Ion Channels and Signaling in the Pituitary Gland

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Endocrine pituitary cells are neuronlike; they express numerous voltage-gated sodium, calcium, potassium, and chloride channels and fire action potentials spontaneously, accompanied by a rise in intracellular calcium. In some cells, spontaneous electrical activity is sufficient to drive the intracellular calcium concentration above the threshold for stimulus-secretion and stimulus-transcription coupling. In others, the function of these action potentials is to maintain the cells in a responsive state with cytosolic calcium near, but below, the threshold level. Some pituitary cells also express gap junction channels, which could be used for intercellular calcium signaling in these cells. Endocrine cells also express extracellular ligand-gated ion channels, and their activation by hypothalamic and intrapituitary hormones leads to amplification of the pacemaking activity and facilitation of calcium influx and hormone release. These cells also express numerous G protein-coupled receptors, which can stimulate or silence electrical activity and action potential-dependent calcium influx and hormone release. Other members of this receptor family can activate calcium channels in the endoplasmic reticulum, leading to a cell type-specific modulation of electrical activity. This review summarizes recent findings in this field and our current understanding of the complex relationship between voltage-gated ion channels, ligand-gated ion channels, and G protein-coupled receptors in pituitary cells. (Endocrine Reviews 31: 0000–0000, 2010)

I. Introduction

II. Pituitary Cell Types
A. POMC-producing cells
B. Heterodimeric glucoprotein-producing cells
C. GH- and PRL-producing cells
D. Nonsecretory cells

III. Ion Channels Expressed in Pituitary Cells
A. Voltage-gated channels
B. Chloride channels and transporters
C. Channels expressed in and controlled by the endoplasmic reticulum

IV. Spontaneous Electrical Activity
A. Spiking and bursting
B. Pacemaking mechanisms
C. Channels involved in spike depolarization
D. A mechanism for bursting
E. Functional roles of spontaneous spiking

V. Signaling by Gap Junction Channels
A. Connexins
B. Pannexins

VI. Signaling by Receptor Channels
A. Cys-loop family of receptor channels
B. Glutamate receptor channels
C. Purinergic receptor channels

VII. Role of GPCRs in the Regulation of Electrical Activity
A. Stimulation of electrical activity by GPCRs
B. Inhibition of electrical activity by GPCRs

VIII. Calcium-Mobilizing Receptors and Electrical Activity
A. The dynamics of Ca2+ release
B. Calcium mobilization and secretion

IX. Summary

The pituitary gland is composed of two embryonically, anatomically, and functionally distinct entities, the neurohypophysis and the adenohypophysis. The neuro-

Abbreviations: AC, Adenylyl cyclase; AMPA, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid; AP, action potential; AVP, arginine vasopressin; BK, calcium-activated big conductance K+ (channels); [Ca2+]ER, Ca2+ concentration in the ER; [Ca2+]i, intracellular calcium concentration; CaCC, calcium-activated chloride (channels); CaV, voltage-gated calcium (channels); [Cl−]i, intracellular chloride concentration; CNS, central nervous system; DAG, diacylglycerol; EAG, ether-a-go-go; ER, endoplasmic reticulum; erg, ergot; ET, endothelin; GABA, γ-aminobutyric acid; GEF, guanine nucleotide exchange factor; GluR, glutamate receptor; GPCR, G protein-coupled receptor; α-GSSG, glutathione peroxide; α-subunit; HCN, hyperpolarization-activated and cyclicnucleotide-modulated (channels); 5-HT, 5-hydroxytryptamine (serotonin); Ih, hyperpolarization-activated current; IP3, inositol 1,4,5-trisphosphate; IP3R, IP3 receptor; Kv, voltage-gated potassium (channels); Kv, inwardly rectifying K+ (channels); Kv, voltage-gated potassium (channels); Na+, background sodium (channels); nAChR, nicotinic acetylcholine receptors; Na+, voltage-gated sodium (channels); Na+, Na+ channel-like protein; NMDA, N-methyl-D-aspartate; PACAP, pituitary adenyl cyclase-activating peptide; PDE, phosphodiesterase; PIP2, phosphatidylinositol 4,5-bisphosphate; PKA, protein kinase A; PKC, protein kinase C; PLC, phospholipase C; PMA, phorbol 12-myristate 13-acetate; POMC, proopiomelanocortin; PRL, prolactin; P2X, ATP-gated (purinergic) receptor (channels); P2Y, ATP-gated (purinergic) receptor (channels); TTX, tetrodotoxin; VIP, vasoactive intestinal polypeptide. Copyright © 2010 by The Endocrine Society doi: 10.1210/er.2010-0005 Received March 3, 2010. Accepted June 2, 2010.
hypophysis includes the posterior pituitary lobe, whereas the adenohypophysis includes the intermediate and anterior pituitary lobes. The posterior lobe is composed of axonal terminals of the hypothalamic magnocellular neurons surrounded by astrocytes, also known as pituicytes. The magnocellular neurons from paraventricular and supraoptic nuclei synthesize vasopressin and oxytocin and transport them to the axonal terminals in the posterior pituitary where they are secreted into the general circulation. The intermediate lobe is populated by melanotrophs, which synthesize and release α-MSH (or intermedins). The anterior pituitary is a heterogeneous gland with multiple cell types that secrete six major peptide hormones necessary for reproduction, lactation, growth, development, metabolic homeostasis, and the response to stress: FSH and LH-producing gonadotrophs, prolactin (PRL)-producing lactotrophs, GH-producing somatotrophs, TSH-producing thyrotrophs, and ACTH-producing corticotrophs. This lobe also contains the non-hormone-producing folliculostellate cells, which are glia-like cells, and endothelial cells that line the capillaries. The adenohypophysis of the fish pituitary is directly innervated by hypothalamic neurons, whereas in other vertebrates such a connection was preserved with the intermediate lobe, and whereas in the anterior lobe central nervous system (CNS) neurotransmitters act as releasing and inhibitory hormones delivered through the portal vessels.

