atrial natriunetic peptide), liver (IGF1)
*Heins	Anterior pituitary Placenta	LH, FSH, TSH hCG gonadotropin-releasing hormone, MSH = melan

**Secretarion of the control of the to ACH = screece, CRH = corticotropin-releasing hormone, GHRH = growth hormone-resonne, LH = to the screece or to the sc TH = deteocorricotropic formone, GH = growth hormone, PTH = parathyroid normone, http://december.com/december

Endocrine Physiology

4. What are some examples of different categories of hormones?

Thyrold Hormones

Amines

Polypeptides

200

Proteins , accommonance

(Oxytocin)

Steroids

(Vitamin D3)

Examples of different categories of hormones. In the case of the protein hormone, each circle represents an amino acid, as shown for the polypeptide hormone. The structure of oxytocin is similar to arginize visopressin. (From Griffin JE, Ojeda SR (eds): Textbook of Endocrine Physiology, 3rd ed. New York, Oxford University Press, 1996, p 7, with permission.)

5. Is there a pattern to the release of hormones?

Hormones are released with a variety of rhythms. Hormones can be released in circudian rhythm, such as cortisol, which peaks at 8 A.M. and reaches its nadir at midnight in diurnal anmals. Hormones can be released in ultradian rhythm, with many regular pulses within a day (e.g., luteinizing hormone [LH]), and even have seasonal rhythms. Hormones can also be released primarily in response to specific stimuli (e.g., suckling-induced prolactin). Pulsatility appears to maintain receptor sensitivity to hormones.

6. What are the general principles of the control of hormone secretion?

The end-product (hormone, metabolite) inhibits the release of the hormone that stimulated the production of the end-product (feedback loop). Most hormones are under negative feedback (the production) of the end-product (feedback loop). Most hormones are under negative feedback (the production) of the end-product (feedback loop). (thermostatic) control. For example, glucagon stimulates glucose production; an increase in plasma glucose shuts off glucagon production.

(ampare and contrast the general mechanism of action of each class of hormone. and contrast use section and action of each class of hormone.

(support and contrast use section (increases or decreases) of an existent cellular section (increases or decreases) of an existent cellular section (increases or decreases). (supplementations) which activates or inhibits a secliked formulates which activates or inhibits a secspecific that alters the function (increases or decreases) of an existent cellular component
of viva an intracellular receptor, which activates specific gene transcellular component
of viva an intracellular component or viva have refer that allers are ceptor, which activates specific gene transcription and trans-defended by or via an intracellular receptor, which activates specific gene transcription and trans-defended by or via an intracellular component [e.g., pump]. Someon of trans-report of the component component is a functionally transcription and transa projection (synthesis of a new cellular component [e.g., pump]), or via an interest component [e.g., pump]), or via an interest component [e.g., pump]). Or pumple of the projection (synthesis of a new cellular component [e.g., pump]). Or pumple of the projection and transfer of the projection (synthesis of a new cellular component [e.g., pump]). Or pumple of the project of the (85-m) and the protein to influence gene expression, and some intracellular hormone receptors also influence gene expression, and some intracellular hormone receptors.

Market and Chothic III			roceptors ma
at his surgenomic in	RECEPTOR LOCATION	SECOND MESSENGER	_
HURMUNE Thoroid Stenid Popide Cuecholamine Popide	Nuclear Cytoplasmic Cell membrane Cell membrane Cell membrane	Transcription Transcription cAMP/cGMP cAMP/cGMP	Slow Slow Fast Fast Fast
1.5			

$_{8.\,List\,some}$ of the types of cell membrane receptors, and give an example of a hormone and System-transmembrane domain receptor: This classic cell membrane receptor is covered

Seen-transmitter 2. The β-adrenergic receptor (catecholamine ligand) is the classic model. a deal in Chapter

a deal in Chapter

Bet receptors interact with another family of proteins (G-proteins) that mediate changes in

The receptors interact with another family of proteins (G-proteins) that mediate changes in Their receptors into the mediate changes in service cyclase activity and cyclic adenosine monophosphate (cAMP) production and turn on te classic phosphorylase cascade. classic prospectors of the provided straining and the provided straining and the provided straining and provided straining and the provided straining and th

round of the phosphorylation of tyrosine on intracellular proteins.

Guanylate cyclase-linked receptors: These receptors (e.g., for atrial natriuretic peptide) grante the production of the second messenger cyclic guanosine monophosphate (cGMP). Cytokine receptor superfamily: Growth hormone (GH) and prolactin are examples of hos-

more that bind to these receptors, which activate tyrosine phosphorylation despite no apparent torology to known protein kinases.

9. Briefy describe the different second messengers that mediate the action of the cell surher hormone receptors. Second messengers quickly transduce and amplify the signal generated by the binding of

the homes to the cell surface receptor. Among these second messengers are cAMP, cGMP, the akim-calmodulin system, and the phosphatidylinositol-diacylglyceride-inositol 1,4,5 triphosplane (IP.) system. The details of each of these can be found in Chapter 2. Briefly, although they arquit different in their biochemistry, the end result of each is the same in that they quickly act on an intracellular element either to inhibit or to activate some function (e.g., enzyme, pump, nembrane potential, calcium release).

ik List and briefly describe some of the types of intracellular hormone receptors.

The intracellular receptors work mostly by altering gene expression. This is why they generated by have a slower onset of action than cell membrane receptors, which quickly activate second between Stroid and thyroid hormones bind either to a cytoplasmic receptor that is translosaid in the nucleus or to nuclear receptors. The binding of steroid to the receptor either liberates the complex from heat-shock proteins (e.g., cortisol) or directly activates the receptor already hand to its respective hormone-responsive elements (HRE) on DNA (e.g., thyroid hormone, esbigst, 125(OH), D). Either way, the activated receptor-ligand forms a pand application. This am, binds to its HRE, and activates transcription of specific genes (mRNA production). This increase is the specific genes (mRNA production). The complex transcription of specific genes (mRNA production). lecused in specific mRNA results in the synthesis (translation) of specific proteins (e.g., enzyme 11. List the general features of the metabolism (clearance) of hormones. List the general features of the latest plasma bound to carrier proteins. Hormones are transported in plasma bound to carrier proteins. Hormones are the compartment; usually only the free (unbound) compared to the compartment of the compartm

 Some hormones are transportment; usually only the free (unbound) component of the circulated from the plasma compartment; it is the unbound hormone that is from the circulated for metabolism. It is the unbound hormone that is from the circulated from th tabolized from the plasma comparation, it is the unbound hormone that is free to exert a bi. ologic action.

gic action.

2. Metabolic clearance is inversely proportional to the percent of total hormone circulating.

A metabolic clearance (long half-life). Metabolic clearance is invested in the bound form. Thyroid hormone has a slow metabolic clearance (long half-life) because it in the bound form. Thyroid hormone in the plasma compart. in the bound form. Thyroid normanic has a character in the plasma compartment protects the circulates > 99.6% bound. Protein binding of a hormone in the plasma compartment protects the circulates > 99.6% bound. Protein officing the free hormone is biologically active and available for hormone from metabolism because only the free hormone is biologically active and available for

metabolism. abolism.

3. Within a class of hormones, metabolic clearance is also inversely proportional to protein 3. Within a class of intimional protein and the steroid cortisol circulates 95% bound and has a slower metabolic clearance than aldosterone, which circulates only 15% bound.

12. Discuss the general principles of endocrine disease.

In general, most disorders that are primarily attributable to hormones result from either their real or apparent underproduction or real or apparent overproduction.

Underproduction:

 Primary underproduction is due to loss of the function of the gland producing the active hormone. An example is destruction of both adrenal glands (primary adrenal insufficiency)

 Secondary underproduction is due to the loss of the hormone that normally stimulates the gland producing the active hormone. An example of this is hypopituitarism, in which the pimitary fails to produce trophic hormones (e.g., adrenocorticotropic hormone [ACTHh. which maintain normal function of a gland (e.g., adrenal cortex).

 Apparent underproduction (target cell insensitivity) is usually due to a receptor defect (mutation) such that, even if the hormone is present, the target cell cannot respond. An example of this is testicular feminization, in which a male genotype (XY) fetus has a mutation in the testosterone receptor and, as a result of loss of androgen activity, develons a fe. male phenotype. Another example is pseudohypoparathyroidism, in which, despite normal or elevated parathyroid hormone (PTH) levels, the target cells for PTH cannot respond.

Overproduction: · Primary overproduction is usually due to a neoplasm (tumor) arising from a cell population that normally produces the hormone such that the hormone is produced in excess regardless of any endogenous signal to stop its production. An example is an adrenocortical

adenoma that produces cortisol even in the absence of ACTH. · Secondary overproduction is due to excess input into the target gland. An example is a tumor arising from normal pituitary cells and producing too much trophic hormone (e.g., ACTH) such that an otherwise normal adrenal cortex is told to produce too much cortisol. Another example is secondary hyperparathyroidism, in which calcium, which inhibits PTH release, is not absorbed properly in the gastrointestinal tract, and PTH release is greatly in-

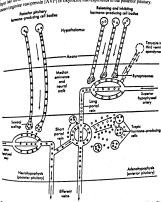
creased to compensate for it. · Apparent overproduction is due to activation of a receptor or cellular component owing to a mutation. Therefore, the function of the target gland is activated even in the absence of normal hormonal stimulation. An example of an activating mutation is Liddle's syndrome, in which the renal epithelial sodium channel is constitutively activated and mimics the effects of too much aldosterone, even though aldosterone is low.

PITUITARY PHYSIOLOGY

13. Describe the functional anatomy of the hypothalamic-pituitary interface. The control of anterior and posterior pituitary hormone release is a classic example of neuroendocrine systems. The anterior pituitary (adenohypophysis, pars distalis) is controlled by hypothalamic releasing or inhibiting (hypophysiotropic) hormones synthesized in parvocellular neurons with cell bodies in nuclei in the hypothalamus (generally medial nuclei such as arcuate and medial

profit in the second bodies increases or decreases the release of stimulatory freezagarcerishilar). (again to see which are released from terminals located on capillaries in the median spirit within a person of the second and the second are transported to the anterior pinnion.

They care the long portal blood vessels and are transported to the anterior pinnion. programment of published years and a partial blood vessels and are transparred to the anterior pituliary. The metastands of judicial the release of hormones from the anterior pituliary. The metastands of judicial the release of hormones from the anterior pituliary. The metastands of judicial the release of hormones from the anterior pituliary. is the control of the anterior pituliary, where provided in the anterior pituliary, where provided in the inhabit the release of hormones from the anterior pituliary. The posterior pituliary of pituliary of pituliary in the place of the pituliary of the pitulia emony, when the separation of pure money when the separation of polygonial and the blood from axion with magnetion of separation of pure money and particular fluorial) muclei of the separation of the separati programment of the supersymmetric and paravertificular (alternal) muclei of the hypothalarma, while the supersymmetric and paravertificular (alternal) nuclei of the hypothalarma, while the supersymmetric and paravertificular (alternal) muclei of the hypothalarma. or dilate of houses not be the fine time of decrease in the release of posterior pituliary hor-ing in the second posterior pituliary hor-lage in a decrease in [AVI] or oxytocin) into capillaries in the posterior pituliary hor-ing and the posterior pituliary hor-We had been considered the posterior plus when the posterior plus and properties in the posterior plus agent organize various examples in the posterior plustary,



he factive al masons of the hypothalamic-pituitary interface and its blood supply. Arrows indicate the dithe state of the s Parayrecrises muse in the posterior pituitary has direct arterial blood supply, whether the properties muse of its blood (containing hypothalamic releasing and inhibiting forces) via portal blood. Fina legal S.U. and its blood (containing hypothalamic releasing and inhibiting story) and ed. St. Louis. String General SM: The endocrine system, In Berne RM, Levy MN (eds): Physiology, 3rd ed. St. Louis, No. 1, 1987. Mody, 1993, with permission.)

ANTERIOR PITUITARY

- Mast are the hormones of the anterior pituitary? Cheproteins (α-subunits identical; β-subunits confer specificity):

 The subunits confer specificity (α-subunits identical; β-subunits confer specificity): operpoteins (α-esabunits identical; β-subunits confer specificity):

 18bnid-dimulating hormone (TSH; thyrotropin): stimulates thyroid hormone synthesis
 ad release

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 Gonadotropins: LH and follicle-stimulating hormone (FSH) Female: stimulates ovarian function and steroidogenesis Male: stimulates testicular function and steroidogenesis Male: summates
2. Somatomammotropins (single-peptide chain with disulfide bonds):

 Somatomammotropins (single-personate growth (via insulin-like growth factor I [IGF-1])
 GH (somatotropin): stimulates somatic growth (via insulin-like growth factor I [IGF-1]) and is counterregulatory to insulin

 Prolactin (mammotropin): promotes lactation in females Prolactin (mammotropin): prolined.
 Propiomelanocortin (POMC) family (precursor for small peptides produced by pos.

translational processing):

 ACTH: stimulates adrenal growth and steroidogenesis ACTT: Summer of the stablished
 β-Lipotropin, β-endorphin: physiologic roles not firmly established

β-Lipotropin, β-encorpini. physical physica

centration in humans; physiologic roles not established

List the factors (hypophysiotropic hormones) involved in the control of anterior pich.

itary secretion. y secretion.

Corticotropin-releasing hormone (CRH) stimulates POMC synthesis and ACTH se-

cretion. Gonadotropin-releasing hormone (GnRH) stimulates LH and FSH secretion.

Gonadotropin-releasing normone (GHRH) stimulates growth hormone release.

 Growth normalic lettering
 Somatostatin (somatotropin release-inhibiting factor [SRIF]) inhibits growth hormone secretion.

Prolactin-stimulating factor probably exists, but its exact nature has not been resolved

Prolactin-inhibiting factor (dopamine) inhibits the release of prolactin.

Thyrotropin-releasing hormone (TRH) stimulates TSH and prolactin secretion.

16. What is the general model of the control of anterior pituitary hormone secretion? The classic model is represented by the control of ACTH release (see figure on next page). Neural input to the hypothalamus increases or decreases the release of a hypothalamic releasing or inhibiting hormone into the long portal system. This hormone is transported to the anterior pituitary, where it increases or decreases the release of a trophic hormone or hormones. These

trophic hormones enter the systemic circulation and exert effects at target glands. The target gland releases a hormone, which has systemic effects. The target gland limits its own release by exerting negative feedback inhibition at the level of the pituitary gland, hypothalamus, or even input to the hypothalamus. Feedback actions medi-

ated by target gland hormones are called long-loop. Short-loop feedback is the inhibition of hypothalamic function by pituitary trophic hormones. Ultra-short loop feedback is the inhibition of hypothalamic function by hypothalamic factors.

17. Is the control of all anterior pituitary hormones the same? No, each is peculiar in its own way. Sometimes it is easier to remember the exceptions (in

bold) to the general model: CRH-ACTH-cortisol Classic system

Dual (stimulatory [GHRH] and inhibitory GHRH/somatostatin-GH-IGE-I

[somatostatin]) hypophysiotropic hormones Majority of negative feedback of thyroid hormone TRH-TSH-T-/T.

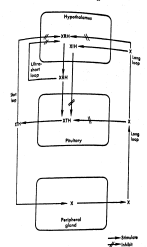
exerted at pituitary (inhibition of TSH)

Two pituitary hormones (parallel system) LH and GnRH-LH/FSH-testes FSH controlled by same hypothalamic factor

Positive feedback of estrogen on LH during GnRH-LH/FSH-ovaries

menstrual cycle Primarily inhibitory hypophysiotropic control Dopamine-prolactin

(dopamine inhibition of prolactin release)



Gasic feedforward and feedback regulation of anterior pituitary hormone secretion. Release (XRH) or its Selective and feedback regulation of anterior pituitary hormone secretare and either stimulate of the selection (XIII) hypothalamic hormones access the anterior pituitary via portal veins and either stimulate of the selection o this fe relate of a pituitary tropic hormone (XTH). XTH then acts at target gland to stimulate peripheral the femiliate of a pituitary tropic hormone (XTH). XTH then acts at target gland to stimulate peripheral the femiliate of a pituitary tropic hormone (XTH). XTH then acts at target gland to stimulate peripheral padiomate (X). X inhibits XTH either directly or by inhibiting XRH or stimulated XIH (long-loop neg-tor feebase. by feedback, XTH may inhibit XTH either directly or by inhibiting XRH or stimulated Art view been been been specified and the stimulate XIH (short-loop negative feedback). It has even been been specified by your statement of the statement of th Record that XRH may inhibit XRH or stimulate XIH (short-loop negative recension). It is shown that the short-loop negative feedback). (From Genuth SM: The cudection is like in the short-loop negative feedback). (From Genuth SM: The cudection is like in the short-loop negative feedback). 1988 a Bene RM, Levy MN (eds): Physiology, 3rd ed. St. Louis, Mosby, 1993, with permission.)

A Defre hypopituitarism.

[&]quot;are appoplinaterism.

All the decreases in anterior pituitary function (although posterior pituitary func
and be decreased). len can be decreased).

What is meant by isolated hypoprimin som.

(An), one or two anterior hormouses are absent. Examples of these are GH deficiency (dwarfing the control of the (July one or two anterfor from was a Grand one or two anterfor from the control of the control

28. (The an example of overactivity (hyperfunction) of an anterior pituitary hormon-Give an example of overactivity (ny per tour rolls) in can lead to hyperproductions. Thinks of the lactotrophs, which synthesize prolactin, can lead to hyperproductions. The state of the lactotrophs and lead to hypogonadism in males and amenorrhea in forms. The Tunivis of the lactotropins, who is a suppression of the lactotropins, who is a suppress LH-FSH roleuse and lead to hypogonadism in males and amenorrhea in females.

POSTERIOR PITUITARY—VASOPRESSIN

21. What is arginine vasonressin (AVP)?

AVP is a neuronormous symmetry of the creates a ring and tail structure. Its structure.

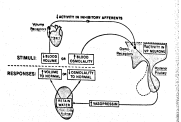
22. Why is AVP also called antidiuretic hormone?

Why is A 1 7 aiso cannot all the two names describe both of its major effects. It is a potent vasoconstrictor, hence vas presslu. At even lower plasma concentrations, it increases passive water reabsorption in the presslu. pressin. At even tower passens can antidiuretic because its effect leads to a concentrated urise

23. Are there any other prominent actions of AVP?

Are there any other production the central nervous system to improve memory, may be AVP appears to nave the state of the state o (a hypophysiotropic effect).

24. Describe the control of AVP. (See figure.)



The two major vasopressin control loops. An increase in the osmolality of the blood stimulates vasopressin release, which increases renal passive water reabsorption (retaining the plasma solvent). A decrease in blod volume stimulates vasopressin release via low-pressure baroreceptors in the heart, which increase plasma volume by increasing renal water reabsorption. Although not shown, an increase in osmolality also increase thirst, thereby uscreasing solvent intake. (From Hedge GA, Colby HD, Goodman RL: Clinical Endocrine Physiology. Philadelphia, W.B. Saunders, 1987, with permission.)

the state of the s to the control loops An true case, in compositive is sensed by osmoreceptors located in the control of a blood-brain burrier. This allows these neural reasons and changes in plantam compositive judgment plantam cofficient. A control of the contro the ambient hypaninamous around in nonodi-brain burier. This bits the cutter in the ambient hypaninamous the cutter is a week small changes in plasma consolidity (usually plasma sodium). A signal from a week small changes in plasma sodium). A signal from a week small changes in plasma sodium, a signal from a week small change in the signal from a secretify or has input to the magnicular vascopersis neurons located primoral. amous the case of the magnocellular vasopressin neurons located primarily in the magnocellular vasopressing of the magnocellular vasopressing of the magnocellular vasopressing vasors of the magnocellular va A signal from a super-controllar nuclei (PVN) of the hypothalamus, Three neurons occard primarily in the super-controllar nuclei (IPVN) of the hypothalamus, Three neurons of the control point of the and the state of t The neurons of the posterior pituitary capillaries, which drain into the systemic croculation, when the posterior pituitary capillaries, which drain into the systemic croculation, which are in the posterior pituitary capillaries, which drain into the systemic croculation. where the posterior pourous empiriors, which drain into the systemic circulation, which was a star reabsorption in the kidney, helping to dilute the increase in plasma through the force and create new water; it just prevents the loss of water from the first plasma. where passive water reasonatement is not souncy, helping to dilute the increase in plasma.

The many half passes not create new water, it just prevents the loss of water from the kidney, and the increases in osmoreceptor activity also stimulate thirst many and an increases in osmoreceptor activity also stimulate thirst ALP dies are considered increases in osmoreceptor activity also stimulate thirst.

Amendment's stimuti:

Amendment's stimuti: Bood volume common as a decrease in end-diastolic volume/pressure/wall stretch. In-banes (Placs in the heart as a decrease in end-diastolic volume/pressure/wall stretch. Inperceptures in the seafferent receptors to the hypothalamus (via a decrease in activity of inper from these afferent results in an increase in vasopressin, which increase pa from more arrowalts in an increase in vasopressin, which increases water reabsorp-

now, and a relative increase in plasma volume. not, and a resource of the protection (via barroeceptors), hypercapnia (via central and peripheral white a hypercapnia (via arterial chemoreceptors), and hypercapnia (via arterial chemoreceptors). (Abors: Arteria in processing via arterial chemoreceptors), pain (via nociception), and nau-agenty-process, hypoxia (via arterial chemoreceptors), pain (via nociception), and nause all increase the release of vasopressin.

& Deire diabetes insipidus. Debre distriction in the property of the prope Public (spans)

The control polyman hypo-estimatic urine) caused by either the lack of AVP or the inability of the processor (processor). It is therefore a nonosmotic diuresis (as opposed to diabetes mellitus).

Make are the types of diabetes insipidus?

name in the constraint in the the novablesize or release AVP. This results in an inability to concentrate the urine. The loss executed teals to an increase in plasma osmolality. The hyperosmolality cannot increase AVP afficient but usually results in a large increase in thirst. In many patients with central diabetes seeks, the patient has high water intake and output but can usually maintain a relatively norne plenu concellin (normonatremia). It is only when water intake is restricted that the severe byennolality becomes apparent.

Nephrogenic (renal) diabetes insipidus is due to the inability of the kidney to respond to serresia Hyperosmolality (hypernatremia) also ensues, and vasopressin is elevated, but the takes cannot respond appropriately.

7. hthere a disease of vasopressin excess?

The syndrome of inappropriate antidiuretic hormone (SIADH) is the overproduction of September of mappingpriate anticonfrection money for the stimuli to vasopressin (e.g., assea) The overproduction of vasopressin (usually from a pulmonary neoplasm) results in case are reassorption, an expansion of plasma volume (hypo-osmolality), and hyponatremia la piera sodium).

ADRENAL GLAND

- La Decribe the functional zonation of the adrenal gland. he arenal gland is composed of layers:
 - The common layer is the capsule.
 - To set here is the capsule.

 The set here is the adrenal cortex, which constitutes approximately 90% of the mass of the adrenal cortex, which constitutes approximately 90% of the mass of the sales.
 - a simul gland and synthesizes steroid hormones. The controlled primary land and synthesizes steroid hormones.

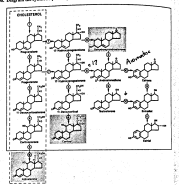
 The controlled primary layer, the core of the adrenal gland, is the medulla, which is controlled primary, by h. by the autonomic nervous system and secretes catecholamines.

ADRENAL CORTEX

29. Describe the histology of the adrenal cortex.

29. Describe the histology or we fractional zountion. The outermost zone is the zona glomman This is a classic securities are the immensionational adulatorems. Next is not zona fraction, which symbiatrus under such earlier securities and the place control of the properties of the place control of the zona factor and the place control of the place control of the zona factor described the zona factor des

30. Diagram the synthetic pathway for the adrenal steroids,



Steroldogeniic pathways in the zona glomerulosa (dotted lines) that produce aldosterone and the zona fascicularreticularis (solid lines) that produce cortisol and adrenal androgens. Major secretory products are shaded. (First Hedge GA, Colby HD, Goodman RL: Clinical Endocrine Physiology. Philadelphia, W.B. Saunders, 1987, which permission.)

Enzymes (abbreviation/gene name) keyed by number to the figure above:

1. Side-chain cleavage (P450scc/CYP1/AI), Rate-limiting step is cholesterol transport in
the mitochondria

3β-Hydroxysteroid dehydrogenase (3β-HSD/HSD3B)

- 3 21-Hydroxylase (P450c21/CYP21) 3. 21-Hydroxylase (P450c11B/CYP11B1) in zona fasciculata (solid box)
 4. 11B-Hydroxylase (P450c11AS/CYP11B2) in the control of 1B-Hydroxylase (P450c11AS/CYP1/B2) in the zona glomerulosa (doned box).
 18-Hydroxylase (P450c11AS/CYP1/B2) in the zona glomerulosa (doned box).
 4-Hostafareticularis does not produce aldosterone under normal
- A Aldostrone symmetry

 A Aldostrone symmetry

 A Aldostrone symmetry

 A Aldostrone under normal conditions,

 The total fasticulation (P450e17/CYP17) in the zona fasticulation

 The Hydroxylase (P450e17/CYP17) in the zon fasciculata/retrocker (P450c17/CYP17) in the zona fasciculata and reticularis only (zona 1/1α-Hydroxylase (P450c17/CYP17) in the zona fasciculata and reticularis only (zona 1/1α-Hydroxylase) gamentost does not produce cortisol)
- stelloss does not produce (P450c17/CYP17). Steps 7 and 8 are catalyzed by same enzyme. Step 8 is 8 17.0 Lysse (P450c17/CYP17). 8 17.20 Lyase (1.2012), Steps 7 and 8 are cataly; Steps 7 and 8 are cataly; Steps 17 are cataly; Steps 17 are cataly; Steps 17 are cataly; Steps 17 are cata ind for steroic and estrogen and estrogen (17-Hydroxysteroid dehydrogenase (17-HSD)

9. 11-11/2007/Julian deliyuroge 10. Aromatase (P450aro/CYP19)

Il. 16a Hydroxylase

1). What is the primary controller of cortisol synthesis?

What is the primary gland increases the synthesis of cortisol acutely and maintains ACTH from the pituitary gland increases the synthesis of cortisol acutely and maintains ACH from use processing and function chronically. ACH binds to a specific cell surface receptor, abspectficial size and control of the state skich via a guanture museus and the mitter benefits a state of the sta less to an increase in cho-less to an increase in cho-less transport from the cytosol into the mitochondria, where the first enzyme-side chain cleav-less transport from the cytosol into the mitochondria, where the first enzyme-side chain cleavknerd massport from use cylinder the cate-limiting step of steroidogenesis is cholesterol trans-ing postect is located. Therefore, the rate-limiting step of steroidogenesis is cholesterol transpot into the mitochondria.

2. What is the primary controller of aldosterone synthesis?

Wiff in the product of aldosterone synthesis involves multiple stimulatory and inhibitory secreta-The common or annual results of the common o protect Conscioury, are good and glomerulosa cell, which, via a G protein, activates phospholi-BEC. Phospholipase C catalyzes the production of the second messengers IP₃ and DAG, which per interpolation of the control of

on into the mitochondria. 33. Are cortisol and aldosterone the most potent glucocorticoid and mineralocorticoid? They are the most potent endogenous steroids of their class. There are several more potent seatetic seroids, such as the glucocorticoids dexamethasone, prednisone, and triamcinotone and te riseratocoticosid 9α-fluorocortisol. Furthermore, some intermediates of endogenous steroidoguesis have biologic activity, such as corticosterone (both glucocorticoid and mineralocorticoid kish) and decaycorticosterone (mostly mineralocorticoid activity). The latter can cause hy-

pression in P450c11B deficiency. 34. How are adrenal steroids transported in the blood?

Seruids circulate in the free (dissolved) form and bind to carrier proteins. The free and bound bens stroid compartments are in equilibrium. In the case of cortisol, about 95% circulates in the board form primarily to corticosteroid-binding globulin (CBG), a high-affinity, low-capacity The and albumin, a low-affinity, high-capacity carrier. The free form is biologically active adis available for metabolism.

E. List the physiologic effects of cortisol. Central nervous system

Lugs

Kiency

Suppresses CRH and AVP Increases food intake Cartiovascular system

Maintains ability to respond to vasoconstrictors Increases gluconeogenesis (glucose synthesis) Necessary for lung maturation and surfactant production in the fetus Pertary

Inhibits ACTH synthesis and secretion Increases glomerular filtration rate

210

Increases resorption/decreases formation Bone Muscle

Increases protein catabolism (increase in gluconcogenic precursors) Decreases insulin sensitivity (decrease in glucose

Immunosuppressive (pharmacologie?)

Immune system

Decreases fibroblast activity and collagen synthesis Connective tissue

Connective tissue

It is well known that cortisol deficiency is a lethal disorder that must be treated promptly. The It is well known that cortisol detections and though it is presumed that the ability to maintain exact biologic reason for this is not known, although it is presumed that the ability to maintain exact biologic reason for this is not known factor. Some of the effects above as product to maintain the main factor. exact biologic reason for this is not known. Some of the effects above are probably relevant

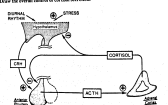
only when the hormone is used in pharmacologic doses.

36. Why is cortisol called a glucocorticold? Why is cortisor came a government of the long-term effects of cortisol is to increase blood glucose (hyperglycemia). It does One of the long-term entered to the control leads to an increase in glucose production in the liver (ghthis in two general ways: First, cortisol leads to an increase in glucose production in the liver (ghthis in two general ways; First, courson seeds to m muscle and glycerol from fat as gluconcogenic concogenesis). The liver uses amino acids from muscle and glycerol from fat as gluconcogenic coneogenesis). The liver uses mining assess that the prevention of the second control prevents insulin-mediated gla-precursors, so, in that sense, cortisol is catabolic. Second, cortisol prevents insulin-mediated glaprecursors, so, in that sense, cottuse is a superior from leaving the plasma compartment. The cose uptake in muscle and fat, which prevents glucose from leaving the plasma compartment. The cose uptake in muscre and tal. The combination of increased glucose uptake leads combination of increased gluconeogenesis and decreased insulin-stimulated glucose uptake leads combination of increased graceine-general to be an important mechanism in maintaining plasma gluose to hyperglycemia. This is thought to be an important mechanism in maintaining plasma gluose levels during a prolonged fast.

37. How is ACTH synthesized?

ACTH is synthesized in pituitary corticotrophs as part of a large precursor molecule, POMC Posttranslational processing of POMC produces big (22 kilodaltons) ACTH, from which ACTH rosmansiational processing the precursor for β-LPH, which is further cleaved to γ-LPH and βand orphin. ACTH contains within it the MSH sequence; hence, when plasma ACTH levels are high (e.g., primary adrenal insufficiency, Nelson's syndrome), skin darkening can occur.

38. Draw the overall control of cortisol secretion.



Regulation of the hypothalamic-pituitary-adrenal (HPA) axis. + indicates that stress stimulates CRH that CRH stimulates ACTH, and that ACTH stimulates cortisol (feedforward control). — indicates that stress stimulates that oction in higher CRH and ACTH and that ACTH stimulates cortisol (feedforward control). — indicates that oction in higher CRH and ACTH and ACTH are action as a feed and act and act are action as a feed and act are actions as a feed and act are actions. hibits CRH and ACTH release (negative feedback). (From Hedge GA, Colby HD, Goodman RL: Clinical En-docrine Physiology, 19:14 docrine Physiology. Philadelphia, W.B. Saunders, 1987, with permission.)

p. beer the four main elements of the HPA axis, perfibr the four man recommend to the hypothalamust The parvocal ular CRH neurons are located priper input lato the try post-manuscript in parvocalidar CRH neurons are located pri-lated and para-versives are more precise input from a variety of sources, in-terior and principal para-versives, and principal prin in mobile plane recommendation in the precise input from a variety of sunce, is as a series of the property and the property while the current of the brain stem (hypotension, hypoxia), and the brain stem (hypotension, hypoxia), where we will release the brain stem (hypotension, hypoxia), when we have the brain stem (hypotension).

or modifiation in the second nature (or protestion, hypoxia).

The modifiation of the portal circulation stimulates ACTH release,

The analysis of the pittitary into the systemic state. **Rel released from the pituitary into the systemic circulation acutely stimulates conti-A TH released from me primony into me systemic circulation acutely stimulates con-tained release, Lang-term elevations in ACTH cause adread hypertrophy, Convenely, and the contained of the conductive defends in a contained to the conductive department of the conductive department of the conductive defends on the conductive department of the conductive departmen and precion. Languerus en ventours in AC 1H cause adreaal hypertropy. Consensely, whether an adversarial of ACTH (accordany adrenal insufficiency, corticosteroid therapy) leads the first and the contract of the contract

s and sample.