This year, we celebrate the 35th anniversary of the discovery that not only neurons and muscle fibers but also endocrine pituitary cells fire action potentials (APs) (1). This pathway, known as the electrical signaling system, is composed of two basic elements: the lipid bilayer and two classes of macromolecular proteins, known as voltage-gated ion channels and ion transporters (2). From the beginning, it was obvious that the role of APs in the propagation of signals along the cells is not of great importance for spherical endocrine cells. However, the discovery of the electrical signaling system in endocrine pituitary cells supported the earlier proposed concept of stimulus secretion coupling (3). The elegance of this concept is that single cells contain all the elements needed for generating Ca^{2+} signals and triggering Ca^{2+}-dependent hormone secretion. The electrical signaling system of endocrine pituitary cells is under intrapituitary and hypothalamic control. Research on chemical signaling within the pituitary cells resulted in the discovery of autocrine and paracrine modes of regulation of pituitary functions and emphasized the role of extracellular ligand-gated ion channels (hereafter called receptor channels) and G protein-coupled receptors (GPCRs) in modulation of electrical activity (4). Communication between pituitary cells through gap junction channels has also been proposed (5).

The main control of spontaneous electrical activity and accompanied voltage-gated Ca^{2+} influx (VGCI) in some pituitary cell types in vivo occurs through hypothalamic releasing and inhibitory hormones acting on pituitary GPCRs signaling through G_{q/11}, and G_{i} signaling pathway. Activated receptors engage variable cellular processes, utilizing G proteins, cyclic nucleotides cAMP and cGMP, and their kinases, protein kinase A (PKA) and protein kinase G. All endocrine pituitary cells have an additional pathway for Ca^{2+} signaling, called the calcium-mobilizing pathway. This pathway is triggered by activation of G_{q/11} coupled receptors and some tyrosine kinase receptors, leading to inositol 1,4,5-trisphosphate (IP_{3}) and diacylglycerol (DAG) production, IP_{3}-mediated Ca^{2+} release from the endoplasmic reticulum (ER), and activation of protein kinase C (PKC) and other signaling pathways. Activated GPCRs also cause a very complex and cell type-specific pattern of changes in the spontaneous firing of APs (6).

The pioneering work on pituitary cell excitability has been summarized in several reviews. The focus in the review by Ozawa and Sand (7) was on voltage-gated ion channels expressed in the pituitary, the effects of TRH on electrical activity and Ca^{2+} signaling, and the characterization of electrical properties of several cell lines. Two subsequent reviews were focused on the plasma membrane and ER Ca^{2+} channels that contribute to Ca^{2+} signaling in pituitary cells (6, 8). Calcium signaling pathways of endocrine pituitary cells have also been reviewed (9). The roles of GPCR-triggered intracellular messengers in hormone secretion were also studied in great detail in (10). Since then, several hundred experimental and tens of theoretical studies have been published describing the structural and functional properties of ion channels expressed in pituitary cells and their roles in spontaneous and receptor-controlled electrical activity, Ca^{2+} signaling, and secretion. In this review, we summarize novel findings on expression, signaling functions, and regulation of ion channels in endocrine pituitary cells.

II. Pituitary Cell Types

The adenohypophysis and neurohypophysis develop from two distinct embryological sources. During craniofacial development, separation of the neuroepithelium (which will become the brain) and the surface ectoderm (which will become the oral epithelium) occurs everywhere except in the middle region forming Rathke’s pouch. The neurohypophysis is derived from the neuroepithelium and originates at the base of the diencephalons. Rathke’s pouch separates from the oral epithelium and forms the adenohypophysis (Fig. 1, inset). Initially, Rathke’s pouch forms a closed epithelial structure with the lumen. Soon after, the
cells from the ventral side of the pouch leave and proliferate to form the nascent anterior lobe, whereas a more limited development of the dorsal wall gives rise to the intermediate lobe. Proliferation is accompanied by the epithelium-mesenchyme transition, with the pituitary-specific transcription factors, Pit-1 (also called Pou1f1) and prophet of Pit-1 (Prop-1) playing important roles in this transition. This stage is closely associated with the initiation of the cell differentiation program (11).

It appears that cell-to-cell contact with the primordial neuroepithelium of the ventral hypothalamus is a critical factor in the differentiation of anterior pituitary cells. The hormone-secreting cells differentiate in a temporal- and spatial-specific fashion under the influence of various transcriptional factors (Fig. 1). In mouse, the expression of the glycoprotein hormone α-subunit (α-GSU) gene is the earliest marker of anterior pituitary differentiation, occurring at embryonic d 11.5 of gestation. Thyrotrophs, somatotrophs, and lactotrophs arise through a common cell lineage determined by Prop-1 and Pit-1 (also known as Pou1f1) transcriptional factors, and mutations of these genes are a cause of combined pituitary hormone deficiency of GH, PRL, and TSH. Terminal differentiation of corticotrophs and melanotrophs is dependent on the T-box transcriptional factor Tbx-19, also known as Tpit. In contrast, Tbx-19 is a negative regulator of the gonadotrophic and Pit-1-independent rostral type of thyrotrophs. Steroidogenic factor-1 and the zinc finger transcriptional factor GATA-2 appear to be important positive regulators of gonadotroph differentiation (11–14). Consistent with this developmental pattern, lactotrophs, gonadotrophs, somatotrophs, and thyrotrophs express neurofilaments NS-68, whereas cells of proopiomelanocortin (POMC) lineage lack the expression of this protein (15, 16).

**A. POMC-producing cells**

The POMC gene is highly expressed in corticotrophs and melanotrophs and is transcribed by the pituitary promoter P1 to an approximately 1200 nucleotide POMC mRNA transcript. In mammals, POMC is posttranslationally modulated by intracellular proteolytic cleavage into an N-terminal peptide, ACTH (1-39) and β-lipotropic hormone in corticotrophs, and into α-MSH, γ-lipotropic hormone, and β-endorphin in melanotrophs. This specific processing of POMC is due to differential expression of prohormone convertases in the two cell types. Corticotrophs only express prohormone convertase-1, which cleaves at a limited number of sites on the POMC prohormone, whereas melanotrophs express prohormone convertase-1 and -2 and more extensively cleave POMC at a number of sites during biosynthesis (17, 18).