A continued from the adrenal gland exerts a variety of systemic serious freedback: Cortisol released from the adrenal gland exerts a variety of systemic serious freedback. Cortisol release by inhibiting the ACTH sensitives as Cart.

A continued from the continued ** Negative feedback a state to the appending the ACTH sensitivity to CRH (at the pidet a maintain it, its interest is own release by inhibiting the ACTH sensitivity to CRH (at the pidet a maintain it. its interest (at the hypothalarmas), and inhibiting insurface at

ACTH sensitivity to CRH (at the pidet a maintain its interest (at the hypothalarmas), and inhibiting insurface at

The property of the pidet a maintain its interest (at the hypothalarmas), and inhibiting insurface at

The property of the pidet a maintain its interest (at the pidet a ma addition. It mans a contract of a minimum and ACH sensitivity to CRI (at the judgest of the hypothalamus), and inhibiting input into the hypothalamus). more & via the limbic system).

& Bearibe the circudian variation in cortisol. Deethe the circumsus various and the state of the control of the control of the control peaks is not because the control of the control peaks is not because the control of In most humans were a summer of the control of the as some \$2.4.M. and as at its coveral an around mounting to the increase in corticol early in the grant was to the covernight fast during sleeping. This pattern is shifted in humans until my be partly they to the covernight fast during sleeping. This pattern is shifted in humans until my be partly they have been also such as the partly of the partly in the partly negist may be purely use. Or me, or m

(name classify the summing one car a name).

The grant term used to describe these stimuli is stress. Stress is difficult to define but can
be grantly term used to describe these stimuli is 4. Can one classify the stimuli to the HPA axis? The grantatern never to construct used attention of suress, ourses as directing define but can partly be drived into two entegories: neurogenic (e.g., anxiety, pain, psychological disturbance). mently be decreased and the company of the company of the configuration bacet and systems: (1937-1931) on project Science, 1937-1931, in new categories are are indeed the predict the magnitude of the ACTH response to specific stimuli, indeed the control of the second of the second

© Outline the general disorders of the HPA axis.

- A Ademocortical insufficiency (not enough cortisol) 1. Primary thass of adrenal function; e.g., Addison's disease) t. running those of auromat auromatic ages Countries a second of ACTH).

 2. Secondary (atrophy of adrenal gland as a result of long-term suppression of ACTH).
- B. Coshing's syndrome (glucocorticoid excess) 1. ACTH-dependent (ACTH induces hypertrophy of adrenal gland)
 - Cushing's disease (pituitary source of ACTH usually from a microadeaoma) b. Eropic ACTH syndrome (nonpituitary source of ACTH; usually from a neoplasm)
- Adressi (autonomous secretion from adrenal adenoma or carcinoma) 2. ACTH-independent
- b. lamgenic/factitious (pharmacologic glucocorticoid therapy)
- C. Adenxonical enzyme deficiencies—congenital adrenal hyperplasia (CAH) 1. 21-Hydroxylase (virilizing; salt wasting)
 - 2. 11β-Hydroxyluse (hypertension)
- 3. 38-HSD (salt wasting)
- 4 17a-Hydroxylase (hypertension)

Q What are the most common symptoms of primary adrenal insufficiency? Reditess

Weight loss Hyperpigmentation Fangue Anoretia Hypotension

history adrenal insufficiency?

All the destruction of by the series of had is the cause of adrenal insufficiency? many adread insufficiency is usually caused by an autoimmune destruction of the adread gland. Secondary adread insufficiency is usually caused by hypopiates of the adread gland. Secondary adrenal insufficiency is usually tuitarism. Abrupt withdrawal of long-term exogenous glucocorricoid therapy also leads to tuitarism. Abrupt withdrawal of iong-term on the HPA axis (negative feedback), ondary adrenal insufficiency because of suppression of the HPA axis (negative feedback), 45. How is the diagnosis of adrenal insufficiency made?

How is the diagnosis of adrenar insurance, and the patient has primary advants. If suspected, a rapid ACTH (cosyntropin) test is performed. If the patient has primary advants are performed in the patient has standard to the patient has primary advantaged to the patient has primary ad 45. How I suspected, a rapid ACTH (cosyntropm) years as personnel at me patient has primary almost insufficiency, the cortisol response to exogenous ACTH is low. If the patient has significant submitted insufficiency, the cortisol response to exogenous ACTH is low because of the patient of the patient has significant and the patient has significant and the patient has significant and the patient has primary almost insufficiency, the cortisol response to exogenous ACTH is low because of the patient has primary almost insufficiency. Il suspension in mustificiency, the cortisol response to exogenous ACTH is low because of advantage on any advantage of the cortisol response to exogenous ACTH is low because of advantage of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to exogenous ACTH. To differentiate between principles of the cortisol response to the cortis insufficiency, the cortisol response to congruence of the insufficiency of the control of the co strophy owing to long-term loss of tropte action of ACTH is usually sufficiently measurement of plasma ACTH is usually sufficient (ACTH et accordary).

ted in primary; low or normal in december that ACTH can be within the normal (reference) range in secondary adrenal itsus. The fact that ACTH can be wanted to that has implications in other consequences of by ficiency is an extremely important concept that has implications in other consequences of by the consequence of byte consequence of the c ficiency is an extremely important concept and population of the p popituitarism (e.g., hypogonaootropu: nypogonaootropu: nypogonaootropu: nypogonaootropu: nypogonaootropu: nypogonaootropu: ny pogonaootropu: ny pogonaootrop to think about it is that it corrusts means that it is inappropriately low for the low certified and fact that the ACTH is not elevated means that it is inappropriately low for the low certified and

46. What are the general symptoms of Cushing's syndrome (glucocorticoid excess)?

- · Facial plethora (red cheeks) and moon face
- Hirsutism
- Hypertension (owing to mineralocorticoid action of cortisol)
- · Myopathy (muscle weakness)
- Myopathy (musics woodness)
 Striae (purple stripes on the abdomen because of skin thinning and stretching and easy · Psychological symptoms (usually depression)

47. How does one screen patients to make the diagnosis of spontaneous Cushing's syn-

drome, and distinguish between ACTH-dependent and independent Cushing's? One or more of the following is usually found in patients with any form of Cushing's syndrome. A 24-hour collection of urine for cortisol is elevated (index of adrenal secretion of cortisol).

- Bedtime salivary cortisol is elevated (due to loss of circadian rhythm; salivary cortisol reflects free [bioactive] plasma cortisol).
- A low dose of dexamethasone given at bedtime indicates that plasma cortisol not fully sunpressed in Cushing's syndrome (test of negative feedback).

To distinguish ACTH-dependent from ACTH-independent Cushing's syndrome, the measurment of plasma ACTH by immunometric assay is usually sufficient. It is low in ACTH-independent Cushing's syndrome (because of cortisol feedback on a normal pituitary) and within or above the normal range in ACTH-dependent Cushing's syndrome. The logic here is similar to that for the normal ACTH in secondary adrenal insufficiency. Pituitary adenomas used to be normal corticorous and retain some responsiveness (albeit diminished) to glucocorticoid feedback. Therefore, although within the normal range, ACTH is inappropriately elevated for the increase in cortisol.

48. Is there a simple way to distinguish between Cushing's disease (pituitary) and ectopic ACTH?

Sometimes it is obvious (big pituitary tumor by magnetic resonance imaging or a lung tumor of radiograph). Occult (radiologically hidden) pituitary and ectopic ACTH-secreting tumors, however. are common. Biochemical testing (e.g., different doses of dexamethasone) is notoriously inaccurate. The only method with sufficient precision involves the measurement of ACTH in the venous outflow from the pituitary (i.e., in the petrosal sinuses) in response to stimulation with exogenous CRH.

49. What is the logic behind the dexamethasone suppression test?

This test was originally designed to diagnose Cushing's syndrome (hypercortisolism). The logic is that a corticotroph adenoma, although arising from a normal corticotroph cell and ex213

and the process of the process the photocorticol recuprior, unto now semantivity to contisol negative feedback. Therefore, and the photocorticol recuprior and the photocorticol release in accordance of the methodoxone (e.g., of 1,000 release in patients with any form of Cushino's management of the photocorticol the first substance of the suppress of the sup and the suppress various excess in patients with any form of Cushing's yndrome.

A suppress of the suppress of See ACTH Sectioning Processing autonomas express sufficient see with low doses of dexamethasone, see approximate the second section of the second section section seems and section se

of the state of th the also been managed our contribute primiting from ectopic ACTHthe set loss a syndrome. A higher dose of decamenhasone (e.g., 8 mg at bedime) is used. The
street colorie is syndrome as the colorie is diseased, because they are
act III secreting primitary adenorass (Caching's diseased, because they are
act III secreting primitary adenorass (Caching's diseased). Cubing 5 Synatrum: A rugue was on our cameransone (e.g., 8 mg at bedinis) used. The second Cubing Synatrum: A rugue was on our cameranson (e.g., 8 mg at bedinis) used. The second Cubing Synatrum: A rugue was one of the second secon The ACTH secreting primary y meananes to usuing a disease), because they store from normal primary to the ACTH secreting primary y meananes (to usuing a disease), because they store from normal primary to the ACTH secretion if mough the ACTH secretion is a secretion in the ACTH secretion if mough the ACTH secretion is a secretion in the ACTH secretion in the ACTH secretion is a secretion in the ACTH secretion in the ACTH secretion is a secretion in the ACTH secretion in the ACTH secretion is a secretion in the ACTH secretion in the ACTH secretion is a secretion in the ACTH secretion in the ACTH secretion is a secretion in the ACTH sec in a superior service service services and services are services and services and services and services are services are services and services are services and services are services and services are services are services and services are s subsection is used. In contact, coups, summar, which did not arise from poliutary correctorages, supports placeoriscoid receptors linked to ACTH secretion. This method also backs precision, and the support of the sup and the secretion with detaments one. The secretion with high-dose detaments and secretion with the secretion with high-dose detaments one, and secretion with detaments one. Therefore a secretion with detaments one. Therefore a secretion with detaments one. Therefore a secretion with detaments one. processor all printery occurs outputs approach row. In secretion with high-dose decumelassor, and secretion with decumelassor. Therefore, although still widely green and secretion with decumelassor. Therefore, although still widely green and secretion with decumelassor. Therefore, although still widely green and secretion with the secretion of the secretion with decumelassor. Therefore, although still widely green and secretion with the secretion with decumelassor. Therefore, although still widely green and secretion with the secretion with decumelassor. Therefore, although still widely green and secretion with the secretion with th suppression test lacks sensitivity and specificity and is used with caution, and the transference suppression test lacks sensitivity and specificity and is used with caution, and the transference suppression test lacks sensitivity and specificity and is used with caution.

a tital is congenital adrenal hyperplasia? What is congenitual and some of the congenitude of Congrainal sufferman experiments as consequently an instancent (usually inherited) in a gene for a consequently experiment of the securiogenic pathway, an instance, carry adversal cannot synthesize adequate control used to be the confidence of th grakesis entyme reasung to a terror canoning pairs in a step of the steroidogenic pathway, in a step of the steroidogenic pathway and the step of the steroidogenic pathway in a step of the steroidogenic path bright, the letta agreeme common agreement and proposed to the state of the state o group of the enzymes before the enzyme step that is blocked.

g. Describe the consequences of the most common enzyme deficiency, 21-hydroxylase

anna... Rezist 17-OH-progesterone cannot be converted to 11-deoxycortisol (cortisol pathway) ed precisions cannot be converted to 11-deoxycorticosterone (aldosterone pathway), both sequentials and adoptions are deficient. The elevation of ACTH increases production of the preasses, which can be converted to androgens. The excess androgens cause virilization in girls ad as lead to ambiguous genitalia in XX fetuses (not sure if phenotype is girl or boy). These dilences be salt wasting because of a deficiency in mineralocorticoid production.

ይ ዝነ do some inherited enzyme deficiencies cause salt retention and hypertension?

The best example is 11β-hydroxylase deficiency. Because cortisol synthesis is impaired, ATH is elevated, which drives steroidogenesis and increases production of the precursor to corist, II-deoxycortisol, and the precursor to corticosterone, 11-deoxycorticosterone, Although a with mineralocorticoid than aldosterone, 11-deoxycorticosterone has sufficient mineralocortirad when elevated to increase renal sodium reabsorption and cause hypertension.

A list the major controllers of aldosterone secretion.

Inhibitory Scientistory Plasma sodium

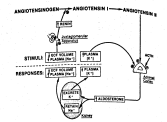
Ang II ACTH (chronically) ACTH (acutely) Atrial natriuretic peptide Places pocassium

A Describe the control of plasma Ang II concentration.

Removed of plasma Ang II concentration.

Removed of plasma sodium and a decrease from the kidney is stimulated by a decrease in plasma sodium and a decrease of ammetes from the kidney is stimulated by a decrease in plasma socious and a function of the cleavage of the line and blood pressure. Renin catalyzes the cleavage of the line and blood pressure. Renin catalyzes the circumstate of the line and blood pressure. he il from the substrate angiotensinogen (see figure, top of next page). Angiotensin I is conend in Aug II by the angiotensin-converting enzyme (ACE). Ang II also directly inhibits renin energy in by the angiotensin-conversing and shown in figure).

Rem don aldosterone help to prevent increases in plasma potassium (hyperkalemia)?
 Account of the advantage of the account of the acc the active at plasma potassium directly stimulates aldusterone synthesis and secretion from the little of the state of the a decree in planner need to prevent increases in processor in processor and secretion of access in planner potassium directly stimulates addusterone synthesis and secretion of the processor in addusterone stimulates renal potassium exerction. push instant brancaman



Highly is inclined version of the mein-inspirentini-aldotencero system. + indicates that reash increases here were not of registerations to support and in and pattern pressure man Age Il directly ordinate advantages reviews. ACPI is a poster aces strainfaiser of aldotencer relates, but this effect was as the several days, we can be a support of the pressure of the pressure from the own the contract of the pressure of the pressure from the own the contract of the pressure of

56. What are the other major renal effects of aldosterone? Aldosterone is called a mineralocorticoid mainly because it increases reabsorption of sodium

in the kidney. Therefore, when sodium intake is low, renin secretion is increased, which leads to an increase in plasma Ang II, which stimulates aldosterone secretion. Aldosterone increases sodium reabsorption to restore plasma sodium to normal.

57. List the major disorders of aldosterone production.

- Hypoaldosteronism

 Primary (loss of zona glomerulosa function; e.g., Addison's disease)
- Primary (loss of zona glomerulosa function; e.g., Addison's disease)
 Secondary (hyporeninemic hypoaldosteronism; loss of renin secretion from the kidney)
- Hyperaldosteronism

 Primary (hyporeninemic): usually caused by a solitary adrenal adenoma (Conn's syndrome)
- Primary (hyporeninemic): usually caused by a solitary adrenal adenoma (Conn's syndrom or nodular adrenal hyperplasia
- Secondary (hyperreninemic): usually caused by renal artery stenosis such that the perfusion
 pressure in the kidney is decreased and the intrarenal baroreceptor stimulates renin release

pressure in the kidney is decreased and the intrarenal baroreceptor stimulates renn release 58. How does one make the biochemical diagnosis of primary hyperaldosteronism?

An increased ratio of plasma addosterone to plasma renin activity, especially in the present of hypokalemia (which usually decreases aldosterone secretion) with the appropriate clinical symptoms (e.g., hypertension), suggests an autonomous production of aldosterone.

ADRENAL MEDULLA

59. Why is the adrenal medulla analogous to a postganglionic sympathetic neuron?

The adrenal medulla is derived from neuroectoderm, is innervated by preganglionic sympathetic neurons, and synthesizes and releases catecholamines.

best the synthesis of adrenal catecholamines.

Be Describe the synthesis of adrenal mechalia and percibe the syntuccion of the adrenal medulla are called chromaffin cells because they contain the relevant cells convert tyrosine to dihydroxyphenylalunine thyrus. gs. Described in the authors are cutted chromalfin cells because they contain the relevant cells in the relevant cells in the relevant cells for the cells convert tyrosine to dihydroxyphenylatanine (DOPA) by the regardles. These cells converted to dopamine. Depunine is converted to dopamine. Depunine is converted to the representative tyrosine hydroxydase. DOPA is converted to dopamine. Depunine is converted to the converted to These cens constructions to university phenomenature (DOPA) by the rega-sers of primary symmetry through the converted to dopumine. Dopumine is converted to the converted to t series e prosine nyurus, access on conveneu to dopamine. Dopamine is convented to test desprise by the enzyme dopamine-β-hydroxylase. Norepinephrise is convented to engineering by the enzyme dopamine-N-methyltransferase (PNMT). acrpiechene by the Companion organistic-ps-hydroxylase. Nore organistic by phenylethanolamine-N-methyltransferase (PNMT), arthrise by phenylethanolamine-N-methyltransferase (PNMT).

4. Which of these enzymatic steps are regulated? Which of these rus, which is tyrosine hydroxylase, which is inhibited by the products of the The major regulated step is tyrosine hydroxylase, which is inhibited by the products of the land of the results of the products of the land of the rus of the ru The major regulared seek of the major regulared seek of the products of the product inhibition). Although somewhat controversial, it is also thought the PNMT with a dernal medulla is increased by cortisol release from the advance of the products of the p princip (end-product nationals) is increased by cortisol release from the adrenal cortex via a solid, in the adrenal medulla is increased by cortisol release from the adrenal cortex via a solid, in the adrenal gland. gracine action within the adrenal gland.

What are the major effects of catecholamines, and what adrenergic receptor mediates these effects?

of Turget Tissues to Catecholamines

Sportse	Tissues to Catecholamines
RECEPTOR TYPE	
B.	Glycogenolysis, lipolysis, gluconeogenesis
	Lipolysis
	Glycogenolysis
	Decreased insulin secretion
	Increased insulin secretion
	Increased heart rate, increased contractility, increase
P ₁	conduction velocity
~	Vasoconstriction Vasodilation in skeletal muscle arterioles, coronary
	Vasodilation in skeiciai musee are
P2	arteries, and all veins
8-	Relaxation
	Decreased contractility
ν2	Sphincter contraction
•	Sphincter contraction
	Detrusor relaxation
	Contraction
	Relaxation
	Ejaculation, detumescence
	Erection?
	Radial muscle contraction
	Ciliary muscle relaxation
	Stimulation
	Piloerection, sweat production
u	Stimulation RL: Clinical Endocrine Physiology, Philadelphia, W.
	RECEPTOR TYPE β ₂ β ₂ β ₂ β ₂ β ₂ β ₃ β ₄ β ₅ α β ₅ α α β ₇ α α α β ₇ α α α α α

dges GA, Colby HD, Goodman RL: Clinical Endocrine P. Stenders, 1987.

Mat are the primary stimuli to catecholamine secretion? Hypoglycemia

Trauma Illness Hetrorrhage

Hypoxia Cold exposure 64. Is there a disease of the adrenal medulia? Is there a disease of the adrendmentorma, which is a catecholamine-secreting tumor, The best appreciated is presented after a gland but can be extra-adrenal (along the sym-These tumors are usually located within the adrenal gland but can be extra-adrenal (along the sympathetic chain).

Endocrine Physiology

65. What are the most common symptoms of pheochromocytoma? Hypertension Headache Excessive perspiration

Absence of all four of these symptoms virtually excludes pheochromocytoma. THYROID PHYSIOLOGY

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Palpitations

66. Describe the functional anatomy of the thyroid gland.

 Follicles: formed by cells that synthesize, store (extracellularly), and secrete thyroid hor. mone Colloid: central space in the follicle where thyroid hormone is stored as a component of

thyroglobulin Parafollicular (C) cells: synthesize and secrete the hormone calcitonin

67. What are the main thyroid hormones? T_s (3,5,3',5'-tetraiodo-L-thyronine) is the main secretory product of the thyroid gland.

 T₁ (3,5,3'-triiodo-L-thyronine) can also be produced by the thyroid gland. Most T₁ is produced by monodeiodination of T₄ in peripheral tissue including target cells. Because T₄ is

significantly more potent than T4, T4 can be considered a circulating prohormone. Reverse T₁ (3.3',5'-triiodothyronine) is found in the blood, although little if any is secreted

by the thyroid. This hormone is essentially devoid of biologic activity and is produced primarily by peripheral monodeiodination of Ta-

68. What is the source of the iodine used by the thyroid gland to synthesize thyroid hormone?

Organic iodine or inorganic iodide (food supplement) in the diet is absorbed into the blood from the gustrointestinal tract. The follicular cell has an iodide (ionic form of iodine) pump, which

traps iodine within the thyroid gland.

its storage space in the colloid.

69. Describe the synthesis of thyroid hormone.

 Trapping of iodide — iodide [I-] pumped from the plasma to the intracellular compartment. 2. Oxidation and organification of iodide (on colloidal side of follicular cell). This is probably the conversion of I" to Io and is catalyzed by the enzyme thyroperoxidase, Io is highly reac-

tive and binds quickly to the ring of a tyrosyl residue of thyroglobulin (see later). 3. Exocytosis of thyroglobulin, which has been synthesized within the cell, into follicular

lumen.

4. Iodination of tyrosine residues within thyroglobulin. This occurs within the follicular lu-

men and is therefore an extracellular reaction. If one carbon of the tyrosine ring is jodinated, this results in 3-monoiodotyrosine (MIT). If two carbons of the tyrosine ring are iodinated, this results

in 3,5 diiodotyrosine (DIT).

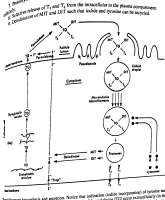
5. Coupling of iodotyrosines (on thyroglobulin molecule) occurs when MIT and DIT come

in contact while still part of the thyroglobulin molecule. If MIT and DIT are coupled, T3 results.

If DIT and DIT are coupled, T, results,

6. Endocytosis of thyroglobulin-containing thyroid hormone. If thyroid hormone is needed systemically, TSH from the pituitary is increased and stimulates recovery of thyroglobulin from 217

policelysis of thyroglobulin. The liberation of T₄ and T₃ from thyroglobulin occurs intraprogly, and T₄ from the intracellular to the plasma compartment,
Selective role are of T₁ and T₄ from the intracellular to the plasma compartment,
Selective role are of MIT and DIT such that include and tyrosine can be



Byrid batteree biosynthesis and secretion. Notice that iodination (iodide incorporation) of tyrosine and Most and storage of thyroid hormone as a component of thyroglobulin (TG) occur extraoellularly (in the blook heres [coloid]). For detailed description of each step, see lext. (From Genuth SM: The endocrine teen in Berne RM, Levy MN (eds): Physiology, 3rd ed. St. Louis, Mosby, 1993, with permission.)

A Why do the thyroid hormones have such a long half-life?

The half-life of T₄ (6 days) and T₃ (1 day) is long primarily because thyroid hormones cirall blook to currier proteins. T_d circulates more than 99,9% bound to thyroid-binding globula fills. is IBO, transhyretin, and albumin. T₁ is slightly less tightly bound (99.7%) and apparently former to the control of the con $k_{0.04}$ is slightly less tightly bound (92.78) and hormone lightly to transity to transity retin. Because little of the total circulating thyroid hormone light that a fac. little is available for metabolism, hence the long half-life.

A list the systemic effects of thyroid hormones. Metabilism: increase basal metabolic rate and oxygen consumption (and therefore in-Class minute ventilation, cardiac output, food intake, carbohydrate metabolism, and heat production)

Growth and maturation: required for normal skeletal growth probably by allowing normal tiffer. nal effects of IGF-1 on bone and normal GH secretion

· Central nervous system: necessary for perinatal maturation and normal reflexes Autonomic nervous system: increase sympathetic activity Temperature regulation: increase thermogenesis 72. How is the circulating thyroid hormone regulated?

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How is the circulating thyroid normous the state of the main feedback loop in this system is T₄-T₃ inhibition of TSH and TSH stimulation of T_C main feedback loop in this system is T₄-T₃ and T₄ inhibit TRH secretion TRH in the system of T_C and T₄ inhibit TRH secretion TRH in the system of T₄ inhibit TRH secretion TRH in the system of T₄ inhibit TRH secretion TRH in the system of T₄ inhibit TRH secretion TRH in the system of T₄ inhibit TRH secretion TRH in the system of TrH The main feedback toop in tuns system = 4 - 3.

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The main feedback toop in tuns the set-point for Ta-Ta negative feedback.

Endocrine Physiology

Hypothalamus TSH (+)

Regulation of the hypothalamic-pituitary axis, + indi. cates that TRH stimulates TSH and that TSH stimulates thyroid hormone release. T4 is converted to the more potent T₂ in the liver and target tissue. - indicates that both

Ts and T4 inhibit TSH release (negative feedback). (From Goodman HM: Basic Medical Endocrinology

2nd ed. New York, Raven Press, 1994, with permission.)

. Thyroid-stimulating immunoglobulins (TSI): These are antibodies produced under abnormal conditions (e.g., Graves' disease) that are directed against TSH receptors but result in activation by mechanisms similar to TSH.

· Thyroid nerves: These may modulate the sensitivity to TSH.

73. Other than TSH, are there other factors that regulate thyroid function?

. Iodine: Although chronic iodine deficiency leads to a decrease in thyroid hormone and a TSH-mediated increase in thyroid size, an increase in iodine can also decrease thyroid hormone secretion by the paradoxical Wolff-Chaikoff effect. This is due to a decrease in the organification of iodide and may be protective against iodine-induced hyperactivity of the thyroid gland. Excess iodine may also decrease the secretion of thyroid hormone possibly by decreasing the sensitivity to TSH.

74. Outline the general disorders of thyroid gland function. A. Hypothyroidism (too little thyroid hormone) 1. Primary

- a. Hashimoto's thyroiditis (autoimmune)
- b. Iodine deficiency
- 2. Secondary-Hypopituitarism
- B. Hyperthyroidism (too much thyroid hormone secretion) 1. Primary (thyrotoxicosis)
 - a. Endogenous (Graves' disease-TSI) b. latrogenic (overuse of exogenous thyroxine)

2. Secondary-TSH secreting tumors (very rare)

Weight gain Constinution

Myxedema

15. What is a goiter? What is a goiter?

What is a goiter?

What is a goiter?

What is an enlargement of the thyroid gland. It can be due to hyperthyroidism (e.g., TSI Apiller is an enlargement by hypothyroidism (e.g., TSI Apiller is a custing decreased T amount of the control of th Mar is an enlargement of the control ground. It can be due to hyperthyroidism (e.g., TSI

A few is said or hypothyroidism (e.g., iodine deficiency causing decreased T, production
in the control of the

what are the symptoms of hypothyroidism? what are any property and any property a Hoarse voice

Cold intolerance Slow reflexes

Cool, dry skin Muscle cramns Coarse hair

7. Define myxedema.

peline mysedemu.

Define mysedemu.

As infiltracion of the skin and subcutaneous tissue with mucopolysaccharides occurs leading As infiltracionoc. usually of the face, hands, and feet. As inhutation or the same associations tissue with the same and same and feet.

Is there a specific concern if hypothyroidism occurs in the neonate?

Is there a specific concern in in prompt common occurs in the neonate?

Is there a specific concern in in prompt, common occurs in the neonate?

You concern and a puffy face with protruding toneue. The month is a concern a puffy face with protruding toneue. Yes, congenital nyponyryomana variationary arount untreated is characterized by dwarfism. Yes, congenital nyponyryomana variation from the preneed reardation, and a purity new man processing surgue. The mental retardation can be pre-regal reardation, and a purity new man special state of the mental period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained when thyroid hormone is administered in the neonatal period (and contin-ted or maintained or maint egid or minimized when unytone monutones a communication in the neonatal period (and contin-egid or minimized when unytone monutons for hypothyroidism (elevated TSH) is the standard or incuptort life). Testing of all newborns for hypothyroidism (elevated TSH) is the standard

R is patient is suspected of having hypothyroidism, is there a simple way to distinguish

ginery thyroid dysfunction from hypopituitarism? sary myrum and the standard of TSH using a third-generation supersensitive assay reliably distinguishes pri-REPORT TO SH) from secondary (normal or low) hypothyroidism.

8. Why can hypopituitarism lead to hypothyroidism if TSH is in the normal range? As in secondary adrenal insufficiency, this is an important concept. If the hypothalamic-

object-throtroph function were normal, a low circulating T₄ should lead to an elevated TSH. TISEs not devated, the low thyroid hormone is due to hypothalamic-pituitary dysfunction.

II. Her does one assess functional hypothyroidism if most of the circulating thyroid hornote is bound (not biologically active)? Measure free T.

 $\mathbb Z$ is there a common condition that causes a discrepancy between free and total $T_4?$ The most common explanation is a change in circulating TBG concentration. For example,

Exergisary (or with estrogen therapy), TBG is elevated, which increases total T_4 . Because the hydralanic-pituitary system is normal in most pregnant women, once the new binding sites are sarran, free T4 is properly regulated and maintained within the normal range.

& List the symptoms of hyperthyroidism.

 Elevated basal metabolic rate · Heat intolerance

• Warm skin

· Excessive perspiration Weight loss (despite an increase in the intake of food)

· Loss of muscle mass · Hypertension

· Tachycardia (a sympathomimetic effect)

E_{aphthalmos} (protruding eyeballs; occurs in Graves' disease)

84. Is there a simple method to diagnose hyperthyroidism?

Is there a simple method to unquinose ...,

Because TSH-secreting tumors are exceedingly rare, suppressed TSH is used as a screening tumors are exceedingly rare. Because TSH-secreting tumors are exceeding.)

Because TSH-secreting tumors are exceeding.

Because TSH-secreting tumors are exceeding.

Because TSH-secreting tumors are exceeding. the need for TRH testing.

85. What are the treatment options for patients with Graves' disease?

- Surgical removal of thyroid gland (thyroidectomy)
- Radioactive iodine administration (ablation)
- Radioactive former authorized to the state of the state o

86. Summarize the thyroid findings in primary hyperthyroidism, primary hypothyroidism. and pregnancy.

	NORMAL	HYPERTHYROID	HYPOTHYROID	PREGNANT
Total T ₄	N	1	1	1
TBG	N	N N	· N	i i
Free T ₄	N	1	* i 🗜	N
TSH	N		T	N

87. Discuss the thyroid findings in primary hyperthyroidism.

The main defect is excess secretion of T₄ and hence an increase in total and free T₄. TSH is suppressed in primary hyperthyroidism because of negative feedback inhibition by free T.

88. Discuss the thyroid findings in primary hypothyroidism.

The main defect is a failure to produce T₄ normally. Therefore, total and free T₄ are decreased, TSH is increased because of the loss of the negative feedback inhibition by free T.,

89. Discuss the thyroid findings in pregnancy.

The increase in estrogen during pregnancy is probably due to an increase in TBG synthesis in the liver. (Oral contraceptives can cause a similar effect.) Total Ta is increased because of an increase in the number of available binding sites on TBG. Assuming normal thyroid and pituitary function, free T, and TSH levels are regulated and maintained within the normal range. Hyperthyroidism may occur during pregnancy and postpartum, and it is extremely important that true endogenous hyperthyroidism be distinguished from a normal elevation of total T, during pregnancy because of an increase in TBG.

ENDOCRINE CONTROL OF GROWTH AND DEVELOPMENT

90. Summarize the hormonal regulation of growth.

Prenatal growth is not well understood. It is thought that insulin or insulin-like factors may be involved because women with increased blood glucose (diabetic hyperglycemia) tend to have larger infants (possibly as a result of fetal hyperinsulinemia). Clearly, other unknown factors influence fe-

tal growth. Hormonal control of growth up to about 1 year of age is also not well understood. Juvenile growth (from age 1 year to puberty) is thought to be influenced by the GH axis (and its intermediates), thyroid hormone, and insulin. Much of the effect of thyroid hormone appears

to be due to its maintenance of normal GH secretion. Puberty is a time of dramatic changes in growth and development. The increase in sex steroid production (androgens in males and estrogens in females) stimulates the pubertal growth sport The major mechanism appears to be sex steroid-induced growth hormone secretion, although

EXERCISE

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April of the epiphyseal (growth) plates of long bones.

April of the epiphyseal (growth) plates of long bones. where factors me amounts. See secrosis also terminate the spin factors of the opiphyseal (growth) plates of long bones.

get light the hormones influencing normal growth, the light the hormones influencing normal growth, the major controlled the light the l pointe the hormones management and growth.