Corticotrophs are the first anterior pituitary cells to differentiate during embryonic development; they are derived from the intermediate pituitary but are scattered throughout the anterior lobe in adult animals (Fig. 1). It has been reported that these cells comprise 2 to 15% of AP cells in rats. The large range could reflect age and sex, and also the methods used for identification. The main control of ACTH release is mediated by CRH, which is secreted by paraventricular neurons that project to the median emi-
nence and release CRH into the hypophyseal portal system. In corticotrophs, CRH binds to G\textsubscript{\alpha}\textsubscript{11}-coupled CRH receptors and facilitates spontaneous electrical activity and ACTH release. In addition to CRH and the CRH family of peptides (urocortin 1-3), arginine vasopressin (AVP) directly stimulates ACTH release and acts in synergy with CRH to potentiate ACTH release (19). Glucocorticoid receptors are expressed in corticotrophs and CRH neurons and contribute to negative feedback actions of glucocorticoids on ACTH secretion (20). There is one corticotroph mouse cell line available, called AtT-20 cells. Like corticotrophs in primary culture, AtT-20 cells synthesize POMC and have been extensively used to study the processing of POMC. These cells can also package ACTH into secretory vesicles and were originally used to define the constitutive and regulated secretory pathways. These cells express glucocorticoid, somatostatin, IL-1, dopamine, histamine H\textsubscript{3}, and muscarinic cholinergic receptors (21).

Melanotrophs are the only secretory cells present in the intermediate lobe and account for more than 95% of the cells found in this lobe. In contrast to the anterior pituitary, which is richly vascularized, the intermediate lobe contains very few blood vessels but is supplied by nerve fibers originating from the hypothalamus. Mammalian melanotrophs are electrically excitable cells, and spontaneous electrical activity is sufficient to trigger release of POMC-derived peptides. Such secretion is primarily regulated by dopaminergic neurons that originate in the mediobasal hypothalamus and directly innervate the intermediate lobe. Dopamine tonically inhibits the synthesis and release of POMC peptides by activating D\textsubscript{2} receptors, leading to inhibition of electrical activity and Ca\textsuperscript{2+} signaling. Connections between dopamine-secreting neurons and the intermediate lobe in rats are established during the first postnatal week (22). Mammalian melanotrophs also express GPCRs for \gamma-aminobutyric acid (GABA) (23), prostaglandin E\textsubscript{2} (24), and serotonin (5-hydroxytryptamine; 5-HT) (25), whereas frog melanotrophs also express receptors for TRH, neuropeptide Y (26), acetylcholine (27), and adenosine (28). Cells from the melanotroph cell line mIL39 express POMC and dopamine D\textsubscript{2} receptors and thus could represent a good cell model for studies on dopaminergic regulation of melanotroph functions (29).

**B. Heterodimeric glucoprotein-producing cells**

Thyrotrophs and gonadotrophs express the 92-amino acid-long \alpha-GSU, which is needed for the formation of TSH, FSH, and LH (and also chorionic gonadotropin) heterodimers with hormone-specific \beta-subunits. Thyrotrophs are the smallest subpopulation of anterior pituitary cells, representing less than 10% of cells in the gland, and are regionally localized within the anteromedial and lateral portions of the gland. The \beta-TSH subunit containing 110 amino acids is unique for thyrotrophs and confers specificity of biological actions. TSH is packed into secretory vesicles, which are small (50–100 nm in diameter). The hypothalamic control of thyrotrophs is mediated by TRH, which is released by neurons localized in the hypothalamic paraventricular nucleus. TRH acts as an agonist for G\textsubscript{\alpha}\textsubscript{11}-coupled TRH receptors expressed in thyrotrophs and lactotrophs. In thyrotrophs, TRH stimulates TSH release, as well as the transcription of both \alpha- and \beta-subunits, whereas T\textsubscript{4} and T\textsubscript{3} suppress transcription. The feedback regulation occurs primarily in the pituitary (30). Somatostatin and dopamine have small suppressive effects in TSH-secreting tumors. Secretion of TRH is also controlled by numerous autocrine and paracrine factors, including endothelins (ETs) acting on ET\textsubscript{A} receptors (31). There are two thyrotroph cell lines: \alpha-TSH cells, secreting the \alpha-subunit and missing some of the Pit-1 transcript isoforms; and T\alpha-T-1 cells, expressing both \alpha- and \beta-subunits and the transcriptional factor Pit-1 (21).

Gonadotrophs constitute about 10–15% of the anterior pituitary cell population and are localized throughout the anterior lobe, frequently adjoining lactotrophs. They secrete the gonadotropins LH and FSH, which are packed in secretory vesicles of about 200 and 300 nm in diameter, respectively. The decapetide GnRH is the main agonist for these cells. GnRH-secreting neurons are dispersed within the mediobasal hypothalamus and preoptic areas, but organize functionally as a pulse generator, delivering GnRH in portal blood every 30 min in rodents; this release is influenced by numerous factors, including the age and gender of the animals, and the stage of the estrous cycle (32). In gonadotrophs, GnRH binds to G\textsubscript{\alpha}\textsubscript{11}-coupled GnRH receptors, leading to the release of both LH and FSH in a pulsatile manner (33). GnRH also affects FSH\beta and LH\beta transcription (34, 35). In addition to GnRH receptors, gonadotrophs also express functional receptors for pituitary adenylyl cyclase-activating peptide (PACAP) (36), ETs (37), AVP (38), and substance P (39), which contribute to gonadotropin synthesis and secretion. Inhibins, activins, and follistatin are important regulators of gonadotropin synthesis and secretion. Sex steroids exert negative feedback on gonadotropin release via GnRH neurons and directly at the pituitary level; estradiol can also exert a positive action on pituitary gonadotrophs (40, 41).

There are four mouse cell lines developed by Mellon and colleagues (42, 43): \alphaT1-1, \alphaT3-1, L\betaT2, and L\betaT4. The \alphaT1-1 is not a gonadotroph cell line but rather is an anterior pituitary precursor cell line that expresses \alpha-GSU and probably represents a progenitor for the thyrotroph and gonadotroph lineages. The \alphaT3-1 cell type represents a later stage in cell differentiation, corresponding to the early embryonic gonadotroph cell lineage. It expresses...
GnRH receptors and α-GSU, which is secreted in a constitutive manner, but does not express LHβ and FSHβ. These cells were used to clone the GnRH receptor and were extensively used for studies on GnRH- and PACAP-dependent signaling and gene regulation as well as the roles of various homeobox genes in pituitary cell differentiation. The LβT2 and LβT4 cells have more mature gonadotroph phenotypes because they express LHβ and FSHβ-subunits, as well as steroidogenic factor-1 and GnRH receptor (for references, see Ref. 21).

C. GH- and PRL-producing cells

A fraction of anterior pituitary cells, known as mammomatotrophs, secrete both GH and PRL and are probably transitional cells capable of becoming somatotrophs or lactotrophs (Fig. 1). Somatotrophs are the most common cell type in the anterior pituitary, representing up to 50% of cells, and are localized predominantly in the lateral portions of the anterior lobe. They synthesize GH, a single-chain polypeptide containing 191 amino acids, which is packed into secretory vesicles of variable sizes. Two hypothalamic neuropeptides, GHRH and somatostatin, play major roles in the control of GH synthesis and release. GHRH is a 44-amino acid peptide secreted by neurons in the arcuate nucleus of the hypothalamus that acts as a native agonist for GHRH receptors. These receptors are coupled to the G_s signaling pathway (44). Somatostatin is a 14-amino acid neuropeptide that is secreted by neurons in the periventricular nucleus of the hypothalamus and is delivered to pituitary cells by a portal vascular system. It binds to receptors coupled to the G_{i/o} signaling pathway (45). These cells also express receptors for ghrelin (46), PACAP (47), and ETs (48).

PRL is a single-chain protein of 198 amino acids that is similar in structure to GH. It is synthesized and released by lactotrophs, which account for 10–25% of pituitary cells. This is a nonhomogeneous group of cells (some have large and irregular dense-core vesicles, and others have small round vesicles) that secrete PRL due to spontaneous electrical activity. These cells can be separated into subpopulations based on morphology (49) or density (50). Consistent with the high basal level of PRL secretion, the predominant hypothalamic influence is inhibitory rather than stimulatory and is mediated by dopamine D_2 receptors coupled to the G_{i/o} signaling pathway (51). These cells also express ET_A receptors, which transiently stimulate PRL release, followed by sustained inhibition (52, 53). On the other hand, TRH, angiotensin II, oxytocin, ATP, acetylcholine, and 5-HT stimulate PRL release (54). Estrogens stimulate PRL gene transcription and secretion, and prolonged estrogen treatments lead to an increase in the number of lactotrophs (55).

There are several cell lines that secrete GH, PRL, or both hormones. GH_3 cells produce PRL and GH and express TRH, vasoactive intestinal polypeptide (VIP), and epidermal growth factor receptors but not dopamine D_2 receptors. GH_3C_1 cells are derived from GH_3 cells and produce only PRL. Both cell types have been very useful in the characterization of electrical activity and the channels involved. There are four MtT cell lines; among them, MtT/S cells are pure somatotroph cell lines expressing high levels of GH and GHRH receptors, whereas MtT/SM are mammomatotrophs. On the other hand, the 235-1 lactotroph cell line has the PRL gene, but not the GH gene. These cells do not express TRH and D_2 receptors, but secrete PRL in a Ca^{2+}-dependent manner. The MMQ cells also secrete PRL only in a Ca^{2+}-dependent manner, but in addition express functional D_2, ET_A, and oxytocin receptors and thus could be used for studies on the receptor-controlled electrical activity and secretion (21).

D. Nonsecretory cells

Folliculostellate cells from the anterior lobe are derived from the neuroectodermal cells and are nonendocrine cells devoid of secretory granules. These cells express the neuronal marker S-100 protein and glial fibrillary acidic protein, reflecting their neuroectodermal origin. It has also been suggested that folliculostellate cells develop from marginal layers of the pars tuberalis and pars intermedia. Although they represent only 5–10% of the anterior pituitary cells and are sparsely distributed within the gland, folliculostellate cells make a complex three-dimensional anatomical network extending over the whole gland. It has been suggested that these cells express gap junction channels and play important roles in intercellular communication (56, 57). There are several folliculostellate cell lines, including human PDFS cells, mouse TtT/GF and Tpit/F cells, and rat FS/D1 h cells. Like native cells, these immortalized cells also express muscarinic acetylcholine, β-adrenergic and PACAP receptors positively coupled to cAMP production, and Ca^{2+}-mobilizing angiotensin II receptors (21).

Pituicytes constitute approximately 30% of the posterior pituitary volume, and are the only resident cells. It is well established that neurosecretory axons are capable of making synaptoid contacts with pituicytes. GABA-containing axons also terminate in synaptoid contacts either on pituicytes or the neurosecretory axons. Other neurons can make direct synaptoid contacts with pituicytes. These cells express receptors for GABA, as well as for AVP, κ-opioid, nucleotides, atrial natriuretic peptide, 5-HT, bradykinin, angiotensin II, and ETs. Cultured pituicytes express several Ca^{2+}-mobilizing receptors, the activation of which leads to the generation of Ca^{2+}-waves, either through gap junctions that are expressed in these cells or via the release of ATP and activation of purinergic receptors. Pituicytes also express IL-1 receptors and release IL-6. Such a complex expression pattern of receptors indicates that released neurohormones can act on
pituicytes (58). The physiological role of activated pituicytes is unknown, but they could trigger release of various neuroactive substances and influence the release of AVP and oxytocin. For example, it is known that ATP released by pituicytes activates purinergic receptor channels expressed in vasopressinergic terminals and stimulates hormone release (59, 60).