18 still stimulates [GP-1, the major controller of somatic growth.

18 still stimulates in necessary for normal central in the stimulates are still stimulated by the stimulation of the stimulation

Remarks initialists for the first simulates for normal central nervous system development and for the ration of IGF-1; it stimulates GH secretion. Thereto are the stimulates of the secretion. nemetal action of ROT manufacture and terminate pubertal growth spurt. They are necessary for General Secretion (particularly androgens).

Gonzali secretion (particularly androgens). Isulia stimulates fetal and postnatal growth.

Insulla stimulases result of the stimulates and decreasing effects of growth of the stimulates of bone. factors on growth plates of bone.

 Bies GH directly increase growth velocity in children?

 Bies GH directly increase growth velocity in children? Boes GH directly mer uses of the growth-promoting effects of GH are mediated by IGF-The cylinesis and release of IGF-1 from the liver and its local production in GH-target tissues

The sylinesis and release of IGF-1 from the liver and its local production in GH-target tissues

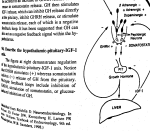
The sylinesis had release of IGF-1 was originally called sulfation factor because it incomes The synthesis and remains the synthesis and production in GH-target tissues as similarity by GH. IGF-1 was originally called sulfation factor because it increased incorporational sulfate into bone. It was then called somatomedia C because it increased incorporation of the synthesis of the synth as straighted by Orn. Co.

It was then called somatomedin C because it mediates the safe chandroxin sulfate into bone. It was then called somatomedin C because it mediates the safe chandroxin G(H). Recause somatomedin C was cabanasa. gire of chandrouin surrant and the second common control of the subsequently found to have sequence that of surrant part of the second control of the seco the set sometime representation of the sequence of the sequenc agrants are growth factor binding proteins (IGFBPs).

SLEEP RHYTHING

21 Describe the control of GH secretion. hypothelamic control of GH secretion involves andstory and inhibitory hypophysiotropic factors. Secrets in the hypothalamus synthesize GHRH, with simulates GH release, and somatostatin, which inhibits GH. These neurons receive inputs for higher brain centers and stress pathways much Birthe HPA axis. An increase in GH secretion can bracebed to an increase in GHRH release or a demuse in somatostatin release. GH then stimulates IEF release, which can inhibit GH release directly ate pinitary, inhibit GHRH release, or stimulate mustain release, each of which is a negative imback loop. It has been suggested that GH can Sout as a negative feedback signal within the hyoodubress

adjust inhibition of GH.



Endocrine Physiology 222

95. List some potential negative feedback loops. GHRH stimulates GH, which inhibits GHRH, which decreases GH.

 GHRH stimulates GH, which stimulates IGF-1, which stimulates somatostatin, which inhibits GH release.

hibits GH release.

Somatostatin decreases, which increases GH release, which increases IGF-1, which stimulates somatostatin release, which decreases GH release.

96. What central and systemic factors are involved in the control of GH release? Inhibition of GH

Stimulation of GH Increase in plasma glucose Decrease in plasma glucose (hyperglycemia)

(hypoglycemia) Increase in plasma free fatty acids Decrease in plasma free fatty acids Cortisol (exogenous or endogenous)

Increase in amino acids (e.g., arginine) Pregnancy Stage 4 (deep) sleep Exercise

97. Place factors involved in the control of GH release into context. GH has direct effects on intermediate metabolism. It is a counterregulatory hormone to insulin

and stimulates glucose production. Therefore, it makes sense that plasma glucose would inhibit GH release because this forms yet another feedback loop: GH stimulates plasma glucose, which inhibite GH release, or a decrease in glucose stimulates GH, which increases plasma glucose.

 Because one of the main effects of GH (via IGF-1) is to increase linear growth, it makes sense that ingestion of the building blocks of protein (amino acids) stimulates GH re-

lease Fasting and prolonged caloric deprivation require mobilization of endogenous fuel stores

to mobilize glucose and fatty acids.

 GH secretion is stimulated during deep sleep. This is extremely important in children and may be one reason why sleep deprivation may lead to short stature. It is well known that endogenous Cushing's syndrome (hypercortisolism) and exogenous

glucocorticoid therapy decrease growth velocity in children. Children receiving high doses of potent glucocorticoids (prednisone therapy) for rheumatoid arthritis or to prevent transplant rejection are often much shorter than predicted.

98. Classify the direct biologic actions of GH.

. In adipose tissue: GH decreases glucose uptake and increases lipolysis, leading to a decrease in adiposity.

. In muscle: GH decreases glucose uptake and increases amino acid uptake and protein

synthesis, leading to an increase in lean body mass. . In the liver: GH increases gluconeogenesis (glucose secretion) and increases IGF-1

release

99. Summarize the direct biologic actions of GH. Most of the direct actions of GH are on intermediate metabolism. GH results in hyperglycemia because of decrease in glucose uptake and increase in glucose production (counteracts effects of insulin). GH also stimulates increase in muscle mass. The net results of these

two effects are a decrease in adiposity and increase in muscle mass. This explains the abuse of GH by bodybuilders and competitive athletes. GH may also directly increase epiphyseal growth, although most of this effect is mediated by local production of growth factors (such

as IGF-1). 100. What are the direct effects of IGF-1? IGF-1 stimulates an increase in organ size and function. For example, it ensures that as a child grows rapidly, the heart, lungs, kidneys, and other structures grow as well. IGF-1 also has dra22 on the chondrocytes in bone; it increases transcription, protein symbasis, conduction, and cell size and number, all of which lead to an increase in lines, and cell size and number and the conduction of the Briefly outline the pattern of linear growth from conception to adulthood.

Fetal growth
Fetal growth

- Peaks at about 4-6 months' gestation 1. Peaks at as high as 12 cm per month
 - B. Juvenile growth
 - Juvenile Brown prenatal peak until about 2 years of age
 - Declines from prenate the providing provided by the providing prov
 - C. Pubertal growth spurt
 - Simulated by at about 10 years of age in girls and 13 years of age in boys
 Usually starts at about 10 years of age in the same family.
 - 2. Island same family variable between subjects even within the same family
 - 3. Variable Detection 12 years of age in girls and 14-15 years of age in boys
 4. Usually peaks at about 12 years of age in girls and 14-15 years of age in boys 4. Usually peaks at account 1991 and 15-16 years of age and most males by 17-18 years

 5. Most females reach adult height by 15-16 years of age and most males by 17-18 years
 - of age
 of age
 for age
 of age
 of age
 of age
 of the pubertal growth spurt caused by gonadal steroid-induced fusion of
 the country plates of the lone banes

11. List the general disorders involving GH.

- GH deficiency:
- GH debreur.

 Hypophuitarism, isolated GH deficiency—leads to short stature in children (dwarfism) and is treated with recombinant human GH injections • Old age - controversial subject
- GH insensitivity—Laron dwarfism (high GH, low IGF-1)
- CH erress—GH-secreting pituitary adenoma:
- · Childhood-gigantism
- Postpubertal—acromegaly (acral enlargement, soft tissue overgrowth, insulin resistance) being to hyperglycemia and hyperinsulinemia)

III. Way does GH excess have two different names depending on the age of onset?

Although the cause is the same (almost always a GH-secreting pituitary adenoma), the physal appearance is quite different. GH excess before puberty leads to greatly increased prowth vekery and a greatly increased final adult height (gigantism). If GH excess commences after pubetal fasion of the epiphyseal plates, linear growth is not restarted and final adult height does not dage: Acromegaly (from the Greek akron ["extremity"] and megas ["large"]) is characterized is connective tissue proliferation.

- IX List some of the features of acromegaly. Soft tissue swelling, particularly in hands and feet
 - Skin thickening
 - Increased sweating

 - Bony changes (cortical thickening, osteophyte proliferation, mandible enlargement leadits to a protrusion of the lower jaw [prognathism]) herve entrapment (owing to bone and connective tissue overgrowth)
 - Organomegaly (large liver and kidneys) · Insulin resistance

its. How does one diagnose GH deficiency?

- Becare of the episodic nature of GH secretion, a single plasma measurement is not particbelow of the epsodic nature of GH secretion, a single plasma measurement is not below the bound of the plasma measurement in the secretion of ***Colal. Usually, some kind of stimulation test is performanted with infusion, or a sleep study to measure GH during stage 4 sleep.

Endocrine Physiology 224 106. How does one diagnose acromegaly?

How does one diagnose acromegany.

Measurement of several elevated plasma IGF-1 levels is probably the most common cur. rent approach. Comparison of photographs from different ages is often helpful.

107. What is the treatment for acromegaly? Pituitary surgery to remove the GH-secreting tumor

 Treatment with somatostatin analogue Radiation therapy of the pituitary

Glucose production

ENDOCRINE PANCREAS

108. Describe the anatomy of the endocrine pancreas. Describe the anatomy of the careful describe digestive enzymes into the gastrointestinal tract)

The pancreas is both an exocrine (secretes digestive enzymes into the gastrointestinal tract)

and an endocrine organ. The endocrine component of the pancreas consists of several million clusters (islets) of cells called the islets of Langerhans.

109. What are the major hormones secreted by the islets and from what cell type? Insulin is secreted by B cells (also known as β cells)—approximately 75% of islet Glucagon is secreted by A cells (also known as α cells)

Somatostatin is secreted by D cells (also known as δ cells)

110. What is the major secretory product of the islets of Langerhans, and how is it synthe-

cized? The protein insulin, the storage hormone, is synthesized as a prohormone called proin-

sulin. Posttranslational cleavage of proinsulin produces insulin and C-peptide (connecting), Al-

though C-peptide has minimal, if any, biologic activity, its measurement is generally used as a marker for islet cell function because it is released with insulin.

111. List the major components of intermediate metabolism under endocrine control.

Glycogenolysis (breakdown of glycogen to glucose)

Gluconeogenesis (synthesis of new glucose from precursors) Glucose consumption Glycolysis (burning of glucose for energy production) Lipogenesis Fat storage

Lipolysis Fot breakdown Ketogenesis (oxidation of fatty acids to ketone bodies) Ketone production 112. Categorize the effects of insulin.

Generally, insulin promotes the storage (anabolic effect) of circulating sugar, amino acids,

and fat and prevents the breakdown (anticatabolic effect) of these stores.

ANABOLIC

ANTICATABOLIC

Effect on liver Increases glycogen storage, synthesis Inhibits glycogenolysis, ketogenesis, of very low-density lipoproteins glucogenesis

(VLDL), glycolysis Increases amino acid uptake and pro-Inhibits glycogen phosphorylase

Effect on muscle tein synthesis and increases glucose transport and glycogen synthesis Effects on fat Increases glucose uptake and tri-

Inhibits lipolysis glyceride storage

113. In question 112, why was hepatic glycolysis (glucose consumption) listed as anabolic? The main effect of insulin within the hepatocyte is to increase glucose uptake and then to store it as efficiently as possible. Therefore, one of the goals of insulin is to maintain free intraand the photose concentration and inspiratory is low as possible. If glucose cannot be glucose cannot be glucose cannot be glucose (glycolysis), used with process is linked to glucose uptake, it is considered anabolic. estant ended to glucose uptake, it is considered anabolic.

Because glices had to pick one primary effect of insulin, what would it be?

If the had to pick one primary effect of insulin, what would it be?

If the had in increases glucose uptake and leads to a decrease in bin. ff sare had to pick one promise. The same state and leads to a decrease in blood glacose (plasma glucose), tradin increases glucose uptake and leads to a decrease in blood glacose (plasma glucose). Ill. How is the release of insulin controlled?

How is the directly stimulate insulin release: Figures must under a glucose, amino acids, fatty acids, ketones)
Fixed metabolites (glucose, amino acids, fatty acids, ketones)

Food memoranes (increase sensitivity of B cell to glucose)

Gistrointestinal hormones (increase sensitivity of B cell to glucose)

• Glucation • ACCOUNT OF THE ACCO Factors that induced y more continuous (cortisol and GH induce peripheral resistance to insulin a Counteregulatory hormones (which arimulate in the counteregulatory) in blood almones (which arimulate in the counteregulatory).

Counterrogument, control glucose [which stimulates insulin release]) Factors that inhibit insulin release:

Figures that the constant of t Catecholamines (epinephrine and norepinephrine)

16. Describe the effects of glucagon. DESCRIPTION OPPOSES (counterregulates) insulin and is therefore catabolic. The main role of

Gaccagus upposed to the state of the state o gacagon are promotes the conversion of circulating free fatty acids to ketoacids.

117. Are the factors that regulate glucagon secretion basically the opposite of those that regsiste insulin?

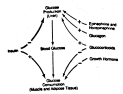
Yes with notable exceptions. Glucose, ketones, and free fatty acids all inhibit glucagon rebut, as one would expect. Amino acids, however, actually stimulate glucagon release. A way to member this is to consider a carnivore in the wild. The hyena can ingest up to 30% of its body negs when it eats, for example, a zebra. This represents a huge protein (and potassium) load. is the metabolites of protein digestion (amino acids) are absorbed (without concomitant glucose abzepise), massive insulin release would lead to hypoglycemia and, possibly, loss of con-Kinzaess. Therefore, amino acid stimulation of glucagon makes sense to counteract the hypo-Plantic effect of insulin when consuming a protein meal. Glucagon also increases amino acid मार्थित in the liver (for gluconeogenesis), so it makes sense that glucagon is stimulated by amino

IR. What is the derivation of the name somatostatin as it refers to an islet cell hormone? The 14-mino-acid peptide somatostatin was first identified as a neurohormone in the hy-

Medianes that inhibits GH (somatotropin) release—hence the name somatostatin. The identical beinger use imments GH (somatotropin) release—hence the name someone was assequently identified from the D cells of the islets of Langerhans and found to in-Milbon insulin and glucagon release.

Bearibe the hormonal maintenance of blood glucose.

As ear end of the spectrum is a state of total glucose consumption (fed state), and at the of the ctd in a state of total glucose consumption (red security to the sale of total glucose consumption). Insulin and the counterregulatory beautiful as state of total glucose production (fasted state). Insulin and the counterregulatory beautiful as the count tength is a state of total glucose production (fasted state). Insulin and the counterleg-tength regulate the balance between glucose consumption and glucose production. In the fed the media is the balance between glucose consumption and glucose production. In the fed replite the balance between glucose consumption and glucose production. It is insulit is simulated and promotes glucose uptake in muscle and adipose tissue (storage). the folds is simulated and promotes glucose uptake in muscle and adipose usue tester the folds that, insulin is low, allowing catabolism. Furthermore, the counterregulatory both the solid to the counterregulatory both the solid to the soli the plant uptake (decreases insulin sensitivity in muscle and fat) and (2) increases hepatic buyes polymers. those production (eluconeogenesis).



tangemen of the regulatory (until mil and conterregulatory) between his the fasted state (upw.01), institues, which should be joulve production, and the contempregulatory between sea or elevated shorts distinguished to the contempregulatory between the contempregulatory between the contempregulatory between the contempregulatory between miles (consumption). The contempregulatory between miles (consumption) that contempregulatory between miles (consumption) the contempregulatory between miles (consumption) and contempregulatory between miles (consumption) and consumption of the contempregulatory between miles (consumption) and contempregulatory between miles (consumption) and contempregulatory between miles (consumption) and consumption of the contempregulatory and consumption of the contempregulatory and consumption of the consumption of the contempregulatory and consumption of the con

120. What is the insulin-to-glucagon ratio? In the normal individual, this reflects degree of fed versus fasted state. If the individual is

in the feet state, the insulin-to-glucagou ratio is high, which induces anabolic enzymes and inhibits catabolic enzymes. If the individual is in the fasted state, the insulin-to-glucagon ratio is low, and catabolic enzyme activity predominates.

121. Describe the pattern of glucose, glucagon, and insulin during a typical day in a normal person. A mixed meal (carbohydrate, protein, fat) increases glucose. Insulin increases in resonse

which simulates glaces uptake and lowers blood glaces. After some meals, installar levels amaly decline which body glaces is still elevants. This decrease in installar may be due to generica scion of acoustosation within the idet cells and probably prevents too large a decrease in glaces after a mol of acoustosation within the idet cells and probably prevents too large a decrease in glaces after a mol of acoustosation within the idet cells and probably prevents too large a decrease in glaces after a most present to probably the control of a second control of a second control to decrease from its peak after a most low below baseline. The attenuation of the installar response after a most allows places and glaces to normalize granularly without going significantly below bastlevish.