III. Ion Channels Expressed in Pituitary Cells

A. Voltage-gated channels

Functional analysis, homology cloning, and the sequencing of genomes of several species have revealed that voltage-gated channels are one of the largest groups of signal transduction proteins. These channels have been classified into two major subgroups: a superfamily of more than 140 members of voltage-gated cation channels that share structural similarity; and a small family of structurally different voltage-gated chloride channels. The superfamily of voltage-gated channels includes sodium (Na⁺) (Fig. 2), calcium (Ca²⁺) (Figs. 2 and 3), and potassium (K⁺) channels (Figs. 4–6), as well as numerous less selective channels. These channels are composed of the pore-forming α-subunits and auxiliary subunits. The α-subunits of Na⁺ channels and Ca²⁺ channels have similar amino acid sequences and folding, whereas the pore-forming subunit of K⁺-channels is smaller, but with obvious homology to Na⁺ and Ca²⁺ channels. In cation-selective channels, the part of the pore known as the ionic selectivity filter is able to distinguish among Na⁺, Ca²⁺, and K⁺. The majority of voltage-gated channels contain voltage sensors, charged transmembrane (TM) helices or segments (S) that sense the electrical field in the membrane and drive conformation changes, leading to opening and closing of the gates near the mouth of the pore (61).

Combined pharmacological, electrophysiological, and molecular biology experiments have revealed that there is a high diversity of K⁺ channels, whereas Ca²⁺ and Na⁺ channels are less diverse. Among voltage-gated channels, the structures of inwardly rectifying K⁺ (Kᵢ) channels and bacterial K⁺ channels, known as KcsA, are the simplest. The α-subunit of these channels is a tetramer, each monomer of which contains two TM segments (S), with a reentry P loop in between (Fig. 4). The primordial members of the voltage-gated cationic channel superfamily were probably 2TM domain K⁺ channels. The K₂P channels are known as leak K⁺ channels, and these channels have two P loops and 4TM domains, a topology similar to a tandem fusion of two Kᵢ channels. These channels do not have a voltage sensor. The Kᵢ channel α-subunit monomer contains the two TM segments found in Kᵢ channels plus an additional four TM segments (Fig 5). Cyclic nucleotide-gated (CNG) channels, hyperpolarization-activated and cyclic nucleotide-modulated (HCN) channels, transient receptor-potential (TRP) channels, and some members of Ca²⁺-activated K⁺ channels (KCa) also have this type of architecture (Figs. 6 and 7). Like Kᵢ channels, all 6TM domain channels are homo- or heterotetramers of principal subunits, fre-

FIG. 2. Na and Ca channels. Top, Structural TM folding model of Na and Ca channels. In this and the following models, α-helices are illustrated as cylinders and the extracellular and intracellular chains of amino acids as continuous lines. The positively charged 4S domain illustrates the voltage sensor, and the SS and S6 domains contribute to the formation of the channel pore. Bottom, TTX and saxitoxin (SAX) sensitivity of Na-α subunits. Rectangles show the subunits whose presence was identified in pituitary cells at the mRNA level. TTX-sensitive Na currents have been identified in all endocrine pituitary cells.

FIG. 3. Classification of Ca channel α-subunits. HVA, High-voltage activated; LVA, low-voltage activated; DHP, dihydropyridines; IVA, w-agatoxin; GVIA, w-conotoxin; SNT, SNK-482. HVA and LVA currents have been detected in all endocrine pituitary cells. Rectangles indicate the mRNA transcripts for Ca-α subunits in pituitary cells. Immunocytochemical studies showed the presence of Ca,1.1, 1.2, 1.3, 2.1, 2.2, 2.3, and 3.1-α subunits in these cells.
quently associated with auxiliary β-subunits. The 6TM domain of these channels is doubled in the two pore channels and quadrupled in Nav and Cav channels. In all of these channels, the 4S in the 6TM domain serves as a voltage sensor, whereas the pore loop between 5S and 6S in each domain determines ion conductance and selectivity (61, 62).

1. Voltage-gated Na\(^{+}\) channels

Mammals express nine genes for the Na\(^{+}\) channel α-subunit, termed Na\(_1\).1–Na\(_1\).9, and closely related Na\(^{+}\) channel-like proteins (Nax) with approximately 50% structure similarity with Nav\(_1\) channels. The α-subunit contains four homologous domains, each consisting of a 6TM domain and a reentry P loop between 5S and 6S, which contains the tetrodotoxin (TTX) binding site, a voltage gate and sensor, and contains the sites for phos-

FIG. 4. K\(_\text{ir}\) channels play important roles in the control of resting membrane potential and agonist-induced inhibition of spontaneous electrical activity in pituitary cells. Left, Structural TM folding model of K\(_\text{ir}\) channels. Right, The 15 known members of K\(_\text{ir}\) channels are divided into three groups, based on their regulation. Rectangles indicate K\(_\text{ir}\)-α subunits identified in pituitary cells. The presence of K\(_3\).3.1 and 3.2 has also been confirmed by Western blot analysis.

FIG. 5. K\(_\text{v}\) channels reduce excitation in pituitary cells. Top, Structural TM folding model of K\(_\text{v}\) channels (left) and tetrameric organization of α-subunits (right). Bottom, Families of K\(_\text{v}\) channels. Rectangles indicate K\(_\text{v}\)-α-subunits for which mRNAs were identified in pituitary cells.

FIG. 6. Two types of K\(_{Ca}\) channels are expressed in pituitary cells. Top left, SK channels have similar TM organization as K\(_\text{v}\) channels, but are not voltage-regulated. Top right, BK (maxi) channels have an additional TM domain (S0) and are regulated by both voltage and calcium. Bottom, Phylogenetic tree for K\(_{Ca}\) channels. The K\(_{Ca}\).1.1 mRNA transcript was found in pituitary cells, and the presence of SK and BK currents in pituitary cells was confirmed using specific blockers of these channels.