122. What is the flow of fuel during a prolonged fast, and what hormones are responsible? In a prolonged fast, insulin is low, and the counterregulatory hormones are increased. Hepatic

photonogenesis is the prime source of glucose (180 gidsy), which is consumed by the certail see was system (which does not require insulin to maintain glucose grades), blood cells; and, to some center, mucke, back, shape, and other organ. The substrate for bepaire gluconogenesis is synferbyle organic source of the state of the supplying among socials to the liver to be used in glucose synthesis; GH-induced and catecholumeinduced lipids susupplies glycerul for beguite glucoseogenesis and fairly social. There fairly sakes the used by the best, knower, and muscle for for the and are also converted to decrease in the little.

123. What would be the consequence of either an inability to secrete insulin or an inability to respond properly to circulating insulin?

- Diabetes mellitus (too much sweet urine):

 Type 1: the absence of insulin itself
- Type 1: the absence of insulin itself
 Type 2: a resistance to insulin action

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Corn can result in a failure in glucose uptake leading to hyperglycemia and a glu
Elbe (corn can result in a failure in glucose uptake leading to hyperglycemia and a glurates sexuit in a gradit sexuit in a gradit sexuit (asmotic) diuresis.

or the two forms of diabetes mellitus.

(**Characterine the two forms of diabetes mellitus (T1DM) has also 't was Characterize the two rounes of consecution mentius.

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(hardrefize the two rounes of consecution mentius.) the first measurement is considered in small production of the second of 117 om militars recursion to the B cells of the sists that cornally produce media.

200 of the minimum destruction of the B cells of the sists that cornally produce media.

200 of the minimum destruction of the B cells of the sists that cornally produce media.

200 of the minimum destruction of the breakform continue.

200 of the product of the breakform continue. control de movimente consistence of the control of the islets that accomally produce media, and the moviment of the control of ment of the control o of the second section of the second s surface. This is why TIDM can be considered "tissue staration in the face of plany."

The face of plany. The by space. This is way a live of plants of the property of the face of plants, and the face of plants of the face of plants. The face of plants of the face of the fa idabetes menuna variante anno acult-oract or non-insulin-dependent in the control acult-oract or non-insulin-oract or non-insulin-oracl or non-insu powers memory security are property and is not due to a lack of inposition of the syndrome. NIDDM is currently thought to be an inadequate
the first early in the onset of the syndrome supposed to be an inadequate
the first early in the onset of the syndrome at worst because type 2 diabetes mellions conhave been of conflusing one at worst because type 2 diabetes mellitus can be treated with being a kest of a conflusing one at worst because type 2 diabetes mellitus can be treated with being a kest of a conflusing one at worst because type 2 diabetes mellitus can be treated with being a kest of a conflusion of constant of the conflusion of the conflusi acquite the set of a continuing one or more no considered a syndrome of severe insulin under certain circumstances. Although the pathogenesis of T2DM distribution of considered a syndrome of severe insulin section of the considered a syndrome of severe insulin section. parachagic does of mount in the considered a syndrome of severe insulin resistance. That is, the sometime to be considered a syndrome of severe insulin resistance. That is, the sometime to be considered and the considered of the control of the in the second sec as and glucose uptake is decreased.

(K. Define diabetes mellitus. shows toponse to oral glucose that is unexplained by other factors.

116. Discuss an example of diabetes mellitus as formally defined.



Thou and placese tolerance test in normal and diabetic subjects. The diabetic subject has fasting (0 hour) replaces interactive test in normal and diabetic subjects, the graces subjects insulin, which lowers blood replaces a few normal subject responds to an oral glucose load by increasing insulin, which lowers load to charac-Process the rormal subject responds to an oral glucose load by increasing insulin, usose load is charac-fracts (but sounly not below) control. The disbetic subject's response to an oral glucose load is charac-tagely have backy, hypopycemia because of a failure to release insulin (type I diabetes mellitus) or the failure to rewas alonguedy to the insulin released (type 2 diabetes mellitus). (From Goodman HM: Basic Medical Enheapenedy to the insulin released (type 2 diabetes mennus). (***

***Control of Philadelphia, Lippincott-Raven, 1994, with permission.)

^{12), Hore is TIDM treated?}

- loain therapy is the mainstay of treatment. Various preparations of insulin are available to delice platmacokinetics, such that the patient can fairly accurately duplicate a normal patient of the patient can fairly accurately duplicate a normal patient of the patient can fairly accurately duplicate a normal patient of the patient can fairly accurately duplicate and patient can fairly accurate and patient can fairly accurate and patient can fairly accurately duplicate and patient can fairly accurately du
- by if bodin action during the day. Nothing substitutes for a normal endocrine pancreas as of its of section action during the day. Nothing substitutes for a normal endocrine pancreas as of its of section. in the company, most type 1 diabetics exhibit morbidity from the disease.

128. What happens if insulin therapy is not given to a patient with type 1 diabetes?

What happens it insulin therapy so the grand synthesis are decreased, while glycogen.

Carbohydrate. Glucose transport and glycogen synthesis are decreased, while glycogen. Carbohydrate. Glucose transport and gryengens olysis and gluconeogenesis are maintained. This leads to hyperglycemia, glucosuria, polyuria olysis and gluconeogenesis are maintaines. The circulatory system to maintain systemic (osmotic diuresis), dehydration, and a failure of the circulatory system to maintain systemic perfusion.

sion.

Lipid. Lipogenesis is decreased, and lipolysis is increased. This leads to hyperlipemia, In. Lipid. Lipogenesis is decreased, and input such a such as the such

his leads to severe metabone actions.

Protein. A decrease in amino acid uptake and protein synthesis and an increase in protein.

This is manifest as Protein. A decrease in animo acid appearance and urine. This is manifest as a negative

The end result is the patient loses large quantities of calories, amino acids, water, and bicarbonate in the urine. This is manifest as extreme weight loss, weakness, hyperglycemic shock,

129. What is the pathogenesis of type 2 diabetes mellitus?

The first defect is probably a decrease in the sensitivity to insulin (insulin resistance), which appears to be an inherited propensity. If this occurs without weight gain, the islet cell can usually compensate by increasing insulin secretion. If the patient gains weight and insulin resistance worsens, the islet cell response is inadequate, and hyperglycemia occurs. Eventually the insulin response to hyperglycemia wanes, and the symptoms worsen. There is usually adequate insulin secretion to prevent ketogenesis in the liver, although there is not sufficient insulin to shut off hepatic gluconeogenesis.

130. Compare and contrast type 1 and type 2 diabetes mellitus.

·	TYPE I DIABETES MELLITUS	TYPE 2 DIABETES MELLITUS	
Pathogenesis	Loss of islet cell function	Resistance to insulin	
Age of onset	Usually < 30 years	Usually > 40 years	
Ketoacidosis	Common	Uncommon	
Body weight	Very thin	> 80% obese	
Prevalence	0.5%	2-4% (may be higher)	
Genetics	Approximately 50% concordance in twins	> 95% concordance in twins	
Autoimmune	Yes	No	
Treatment	Insulin	Diet, hypoglycemia agents, appetite suppression, weight loss, exercise insulin (sometimes)	
Symptomatic	Usually	Not usually (at least, at first)	

131. List other conditions that can resemble diabetes mellitus.

Secondary diabetes

- Pancreatic disease
- Excess in the counterregulatory hormones such as GH (acromegaly), cortisol (Cushing's syndrome), catecholamines (pheochromocytoma)
- Drugs

Syndrome X (also known as syndrome of insulin resistance, subclinical diabetes, or the metabolic syndrome)

Gestational diabetes

 Glucose intolerance (hyperglycemia) usually only manifest during pregnancy. It is probably due to placental hormones (e.g., human chorionic somatomammotropin, placental steroids).

HORMONAL CONTROL OF CALCIUM HOMEOSTASIS

White keakelum flux so tightly regulated? HAVE selection flux so transportant cation in many intracellular and extracellular processes, (3-3-4) is an extracelly intracellular processes, (3-3-4) in the contract of the When a secretary important curves in many intracellular and extracellular processes.

"Second solid secretary for normal mineralization of bone, blood clotting, and plasma in the second secon " is a survey of the contract of processes, includ-ated and contain makes unknown, are secretion of hormone, neutransparies, and di-stances are arranged action potentials and retinal function; maintenance of transport of son-ter are actions and action potentials and retinal function; and cell growth and division. outputs and return potentials and return incident maintenance of the part of t

the bands form does calcium circulate in the plasma? la wass on protein (primarily albumin) In the free junized state

B. Describe daily calcium balance. poertie daily consumer the control compartment with which all other the curacefular fluid (including plasma) is the central compartment with which all other the curacefular fluid (including plasma) is the central compartment with which all other the curacefular fluid (including plasma) is the central compartment with which all other the curacefular fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment with which all other the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid (including plasma) is the central compartment of the curaceful fluid fluid (including plasma) is the central compartment of the curaceful fluid flu The caractelatar near time to the control of the caracter compartment with which all other the caracter compartments and their hormonal conon are as follows.

Gastrolntestinal tract. This is the primary site of calcium absorption. For example, out of mers are as follows:

Gardulatestimal tract. This is the position y area or carcium absorption. For example, out of Gardulatestimal tract. This is shown 400 mg (40%) is absorbed. The absorption of calcium in the atting of diethy calcium, about 400 mg (40%), is absorbed. The absorption of calcium in the atting of the carcinomatic and the calcium in the carcinomatic and the carcinomatic in the state of dictary calcium, account on the core, is associated. The absorption of calcium in the state of dictary calcium, account on the component of the vitagradient of the production of 1,25(OH),D is stimulated by PTH. Above 200 production at the Land Section of 1,25(OH)₂D is stimulated by PTH. About 300 mg (out of 1000 mg) production of 1,25(OH)₂D is stimulated by PTH. About 300 mg (out of 1000 mg) production of the vital state of from the extracellular fluid compartment to the as Digitizary, recommended to the extracellular fluid compartment to the gastrointestinal tract girl of challenge the repical net calcium absorption per during rearriof carcium of the typical net calcium absorption per day is about 10% of the calcium is seniors. Therefore, the typical net calcium na screams per may is about 10% of make although this can be changed dramatically by vitamin D excess or deficiency,

a absorb was can be primary storage site for calcium (approximately 1 kg; ~99% of total body Bone, Borne to use primary sources and total body with the plasma compartment. Bone accretion also an Calcium in bone is actively exchanged with the plasma compartment. Bone accretion distinct caronin in some process. Reclamation of calcium from bone is a process called repersons in a concess of the long term (steady state), bone formation and recorpsequences and to a sequence of the sequence of page a formation or formation is decreased without a decrease in resorption ultimately results alos of hone (e.g., osteoporosis).

Kidney, Calcium is filtered (about 10,000 mg/day) as part of the glomerular filtrate. The tites has developed efficient mechanisms for reclaiming this filtered calcium from tubular hid-reabsorption - which is stimulated by PTH.

Beekee, PTH increases extracellular calcium concentration directly by increasing calone reception from bone and increasing calcium reabsorption from renal tubular fluid and insuch by increasing calcium absorption in the gastrointestinal tract via $1,25(\mathrm{OH})_2\mathrm{D}$.

Ut. How is the secretion of PTH controlled?

PTH, produced by the parathyroid glands, is one of the only hormones whose secretion is wheel by an increase in extracellular calcium. In a simple feedback loop, a decrease in plasma abanesuls in an increase in PTH, which increases bone resorption, calcium reabsorption, and disan absorption (via 1.25(OH)₂D), all acting to restore plasma calcium. The converse is also the a that an increase in plasma calcium suppresses PTH release, which decreases the resorpto naisopion, and absorption (via 1,25(OH),D) of calcium, allowing plasma calcium to de-PTH is the most important acute controller of plasma calcium.

bk. Box do the parathyroid cells detect small changes in extracellular (plasma) calcium? her cells express a receptor with an extracellular calcium-sensing component and a 7beamstrate spanning domain; the receptor is G-protein coupled. This receptor acts via phos-Refine C and also by inhibiting adenylate cyclase. Parathyroid hormone is one of the few Page excess is inhibited by an increase in calcium.

137. Other than PTH and 1,25(OH)2D, is there another calcium? int ?

Calcitonin, produced by the parafollicular cells of the thyroid gland, inhibits bone resormer. 138. Describe the pathway that produces 1,25(OH)₂D.

The endogenous vitamin D steroidogenic pathway. Conversion of vitamin D to 25(OH)D in the liver is relatively unregulated while activation of 25(OH)D to 1,25(OH).D in the kidney is highly regulated (e.g., stimulated by PTH). Vitamin D can also be obtained in the diet as choleculciferol (animal vitamin Di) or enco calciferol (plant vitamin D2). (From Griffin JE, Ojeda SR (eds): Textbook of Endocrine Physiology, 3rd ed New York, Oxford University Press, 1996, with permission.)

1. The vitamin D (calciferol) pathway is a steroidogenic pathway catalyzed by a series of cytochrome P-450 enzymes. There are two forms of vitamin D in the diet: animal vitamin D₁ (cholecalciferol) and plant vitamin D2 (ergocalciferol). In addition, vitamin D3 can be liberated from the skin from 7-dehydrocholesterol via the action of ultraviolet light.

 Once vitamin D₂ or D₃ reaches the plasma compartment, it is converted to 25(t)Hip
 Once vitamin D₂ or D₃ reaches the plasma compartment, it is converted to 25(t)Hip 2. Once vitamin 2.

2. Once vitamin 2.

2. Once vitamin 2.

3. Once vitamin 2.

4. Once vitamin 2.

4. Once vitamin 2.

5. Once vitamin 2.

6. Onc he down of 25 nyama years and one in the over. This is a relatively unregulated step, at the devated 1.25(OH), D is thought to inhibit this step (end-product inhibition). In physio, and the down of is the second of L2(UII) LV is throught to innoin this step (end-product inhibition). In physio-body devarrations, 25(OH)D has little biologic activity, whereas it may have calciumyie ef-the devaled. nes when elevated. when elevated.

when elevated to the active form, 1.25(OH), D, by 1-hydroxylase enzyme heated

\$25(OH) is activity of 1-hydroxylase is increased by PPH and increased enzyme heated

 3. 25(OH)D is accounted to the activity of 1-hydroxylase is increased by PTH and inhibited by plasma phase of the activity of 1-hydroxylase is increased by PTH and inhibited by plasma phase in the plasma phase in white. The activity of insponsorates is increased by PTH and inhibited by plasma phase is in 120(H). D (end-product inhibition). 25(OH)D can also be inactivated to 24.25(UH). D and also be inactivated to 24 pear and production in the kidney.

what is the best method to assess the activity of the vitamin D pathway? what is the Dest. included the pathway.

Measurement of serum 1,25(OH), D. the active component of the pathway.

What is the best method to assess vitamin D intake and stores? What is the uses a secum 25(OH)D because it reflects the summation of vitamin D from disand sources available for activation to 1,25(OH),D.

	ns of the major calcium	
HORMONE Parathyroid hormoste	Bone Kidney	Calcium and phosphate resorption Calcium reabsorption Phosphate reabsorption Conversion of 25(OH)D to 1,25(OH)A Calcium and phosphate resorption
Calcinnin	Bone Kidney	Calcium and phosphate reabsorption
Vancin D [1,25(OH) ₂ D]	Bone Gastrointestinal tract	Calcium and phosphate absorption

Discuss other hormones that affect bone and calcium metabolism.

Goadal steroids. Androgens and estrogens are necessary for the pubertal growth spurt and dour of the epiphyseal (growth) plates in bone and, therefore, before adulthood, favor bone for Box In earlit, estrogen decreases one resorption (probably PTH-mediated) and therefore Process desired to the control of th hospitalism) is characterized by a loss of bone mineral density (osteoporosis).

Glecoorticoids. Although cortisol is necessary for normal skeletal growth, cortisol in exonecorricoids. Although cortisol is necessary for normal sketcia growin, consistent of a monocorricoids and the consistent of the consistency of t Propose scheding hypercalciuria and an inhibition of 1,25(OH). D-mediated calcium absorpwas a straintestral tract. The resultant secondary hyperparathyroidism accelerates bone straintestral tract. The resultant secondary hyperparathyroidism accelerates bone forms from the secondary hyperparathyroidism accelerates bone forms from the secondary hyperparathyroidism. a use gastrointestinal tract. The resultant secondary hyperparathyroidism accesses seems before producing tracts and tracts appears to inhibit osteoblastic bone formation directly.

Privide plucocorticoid may induce secondary hypogonadism.

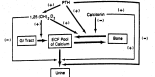
Privide hormone, Lack of adequate thyroid hormone delays ossification of bone growth ruly ricess glucocorticoid may induce secondary hypogonadism. introd hormone. Lack of adequate thyroid hormone delays ossification of rance govern-nesses and can retard bone development in children. Excess thyroid hormone may cause in-

steed home resorption. GH. GH simulates IGF-1, which increases bone formation.

Decribe the overall regulation of calcium balance.

is the steady state, calcium make should roughly equal calcium loss via the gastrointextih the sleedy state, calcium intake should roughly equal calcium loss via the gastron-beat and the time. Calcium intake should roughly equal calcium loss via the gastron-the state of the time. Calcium absorption is increased by 1,25(OH),D, whose production from \$4.55(OH),D, whose production from \$4.55(OH),D, whose production from \$4.55(OH),D, whose production is increased by 1,25(OH). and the time. Calcium absorption is increased by 1.25(OH). D, whose production from the PH Also increases calcium resorption from bone and calcium reabsorption from the photos of the PH also increases calcium resorption from bone and calcium reabsorption from the photos.

* age It plants calcium is high, calcitonin may decrease bone recorption.



Integration of the hormonal regulation of calcium ballines. PTH Increases planns (ECP) calcium by increase in glove encoping, in recentage rehardports of calcium in the kidney, and by increasing real production of 1,25(011), by which stimulates gastrointestinal absorption of calcium. Although circle intends on screen know encoping an pathophologic concentrations and with plannas/fold course, in phylologic role in many (Front Goodman HM: Basic Medical Endocrinology, 2nd cs. Philadelpsia, Lippanorit-Raven, 1994, with permitted).

144. Briefly explain phosphate balance. Phosphate resorption in the gastrointestinal tract accompanies calcium and is increased by

1,2S(OH)₂D. Phosphate resorption also accompanies calcium and is increased by PTH. The main difference between calcium and phosphate balance occurs in the kidney, where PTH increases phosphate excretor. This is why patients with elevande PTH have hypercalcemia and hypophophatemia—they reclaim calcium from the urine while allowing phosphate to be excreted.

145. Discuss the pathogenesis of PTH-dependent hypercalcemia. PTH-dependent hypercalcemia is defined as primary hyperparathyroidism and is usually

one to parally roll adorma. These terms or prometer and are provided in the production of the producti

146. Explain why a patient with elevated PTH has hypercalciuria if PTH increases renal calcium reabsorption.

When plasma calcium is elevated, the filtered load of calcium in the kidney increases. Although PTH does increase tubular calcium reabsorption of calcium, the filtered load of calcium may exceed the renal reabsorptive capacity, and calcium spills into the uringly.

147. What are PTH-independent causes of hypercalcemia? • Vitamin D intoxication

PTH-related peptide (PTHrP) secretion from a malignancy

148. Discuss the pathogenesis of vitamin D intoxication.

Hypercalcentus is not necessarily due to an elevation in 1,25(OH),D but may be due to small but significant biologic activity of 25(OH),D and that elevated 25(OH)D index of increased vitamin D stores) may displace 1,25(OH),D from its plasma carrier protein, increasing its free, but

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The increase in gastrointestinal absorption of calcium increases plasma calcium and possible in market home and possible in market home. The increase in a successful calcium exerction and results in marked hypercalcular and results in marked hypercalcular.

piccus the endocrine causes of hypocalcemia, pieces the encovering gland function leads to primary hypoparathyroldism, Lack of PTH A lack of parathyroid gland function leads to primary hypoparathyroldism, Lack of PTH A lack of PTH and a decrease in passage of the primary hypoparathyroldism. 16. Per col parathyrou game rous connectus to primary hypoparathyroidian. Lack of PPH A loss (address to increase 1.25(OH), 1) and a decrease in gastrointestinal absorption of coloridation, the lack of PPH activity in the kidney prevents the renal resonant coloridation. in a salter to increase and a severage in gastrointestinal absorption of en-gals in a salting, the lack of PTH activity in the kidney prevents the renal response to hypocal, into in addition, the lack of PTH activity in the kidney prevents the renal response to hypocal, into in a series calcium reabsorption. Also, without PTH to inhibit phosphate reads. man in addition, the nace of the man and the man real response to hypocal-tion in addition, the nace calcium real-surption. Also, without PTH to inhibit phenybate real-surption, by-terial successit may cassic.

popersphatemia may ensue. Sophitemia may custo.

Neghitemia may also result from a failure to take in adequate vitamin D (rickets in chilthrowshemia may also result from a failure to take in adequate vitamin D (rickets in chilthrowshemia may customer).

are estremulacia in actults). ostromaticas in an analysis of calcium and vitamin D may also lead to hypocaleenia. General States of the Appendix This is called secondary to provide the secondary hyperparathyroidism is also often as the skilney in an order of assitu calcium. Secondary hyperparathyroidism is also often as This metrous and calcium. Secondary hyperparathymidism is also often a consequence of moster plasma calcium. Secondary hyperparathymidism is also often a consequence of most of the inability to generate 1,25(OH). D and nothing serge to refere plasma carconal season and a per paramyroanism is also often a consequence of the inability to generate 1,25(OH) D and, perhaps, a loss of renal calcium and influe because of the inability to generate 1,25(OH) D and, perhaps, a loss of renal calcium

rebsorptive capacity. 18t List the symptoms of hypocalcemia. LIST the symposium of the control (seizures) nerve

removes
 abnormal electrocardiogram (prolonged Q-T interval)
 Cardiovascular: abnormal electrocardiogram (prolonged Q-T interval)

gt. How can one distinguish between hypercalcentla caused by a PTH-secreting adenoma

and hypercalcennia caused by PTHrP? represervants causes.

The best way is to measure intact PTH. Although PTH and PTHrP have sequence homology. and controlly used assays for intact PTH do not measure PTHrP, PTH does not have to be above tesferore range to suggest primary hyperparathyroidism. PTHrP-induced hypercalcenia should

saroress intact (normal) PTH. ISt. How can one distinguish between vitamin D intoxication and hypercalcemia of maligmany caused by PTHrP since they both have suppressed intact PTH?

Assays for PTHrP provide accurate results. Furthermore, patients with vitamin D intoxicaion usually have elevated 25(OH)D levels (an index of vitamin D stores). Measurement of aptragenous (urinary) cAMP has been done in the past because this is an index of PTH activity ad is increased by both intact PTH and PTHrP.

FEMALE REPRODUCTION (EXCEPT FOR PREGNANCY AND LACTATION)

153, Discuss the factors controlling fetal sexual differentiation

User most circumstances, genotype (genetic sex) and phenotype (sexual characteristics) are team that is, an XX conceptus develops into a female baby and an XY conceptus develops into had bely. The presence of a Y-chromosome (H-Y antigen) induces development of testes. oney. The presence of a Y-chromosome (H-Y antigen) induces development of ownies. The the same tensorement of the male reproductive trust from the wolfand data. Testosterone, which induces development of the male reproductive that the target data. Testosterone is also converted to dihydrotestosterone (DHT) by 5ce-reductive in target has been resourced is also converted to dihydrotestosterone (DHT) by Secretic millerian-base when induces development of male genitalin. In addition, the testes secrete millerian-tests for the secretary of festes, and who induces development of male genitalia. In addition, the testes secret backg factor (MIF), which causes regression of the millerian ducts. The absence of testes, and backgrain ducts. have failed a state (MIF), which causes regression of the millerian ducts. The answers of the development of the state of testosterone, DHT, and MIF, allows wolffian ducts to regress, the development of the state of testosterone. as bease of isstosierone, DHT, and MIF, allows wolffian ducts to regress, inc.

bease restales, and the formation of the female reproductive tract from the mullerian ducts.

the time the histology of the country give insight into its function?

Depola. The number of germ cells (potential oncytes) peaks at approximately 6 million at

Endocrine Physiology 234 monoyauxe (approximately 50 years of age), almost no viable germ cells remain. Primary follicles. These have the potential to start maturation.

Gradian follicle. The dominant follicle is filled with fluid and contains a mature occyte ready for evaluation. It produces large amounts of estrogen and is primed to produce large amounts of progesterone after ovulation. Corpus luteum. This develops from the ruptured follicle after ovulation. It is full of

about 6 months of gestational age and decreases thereafter via a process called atresia. By

Primary follicles. These begin to develop intrafollicular fluid and proliferating steroido.

steroidogenic cells, which produce large amounts of progesterone (and estrogen, to a lesser extent). Atretic follicle. This is a follicle whose occyte was not ovulated but regressed during mat.

uration (nondominant folliele). Retrogressive corpus luteum. If conception does not occur, the corpus luteum "dies " 155. List the margi female andorrine changeathroughout-life.

 HCG (from trophoblast and placenta) and fetal FSH and LH stimulate development of ovarian germ cells.

 LH and FSH burst approximately 4 months postpartum (sexual differentiation of the brain?) Adrenarche -- increase in adrenal androgens at about 8 years of age.

 At onset of puberty (8–10 years old), GnRH pulses from hypothalamus increase, which stimulates LH and FSH and increases ovarian function.

Increase in ovarian steroids induces development of secondary sex characteristics

 Menstrual cycles (menarche) start at approximately 12 years of age. In addition to development of secondary sex characteristics, pubertal estrogens stimulate.

growth spurt (assuming presence of adequate GH). · Estrogens also stop growth spurt by causing fusion of epiphyseal plates in bone.

 At menopause (at approximately 50 years of age), ovaries stop producing steroids. This leads to the absence of menses as well as other physiologic (hot flashes) and psychologic ical changes. Lack of steroid negative feedback leads to an increase in FSH and I.H.

156. How are pearian-steroids synthesized?

gonic cells (theca and granulosa cells).

The pathway is essentially the same as that outlined in the adrenal cortex section, particularly with respect to progesterone production. After 17-hydroxylation and androgen production. androstenedione is converted to estrone, and testosterone is converted to estradiol by the enzyme aromatase. It is generally believed that this process requires the two follicular cell types-thera and granulosa cells - to work in what has been called the "two-cell hypothesis of ovarian steroido-

genesis" (see figure, next page). The theca cell expresses primarily LH receptors. LH stimulates steroidogenesis and large amounts of androgen production. The theca cell is relatively devoid of aromatase activity.

Androgens diffuse through the basal lamina into the granulosa cell. The granulosa cell expresses primarily FSH receptors, although it can express LH receptors just before ovulation. An-

drogens from the theca cell are aromatized to estrogens primarily in the granulosa cell.

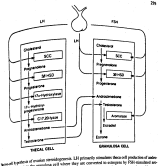
157. Since the gonadal hormones are steroids, do they circulate bound to carrier proteins similar to cortisol?

Estradiol and estrone, because they have undergone the 17,20 lyase reaction, do not resemble cortisol very much and therefore do not bind to CBG. There is another carrier protein called sex hormone-binding globulin (SHBG) that carries estradiol (approximately 38%). Progesterone

does circulate bound to CBG (approximately 18%),

158. If progesterone only binds about 18% to CBG and estradiol only binds about 38% to

SHBG, do these gonadal steroids circulate mostly in the free form? No, because they are bound significantly by plasma albumin, with estradiol having about 60% binding with albumin, and progesterone about 80% binding with albumin. Therefore, estra-



he process represents the granulosa cell where they are converted to estrogens by FSH-stimulated ansake armos una constitue de la also produce progesterone but do not have adequate enzymatic machinery to state across, consumers to androgens. (From Griffin JE, Ojeda SR (eds): Textbook of Endocrine Physiology, 3ed d See York, Oxford University Press, 1996, with permission.)

deland progesserouse circulate approximately 2% free, with about 98% percent bound to carrier mens or albumin.

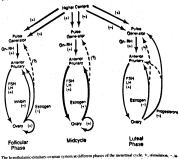
19. Do the ovaries produce peptide hormones? Relaxin: relaxes pelvic ligaments

lehibin: selective inhibition of FSH Activins: selective stimulation of FSH

M. Bow is the hypothalamic-pituitary-ovarian control system similar to and different but the HPA axis? in the propulertal and postmenopausal state, they are similar. Increases in GnRH pulses from

by the chairman and postmenopausal state, they are suntian, increases LH and FSH. These two Armore, strukte ovarian steroidogenesis, which, via negative feedback, decreases the secretion doubt and LH and FSH. Therefore, if ovarian steroids are deficient, LH and FSH are increased herparadoregic hypogonadism analogous to primary adrenal insufficiency); and if hypopitu-Sources, LH and FSH are inappropriately low (analogous to secondary adrenal insufficiency). Between menanche and menopause, there are quite a few differences. The ovaries produce want hemore, called inhibin, which inhibits the release of FSH. Therefore, there are parallel

when the strong called inhibin, which inhibits the release of FSH. Therefore, increase when the strong called inhibin, which inhibits the release of FSH. Therefore, increase and the strong called the strong cal her difference is the existence of positive feedback. During a specific time (midcycle) in the constitute is the existence of positive feedback. During a specific time (muo. stable) (vid. estrogen actually stimulates LH and FSH release. This results in a surge in go-



hibition. Dushed lines indicate hypothetical relationships. Line thickness indicates the intensit of sh tion. (From Goodman HM: Basic Medical Endocrinology, 2nded, Philadelphia, Luyuncus Raves, 1944, and permission.)

161. Outline a typical menstrual cycle. Because this is truly a cycle, day 1 is somewhat arbitrary. Because the major physical sign is the onset of menses, however, this is considered day 1.

- Follicular phase: The emergence of the dominant follicle.
- Menses are induced by decreases in estrogen and progesterone. 2. Increase in FSH on day 28 is induced by loss of steroid negative feedback. Increase in
- FSH on day 28 promotes the maturation of 6-12 primary follicles.
- 3. Increase in FSH on day 28 followed by LH on days 2-5 increases estrogen production 4. One (usually) follicle becomes dominant and increases estrogen production. This estro-
- gen inhibits FSH secretion by negative feedback. Estrogen concentration may correlate with the size of the dominant follicle. 5. The nondominant follicles cannot survive the decrease in FSH and undergo atresia. The
- dominant follicle survives this decrease in FSH because it has increased expression of LH and FSH receptors.
- 6. The system shifts from negative to positive feedback so that the large increase in estregen from the dominant follicle induces the LH surge. Testrogen causes T LH causes T estrogen causes T LH.
 - 7. Preovulatory increase in progesterone potentiates estrogen positive feedback on l H. 8. LH surge (and FSH surge) occurs, with the ratio of LH to FSH increasing dramatically
- This induces resumption of meiosis in the occyte of the dominant follows. 9. FSH surge induces LH receptors on granulosa cells to prepare the follole for marchemation into the corpus luteum.

 Betrogen some Sections as LH reaches its peak. The loss of the LH receptor on the thecateranulosa cells, and the last of the LH receptor on the thecateranulosa cells, and the loss of estrogen positive feedback stimulation. 10. Estrogen starts to decrease as LH reacties its peak. This is hypothesized to be due to possibilition of the Lin control of the Lin control of the Lin control of the Lin control of Lin bull The loss of essuages, possible recupack stimulation of LH terminates the property of progesterone to estrogen may be a negative feedback signal region of progesterone to estrogen may be a negative feedback signal and the company of the expension of the owner from the dominant feetback signal and the company of the expension of the owner from the dominant feetback signal and the company of the expension of the owner feetback signal and the company of the expension of the company of the expension of the owner feetback signal and the company of the expension of the expension of the owner feetback signal and the company of the expension of the owner feetback signal and the expension of the expension of the owner feetback signal and the expension of the owner feetbac ratio of programmer of the ovum from the dominant follicle.

Ovulation occurs owing to prior LH surge.

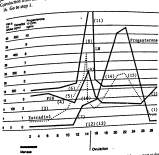
12. Ovulation

Luteal phase: Progesterone (and estrogen) secretion from the corpus luteum.