FIG. 7. Cyclic nucleotide-modulated channels are nonselective cation channels. Top, Structural TM folding model of CNG channels and HCN channels. Bottom, Phylogenetic tree of CNG and HCN channels. The mRNA transcripts for all four HCN α-subunits and CNGA1 α-subunit were identified in pituitary cells, as well as the functional HCN current.
phorylation by protein kinases on the intracellular surface. Na\textsubscript{v} 1.5, 1.8, and 1.9 are TTX insensitive (Fig. 2). Four auxiliary subunits have been identified so far, termed Na\textsubscript{v}β\textsubscript{1}, Na\textsubscript{v}β\textsubscript{2}, Na\textsubscript{v}β\textsubscript{3}, and Na\textsubscript{v}β\textsubscript{4}. They belong to a single family of proteins, which interact with different α-subunits and alter their physiological properties. The main function of Na\textsubscript{v} channels is to depolarize cells and generate the upstroke of the AP, controlling the firing amplitude in excitable cells, including nerve, muscle, and neuroendocrine cell types. In some cells, these channels are solely responsible for the rapid and regenerative upstroke of an AP. In others, they act in conjunction with Ca\textsuperscript{2+} channels to depolarize cells. Na\textsubscript{v} channels are also expressed in non-excitable cells at a lower level, where their physiological role is unclear (63).

The expression of TTX-sensitive and -insensitive Na\textsubscript{v} channels has been extensively studied in endocrine pituitary cells. Electrophysiological experiments revealed that both freshly dispersed and cultured melanotrophs express functional channels composed of TTX-sensitive and TTX-resistant components (64, 65). Single-cell Ca\textsuperscript{2+} measurements further indicated the presence of functional Na\textsubscript{v} channels in frog melanotrophs (66, 67). The TTX-sensitive current has also been identified in rat (68–70), mouse (71), ovine (72, 73), and fish (74, 75) gonadotrophs, as well as in α\textsubscript{T3}-1 mouse gonadotrophs (76, 77). The presence of functional Na\textsubscript{v} channels in gonadotrophs was recently confirmed using mice pituitaries with genetically labeled gonadotrophs (78). Rat lactotrophs (79, 80), somatotrophs (70), corticotrophs (68), and GH\textsubscript{3} cells (81), and fish lactotrophs (82) also express Na\textsubscript{v} channels. GH\textsubscript{3} cells were frequently used as a cell model to study the gating properties of Na\textsubscript{v} channels (83, 84). Thus, it is reasonable to conclude that Na\textsubscript{v} channels are native to all secretory pituitary cells.

There has also been progress in identifying the mRNA transcripts for Na\textsubscript{v} subunits in pituitary cells in various physiological conditions influencing the expression of these channels. Rat melanotrophs express mRNA transcripts of seven α-subunits, including the TTX-insensitive Na\textsubscript{v}1.8 and 1.9 subunit mRNAs and β\textsubscript{1} and β\textsubscript{2} auxiliary subunit mRNAs (65). The mRNA transcripts for the α-subunit of Na\textsubscript{v}1.1, 1.2, Na\textsubscript{v}1.3, and Na\textsubscript{v}1.6, as well as β\textsubscript{1} and β\textsubscript{3} subunits of Na\textsuperscript{+} channels, are present in GH\textsubscript{3} cells (85). The expression of the Na\textsubscript{v}1.7-α-subunit in the rat anterior pituitary was confirmed by in situ hybridization and immunohistochemistry (86). Somatotrophs from GH-green fluorescent protein transgenic mice express mRNA transcripts for Na\textsubscript{v}1.5, 1.8, and 1.9, as well as the TTX-sensitive and TTX-resistant Na\textsuperscript{+} current (87). It appears that the level of Na\textsuperscript{+} channel expression is greater in cultured rat gonadotrophs than in somatotrophs and lactotrophs (70). The level of expression of Na\textsubscript{v} channels in GH\textsubscript{3} cells is inhibited by glucocorticoids (81) and stimulated by long-term activation of the ghrelin-GH secretagogue receptor (88) and activation of L-type Ca\textsuperscript{2+} channels (85). The role of these channels in electrical activity is summarized in Section IV.C.

2. Voltage-gated Ca\textsuperscript{2+} channels

Electrophysiologically, Ca\textsubscript{v} channels are separated into two groups. The first group of channels is known as high-voltage-activated Ca\textsubscript{v} channels because these channels require moderate to strong membrane depolarization to open. Among this group, biophysical and pharmacological studies have identified L-, N-, P/Q-, and R-type Ca\textsuperscript{2+} channels that are distinguished by their single-channel conductance, pharmacology, and metabolic regulation. The second group is known as low-voltage-activated Ca\textsubscript{v} channels because they require less depolarization for activation and subsequent inactivation than high-voltage-activated channels, and a strong membrane hyperpolarization is required to bring them out of steady inactivation. Because of such gating properties, these channels are often referred to as transient or T-type Ca\textsubscript{v} channels (Fig. 3). Purification of calcium channels has identified five subunits: a pore-forming large α\textsubscript{1} subunit and four smaller ancillary subunits: α\textsubscript{2}, β, γ, and δ. Like α-subunits of Na\textsubscript{v} channels, the α\textsubscript{1} subunit of Ca\textsubscript{v} channels consists of four homologous repeats, each consisting of a 6TM domain and a 5P-loop between SS and 6S. In addition to the voltage sensor, gating machinery, and the channel pore, the α\textsubscript{1} subunit also contains most of the known sites of channel regulation by intracellular messengers, drugs, and toxins, including Gβγ domains and multiple PKA phosphorylation sites (89).

Ca\textsubscript{v} channels serve two major functions in cells: electrogenic and regulatory. In some neurons and many neuroendocrine cells, these channels give rise to APs in the same way as Na\textsubscript{v} channels, although typically with slower kinetics and lower amplitude. In other neurons, Ca\textsubscript{v} channels shape the Na\textsuperscript{+}-dependent APs. The regulatory function of these channels is based on Ca\textsuperscript{2+} influx during the transient depolarization, which acts as an intracellular (second) messenger controlling a variety of cellular functions. During a short-term depolarization, Ca\textsuperscript{2+} entry through T-type channels constitutes a disproportionately large fraction of the total Ca\textsuperscript{2+} entry (90). Because T-type Ca\textsubscript{v} channels exhibit rapid and complete voltage-dependent inactivation, however, they are unlikely candidates to promote Ca\textsuperscript{2+} influx of sufficient amplitude to generate global Ca\textsuperscript{2+} signals in neuroendocrine cells firing slow APs. The major function of these channels is electrogenic; at the resting potential, these channels depolarize cells to the threshold level for a Na\textsuperscript{+} or Ca\textsuperscript{2+} spike. In contrast,
high-voltage-activated channels inactivate incompletely and help to keep the cells depolarized for a prolonged period. Such APs increase intracellular calcium concentration ([Ca^{2+}]_{i}) of sufficient amplitude to trigger Ca^{2+}-dependent processes (89, 91).