Luteal phase: bloom is formed primarily by granulosa calls. Lated phase: Programment of the (2) Corpus luteum is a source of progesterone and estrogen increases; the process is depen-led. Corpus luteum secretion of progesterone and estrogen increases; the process is depen-led. Corpus adequate) LH levels.

des (a low (but adequate) LH levels.

blow (but account of the control of is processed and extragen occrease owing to finite life of corpus luteum in the absence is processed open occurs. HCG from the trophoblast rescues the distinction gondotropin levels. (If fertilization occurs, HCG from the trophoblast rescues the distinction and prevents menstruation.) The corpus luteum dies unless fertilization. displaced gonadotropin revers. (It is intraction occurs, HCG from the trophoblast rescues the displaced and prevents menstruation.) The corpus luteum dies unless fertilization occurs and opposition from the trophoblast rescues the corpus luteum. locus locus and processing the trophoblast rescues the corpus luteum.



na menstrual cycle. Days of the cycle are shown across the bottom. Menses start at day 1; ovuhere sunly occurs around day 15. Each step is identified numerically in the text. (Adapted from Speroff L. Samy occurs around day 15. Each step is identified numericarty in use text covariate to the St. Kase NG: Clinical Gynecologic Endocrinology and Infertility. Baltimore, Williams & Wilkins.

Describe the proliferation of granulosa cells during follicular development.

where the proliferation of granulosa cells during follicular developments in FSH white dominant follicle grows, it secretes more and more estrogen despite decreases in FSH to be in the following the control of the co as each follower phase. Local estradiol production within the dominant follicle increases the to the property of the propert By more FSH receptors. Positive feedback of estrogen from the dominant content of the later following seeks, which induces LH receptor expression on granulosa cells. By the later following seeks the back increased greatly Sec just chose value induces LH receptor expression on granulosa cells. By the later increased greatly the property of the dominant follicle have increased greatly that granulosa cells of the dominant follicle have increased greatly that granulosa cells of the dominant follicle have increased greatly that granulosa cells of the dominant follicle have increased greatly that granulosa cells of the dominant follicle have increased greatly that greatly the granulosa cells of the dominant follicle have increased greatly that greatly the granulosa cells of the dominant follicle have increased greatly that greatly the greatly the greatly that greatly the greatly the greatly that greatly the greatly that greatly the greatly that greatly the greatly the greatly the greatly the greatly that greatly the greatly thad greatly the greatly the greatly the greatly the greatly the gr t before ovulation), the granulosa cens of one --

hat. How do changes in the endometrium of the uterus correlate with the phases of the men. streat cycle? cycle? The endometrial cycle is in synchrony with the menstrual cycle and is a hallmark of the extraco arism actions of gonadal steroids.

 Proliferative phase occurs during the follicular phase of the menstrual cycle. Estrogen
 Proliferative phase occurs during the follicular phase of the menstrual cycle. Estrogen Proliferative phase occurs using the stimulates the growth of the epithelial and stromal layers. The thickness of the endo.

stimulates the growth of the epimeria.

The spiral arteries, which are increases, and the uterine glands increase in size. The spiral arteries, which are increases, and the uterine glands increase in size. the primary blood supply for the endometrium, elongate.

 the primary blood suppry on the clutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the Secretory phase occurs during the lutter phase of the menstrual cycle and prepares the secretory phase occurs during the lutter phase occurs during the lut Secretory phase occurs using an example of the conceptus. Progesterone stimulates secretory acity. enklometrum for implantation of the control of the uterine glands, and glycogen production increases. The stroma becomes edematous, and the spiral arteries coil.

 Monstrual phase correlates with the end of the luteal phase of the menstrual cycle. The Meastrust phase correlates the secretion from the corpus luteum induces vasoconstriction loss of gonadat steriou sections to the endometrium. The endometrial lining tspusm?) of the spiral arteries and necrosis of the endometrium. The endometrial lining is sloughed off in the form of menstrual bleeding.

164. Are there other cornervarian actions of estrogen and progesterone?

Oviduets — Estrogen increases cilia formation and contractility, and progesterone increases secretory activity and decreases contractility. Mymmetrium — Estrogen increases growth and contractility, and progesterone decreases

contractility. Cervix - Estrogen induces a watery secretion, and progesterone stimulates the production of dense, viscous secretions.

Vagina -- Estrogen induces epithelial proliferation, and progesterone induces epithelial differentiation.

Breasts - Estrogen stimulates development of the duct system and adipose tissue (e.g., at puberty), and progesterone induces formation of secretory alveoli (e.g., during pregnancy), Bone - Estrogen stimulates and terminates pubertal growth spurt. Estrogen inhibits bone resorption.

Other - Estrogen increases SHBG, CBG, and TBG. Estrogen alters lipid profile.

165. List and briefly describe fertilization of the ovum and implantation of the conceptus. Ovum transport: Ovulated oocyte collected by fibrial end of fallopian tube

Sperm transport and capacitation: Contact with female tract activates sperm function

· Fertilization: Usually occurs in fallopian tube Implantation and placentation: Blastocyst usually implants on endometrial lining approximately 7 days after ovulation

166. What is prenopause?

Menopouse is the age-related cessation of regular menses during the female climacteric when reproductive cyclicity gradually disappears. Usually, menstrual cycles become irregular before they completely stop. Menopause is characterized by a loss of ovarian function probably as a result of exhaustion of available follicles lost because of atresia. Because of the decrease in estrogen production from the ovary, LH and FSH increase owing to loss of negative feedback. In that sense, menopause can be defined as hypergonadotropic hypogonadism,

- 167. What is amenorrhea? Printary amenorrhea: the failure to have menarche (the onset of menstrual cycles at puberty). It is currently believed that the failure to have normal menstrual cycles by the age
 - Secondary amenorrhea: the premature cessation of normal menstrual cycles. Causes include
 - pregnancy, hyperproductinemia, premature menopause, excessive exercise, and weight loss Oligomenorrhea: irregular menstrual cycles.

MALE REPRODUCTION

is the male XX ganotype), is there a relationship between gonadal function and physical field of the female, the development of a male at the female, the development of a male at the female, the development of a male at the female at the fe is much throughout life?

is the many strongerous the development of a male phenotype requires a signal from the Assipposet. The secretion of müllerian inhibitory factor (MIF) induces a signal from the A supposed to me to millerian inhibitory factor (MIF) induces regression of the secretion of millerian inhibitory factor (MIF) induces regression of the supposed allows the wolffian ducts to develop into the internal male and allows the fetal testes induces sometime for the secretion of the sec servering sounds. The servering sounds of the serverin person due to make a more and a new and a more and a make general make

public of testosterone is absent or if there is resistance to the action of testosterone is absent or if there is resistance to the action of testosterone (testicular public of testosterone) phenotype develops. M. H. testing of the control of the transfer o institute the subsequent increase at about 6 months of age (analogous to a similar April Portunition).

After parturnization.

After parturnization in the subsequent increase in testicular steroidogenesis and androgen secretion form in the sexual differentiation of the brain. end in the sexual differentiation of the brain. graft in the Sexual control of the Sexual co

Adequate (estronomental androgens), which usually occurs at approximately 8 years of series harbinger of the onset of puberty.

is a harbinger or age. LH and FSH increase, leading to a marked increase in testicular At 10-14 years of age. At 10-14 years or age.

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At 10spit of the voice, axillary and pubic hair, increased sweat glands), and pubic hair, increased sweat glands).

guberty (deepening of a male climacteric in the elderly is a controversial topic. Although testos-The experience cases as men age into their forties to sixties, this may not represent clinical byceradism, and spermatogenesis can be maintained.

18. Sammarizodorticular steroidogeness. Summariant This is essentially the same as adrenal androgen production, in which pregnenolone and

inguistrius are converted to testosterone. Although testosterone is the primary androgen produced

This is essentially the DHEA and androstenedione by P450c17. DHEA and androstene-spectation are converted to DHEA and androstenedione by P450c17. DHEA and androstenegare can be some youngen produced by the testes, it is not the most potent. Dihydrotestosterone (DHT) is produced from testosterone (DHT) h So eductase primarily in target tissue (peripheral activation). In addition, sesticular androgens as te converted to estrogens in males primarily by peripheral conversion by aromatase. 176. Review the circulating gonadal steroids in the male and state their sources.

> 95% of circulating testosterone is from the testes.

> 80% of the circulating DHT is from peripheral conversion of testosterone.

 > 80-90% of circulating estrogen is from peripheral conversion of precursors. > 90% of circulating DHEA (sulfate) is from the adrenal cortex.

III. List the hormonal and somatic changes during male puberty.

1. GrRH pulses from the hypothalamus increase FSH and LH secretion.

2 LH stimulates testosterone production, which induces development of secondary sex deractenstics.

3. FSH stimulates spermatogenesis.

 ACTH (or some other pituitary factor) increases adrenal androgen production (adrenanche). 5. GH maintains linear growth.

 Testesterone enhances the secretion of GH and initiates the pubertal growth spart. leasterone then causes epiphyseal fusion (termination of the pubertal growth spurt).

What is the significance of the pulsatility of GinRH release from the hypothalamus? on the significance of the pulsarillty of GinRH release from the hypomanantasis, the hallmarks of the gonadotropin control system in males and females is pulsarilly. by me of the hallmarks of the gonadotropin control system in males and tensies to waith released in pulses with about a 90-minute frequency, although this is variable by many and a state of the majors are necessary to the pulses are necessary to the puls being ad within subjects and by time of day (and even seasons of the year). GRH pulses are ne-ber to adwithin subjects and by time of day (and even seasons of the year).

on and within subjects and by time of day (and even seasons of the year). Office I H at first, lead to , the property of the p has a pulse. Constantly high levels of GRRH, although stimulating LTB and photostopic and printerly gonadostrophs and a decrease in LH. This is the basis for using GRRH at the photostopic and a decrease in LH. This is the basis for using GRRH at the photostopic and a decrease in LH. This is the basis for using GRRH at the photostopic and the photostopic and a decrease in LH. **Polica not printiary gonadorrophs and a decrease in L.H. This is the basis for using a printiary gonadorrophs and a decrease in L.H. This is the basis for using a printing the proper advantage of the proper advantage of the proper advantage of the property of the p

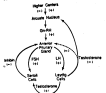
173. Describe the overall equiation of testicular function.

240

Describe the overall degulation or the most part, similar to the HPA big the hyperbalamic-primitary desticular (HPT) axis is, for the most part, similar to the HPA big the hyperbalamic primitary destination regulates the release of pulses of GnRH from the hyperbalamic into the brain regulates the release of pulses. Describe the one of the probability described and the probability of the polyal annies printing the brain regulates the release of pulses of GnRH from the bypacoda. It would input into the brain regulates the release of pulses of GnRH pulses significantly the pulses of the pulses o The hyse surface the brain regument into the anterior pituitary. GraPH pulses stimulate the process and a new post of the process into the long portal veins, which drain into the anterior pituitary. GraPH pulses stimulate in and the new transfer of the Leydig (interneting the Le and FSH release.

cells in the testes. Lri and in the tester.
 SSH stimulates Sertoli cells (in concert with local testosterone) to increase spermay,
 FSH stimulates Sertoli cells (in concert with local testosterone) to increase spermay,

 rest summer or tein (a local factor), and inhibin production, is androgen binding protein (a local factor). androgen binding protein (a rocar ta-androgen binding protein (a rocar ta-tandrogen binding release.



HPT axis. + indicates that GoRH stimulate FSH and L.H. that FSH stimulates Sensites while LH stimulates Leydig cells, and the Leydig cells stimulate Scrtoli cells (Jen Crine). — indicates that inhibin produced from Sertoli cells inhibits FSH release white teas. terone inhibits GnRH release (negative log) back). Direct inhibition of LH by issostrage has not been firmly established (doned line) (From Goodman HM: Basic Medical End. crinology, 2nd ed. Philadelphia, Lippings. Raven, 1994, with permission.)

174. What is the main difference between the HPT and HPA axes?

The testes produce two negative feedback signals: Testosterone inhibits LH (and PSH): hibin inhibits FSH release.

175. Categorize the actions of FSH and LH on the testes.

LH stimulates steroidogenesis (testosterone synthesis and release) from the Leydig (numi tial) cells. Although LH was named for its luteinizing action in the female, its effect in males in crease in androgen) is analogous. In fact, in the past, LH has been called interstitul cell-sine lating hormone. FSH stimulates androgen binding protein from Sertoli cells into the lumen of the serial

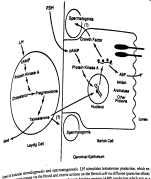
erous tubule. Androgen binding protein acts as a local testosterone sink, which dramstically creases the local concentration of testosterone that is necessary for sperm maturation. FSH also stimulates spermatogenesis. (See figure, next page.)

176. Why are LH and FSH necessary?

LH is mecessary to stimulate testosterone, which has systemic effects and local effects. stimulates spermatogenesis and increases local testosterone concentration by increasing autogen binding protein release from Sertoli cells into tubular lumens.

177. What are the major actions of androgens?

Fetal development: Testosterone stimulates internal genitalia and testosterone stimulates (DHT) stimulates external genitalia. Puberty: Testosterone and DHT increase secondary sex characteristics (musculator.)



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meaning the Medical Endocrinology, and od. Priminogrania. Lippenson-Ravell, 1999, was permission, journal with the control of the Control of

historie closure of epiphyseal plates.

Adultioned: Actions include maintenance of normal skeleton, libido, spermatogenesis, and tester secondary sex characteristics.

IL Outine the profile of male puberty.

Box Is

KCE (YEARS)	PHYSIOLOGIC CHANGES
About S	E. Vindore makele increases in
400E S	Adrenal androgens (DHEA and androstenedione) induce subtle increases in
	secondary sex characteristics (e.g., light mustache).

Mang 9-10 GraPH pulses increase, which stimulates LH and FSH release. This stimulates texticular growth and increases in testosterone.

The patental increase in public and axillary hair starts followed by an increase in pentic growth.
The patental growth spurt has usually started with a peak growth velocity at about

age 14-15 years.
The growth spurt starts to wane.

First adult height is usually achieved (although growth until age 20 is not

179. Is there an event in males analogous to menopouse in females?

Total testosferone levels do tend to decrease as men age but usually remain within the nor. Total testesserone levels do tend to occident to desterone may decrease due to changes in SHBG mail range. More importantly, free (bioactive) testesterone may decrease due to changes in SHBG mal range. More importantly, tree torractory, and making men is not a ubiquitous finding characteristics. Although hypogenadism in aging men is not a ubiquitous finding (like binding characteristics. Annuaga ayrogameters because the special special special special to be adequate for fertility in men in their eighties.

180. What is the most common disorder of the HPT-axis? Hypogonadism (a decrease in testicular function).

181. Discuss the causes of male hypogonadism:

Hypogonadism in males can be generally classified as two types:

- 1. Testleular dysfunction is due to a decrease in testosterone production from the testes. LH and FSH increase because of a loss of negative feedback. Therefore, this is called hypergonado. tropic hypogonadism and is analogous to primary adrenal insufficiency.
- 2. Hypopitultarism is called hypogonadotropic hypogonadism and can be due to an idiopathic decrease in LH and FSH or due to panhypopituitarism. "Hypogonadotropic" may be misleading because LH concentrations are often in the normal range in patients with hypogonadotropic hypogonadism. The LH levels are inappropriately low for the low testosterone.

Another cause of hypogonadotropic hypogonadism is hyperprolactinemia, which is usually due to a prolactin-secreting pituitary adenoma. Elevated prolactin levels inhibit gonadotropin secretion and induce hypogonadism in males (and amenorrhea in females).

182. What are the symptoms of hypogonadism in males?

Symptoms depend on the age of onset.

- Androgen deficiency or insensitivity to androgens in early fetal development leads to varying degrees of ambiguity of the genitalia and male pseudohermaphroditism.
- Prepubertal androgen deficiency leads to limited secondary sex characteristics and eunuchoid skeletal proportions because, even though there is no androgen-mediated pubertal growth spurt, there is also failure to close the epiphyseal plates and the long bones continue to grow. Therefore, the arm span of these individuals is longer than a typical
- · Androgen deficiency after puberty usually results in decreased libido, impotence, and low energy levels. If androgen deficiency continues for longer periods of time, there can be a decrease in facial or body hair.

183. What is the most common cause of male hypogonadism?

Klinefelter syndrome, which occurs in about 0.2% of male births.

184. Describe the genotype and phenotype of Klinefelter syndrome.

The most common genotype is XXY (an extra X chromosome). An XXY genotype usually results from meiotic nondisjunction during gametogenesis. The phenotype usually appears at puberty and includes increased lower-to-upper body segment ratio, gynecomastia, small penis, and sparse upper body hair. The testes do not develop normally and are usually small and fibrotic. The decreased testosterone production usually leads to elevated LH and FSH concentrations.

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