The functional expression of both inactivating and noninactivating Ca_{v} currents is well documented in rat (68–70, 92), ovine (93), and fish (75) gonadotrophs, as well as in genetically labeled mouse gonadotrophs (78) and αT-3 immortalized mouse gonadotrophs (76). These currents are also present in somatotrophs and lactotrophs (70, 94) and GH cells (95–97). In GH_{3} cells, there are multiple conductance levels of the L-type Ca_{v} channels (98), and estrogens stimulate the expression of T-type channels (99). In the same preparation of rat anterior pituitary cells, it appears that Ca_{v} channels are more prominent in rat somatotrophs than in lactotrophs and gonadotrophs (70). Within the same subpopulation of cells, the expression of T-type Ca_{v} channels varies, as is well documented for lactotrophs (100). Mouse (101) and rat (102, 103) melanotrophs also express functional Ca_{v} channels. The properties of inactivating Ca_{v} channels are consistent with the expression of T-type Ca^{2+} channels, whereas the noninactivating Ca^{2+} current in pituitary cells is mediated by dihydropyridine-sensitive and -insensitive Ca_{v} channels (69, 92, 96, 104). The prominent expression of T-type Ca^{2+} channels in somatotrophs is reflected by their contribution to the generation of the high-amplitude [Ca^{2+}]_{i} transients in spontaneously active cells, whereas L-type channels are essential to the generation of both spontaneous and agonist-induced electrical activity and Ca^{2+} signaling in pituitary cells (for details, see Section IV).

Progress has been made in the identification of Ca_{v}-α-subunit transcripts present in pituitary cells. The Ca_{v.3.1} and Ca_{v.3.3} mRNAs were detected in GH_{3} cells exhibiting prominent T-type Ca^{2+} current (105). Several pore-forming subunits of Ca_{v} channels are present in GH_{3}/B_{6} pituitary cells and account for the formation of the T-type of Ca_{v}(Ca_{v.3.1}), L-type (Ca_{v.1.1}, 1.2, and 1.3), and P/Q (Ca_{v.2.1}) type currents. The mRNA transcripts for β1, β2, and β3 Ca_{v} subunits were also detected in these cells (95). Immunocytochemical analysis confirmed the expression of Ca_{v.1.2}, 1.3, 2.2, and 3.1 α-subunits in mouse anterior pituitary cells (106). The disulfide-linked α2δ-subunit was cloned from human pituitary (107). An immunocytochemical study also suggested that pituicytes express Ca_{v.1.2}, 2.1, 2.2, 2.3, and 3.1 subunits (108). In GH somatotrophs, ghrelin and GH-releasing peptide-6 enhanced the expression of the Ca_{v.1.3} pore-forming α-subunit (109). In LβT2 gonadotrophs, leptin increases L-type Ca_{v} channel expression and GnRH-stimulated LH release (110). In guinea pigs, estrogen significantly increases the mRNA expression of the Ca_{v.3.1} α-subunit in the pituitary and the hypothalamus, accompanied by an increase in the peak T-current, which could explain the stimulatory effects of estrogens on burst firing (111). This increase is dependent on the expression of estradiol receptor α (112). Because of their enormous physiological relevance (discussed in Section IV.E), it is reasonable to speculate that other hormones and neuropeptides affect the expression of Ca_{v} channels in pituitary cells and other excitable cells.

3. Inwardly rectifying K^{+} channels

The term “inward rectifier” describes the activation of inward current under hyperpolarization, leading to K^{+} influx, and almost no K^{+} efflux under depolarization. Because of these unusual activation properties, these channels are also known as anomalous rectifiers. K_{ir} channels are expressed in numerous tissues, including brain, heart, kidney, endocrine cells, ears, and retina. They participate in the control of resting potential and are closed by a strong depolarization. There are 15 members of this family of channels that are divided into three groups, based on their regulation (Fig. 4). The majority of channel subtypes are “classical” K_{ir} channels that are controlled by intracellular messengers (K_{ir.1}, 2, 4, 5, and 7). On the other hand, K_{ir.3} channels are regulated by G proteins and K_{ir.6} channels by intracellular ATP. The long cytoplasmic pore of these channels plays a critical role for inward rectification and provides the structural basis for modulation of gating by G proteins and phosphatidylinositol 4,5 bisphosphate (PIP_{2}) (113).

G protein-regulated K_{ir} channels are present in endocrine pituitary cells. In rat pituitary lactotrophs, K_{ir} currents are activated by dopamine (114) and ETs (115), whereas in somatotrophs they are activated by somatostatin (116) and ETs (48). AtT-20 corticotrophs also express K_{ir} channels activated by G protein-coupled somatostatin and muscarinic receptors (117–120). The G protein dependence of activation of K_{ir} currents in AtT-20 cells and human GH-secreting adenoma cells was shown by down-regulation of their expression by antisense oligonucleotides (121, 122). Consistent with the G protein-dependent K_{ir} currents, RT-PCR analysis showed the presence of K_{ir} 3.1, 3.2, and 3.4 mRNA transcripts in female rat pituitary cells (123), and K_{ir.3.1}–3.4 mRNA transcripts in GH_{3}/B_{6} mammosomatotrophs (124). The presence of K_{ir.3.1} and 3.2 proteins in AtT-20 cells was also confirmed by Western blot analysis (118). For details on the role of these channels in receptor-controlled electrical activity see Section VII.B.

Other members of this family of channels are also expressed in pituitary cells but were only partially characterized. In GH_{3} cells, constitutively active K_{ir} channels play an important role in the maintenance of the resting
membrane potential (125, 126). These channels are inhibited by activation of the TRH receptor, presumably through their cross-coupling to the Gs signaling pathway (127). TRH also inhibits K\textsubscript{v} channels in lactotrophs from lactating rats (128). In ovine somatotrophs, GH-releasing peptide-2 reduces K\textsubscript{v} current via the PKA-dependent signaling pathway (129). The presence of ATP-sensitive K\textsuperscript{+} modifier subunits (Kv5, 6, 8, and 9), further increases the maintenance of the resting potential (132). Channels also have inward-rectifying properties and contribute to the functional diversity of this group. The coding regions of Kv3, 4, 6, 7, 9, 10, and 11 gene families are made up of several exons that are alternatively spliced, leading to a further increase in functional diversity (133). It is likely that Kv1.4, 3.3, 3.4, 4.1, 4.2, and 4.3 contribute to the formation of A-type channels (133, 134). The M-channel is made up of several subunits from the K\textsubscript{v}7 (KCNQ) family of \textalpha{}-subunits, and the voltage-dependent activity of this channel is modulated by PIP\textsubscript{2} (135). The members of Kv1, 2, and 3 contribute to the formation of the fast activating delayed rectifier, and the members of Kv10, 11, and 12 generate various EAG channels, including the erg1 channel expressed in cardiac tissues (133).

Qualitative RT-PCR analysis revealed that GH\textsubscript{3}/B\textsubscript{6} pituitary cells express mRNA transcripts for Kv1.3, 1.4, 1.6, 2.1, 2.2, 3.2, 3.4, 4.1, 4.2, 4.3, 6.1, 7.1, 7.2, 7.3, 10.1, 11.1–11.3, and 12.1–12.3 (124). Others reported about the expression of Kv1.5 in pituitary cells (136). There is also evidence that K\textsubscript{v} expression is modulated by hormonal status. Glucocorticoid injection in vivo increases 8-fold the amount of K\textsubscript{v}1.5 mRNA in rat pituitaries (137). The increase in K\textsubscript{v}1.5 (but not K\textsubscript{v}1.4) expression is associated with an increase in a nonactivating component of the K\textsubscript{v} current in GH\textsubscript{3} cells (138). Interestingly, both depolarization and TRH application reduces K\textsubscript{v}1.5 expression in GH\textsubscript{3} cells, increasing cell excitability (136, 139). Thus, hormonal and physiological status can dynamically alter the excitability of pituitary cells on a time scale of hours.

Electrophysiological experiments confirmed the presence of delayed rectifier current in GH cells (96, 140–142) and their regulation by GHRH (143). This current is also present in native rat lactotrophs and somatotrophs (70, 144–146). Mouse \textalpha{}T3-1 gonadotrophs (76), and native goldfish (75), rat (70), and ovine (147) gonadotrophs also express delayed rectifiers, and estrogens transiently increase the expression of these channels (147). Other fish pituitary cell types also have these channels (74, 148). The potential role of delayed rectifier K\textsuperscript{+} channels in electrical activity was examined in various pituitary cells. In GH\textsubscript{3} cells, inhibition of this channel by tetraethylammonium increases the duration of the AP (149) and the amplitude of the spontaneous [Ca\textsuperscript{2+}]\textsubscript{i} transients (125), whereas in native rat lactotrophs, tetraethylammonium does not alter the pattern of AP firing (149). In frog melanotrophs, adenosine potentiates the delayed rectifier K\textsuperscript{+} conductance, leading to inhibition of electrical activity (150).

A-type K\textsubscript{v} channels are also expressed in the majority of secretory pituitary cells. They were identified in GH\textsubscript{3} mammosomatotrophs (142) and \textalpha{}T3-1 gonadotrophs (76). Native fish (74, 75, 148), frog (150), and rat (70, 144, 145) pituitary cells also express A-type K\textsubscript{v} channels. In frog melanotrophs, adenosine potentiates these chan-
nals (151). Direct comparison of rat lactotrophs, somatotrophs, and gonadotrophs indicates that the expression level of the A-type Kv channels is much higher in lactotrophs and gonadotrophs than in somatotrophs (70). In contrast, high levels of expression of these channels have been observed in ovine somatotrophs and may contribute to the regulation of AP firing and hormone secretion (152). The participation of the A-type K^+ channel in regulating AP firing in other anterior pituitary cell types is unclear. In rat lactotrophs, for example, they do not appear to participate in AP generation (149), which may be due to their prominent inactivation at the resting membrane potential in these cells.

The M-type K^+ current has also been identified in lactotrophs, where it is inhibited by TRH, leading to an increase in the firing frequency during sustained stimulation (153). The M-type current resembles one generated by erg channels (154), which are also expressed in GH mammosomatotrophs and native rat lactotrophs (155). Blockade of erg channels by E-4031 causes depolarization of the membrane potential of about 5 mV, facilitating the release of PRL (156). In our hands, E-4031 did not alter basal PRL release in perfused pituitary cells measured by RIA, in contrast to Ca^2+ in concentrations that inhibit Kv channels (157). Erg channels are inhibited by TRH through an unidentified intracellular messenger (155, 158). TRH was also able to inhibit erg1, erg2, and erg3 channels, as well as human erg, when expressed in HEK293 cells (159). Erg currents are also expressed in MMQ lactotrophs, and their blockade facilitates AP firing and PRL secretion (160). Functional channels were identified in mouse gonadotrophs, and GnRH inhibits these channels through a still uncharacterized signal cascade (161). Human PRL-secreting tumors also express human erg, and they are functionally coupled to PRL secretion (162).

5. Calcium-activated K^+ channels

K_{Ca} are the third major group of K^+ channels and are composed of two families (Fig. 6). One family of these channels includes three small-conductance [calcium-activated small K^+ channels (SK)] channels (K_{Ca}2.1, 2.2, and 2.3) and one intermediate-conductance channel (K_{Ca}3.1). Splice variants have also been identified for SK channel genes. These channels have a general topology similar to that of Kv channels but show little voltage dependence and are activated by Ca^{2+} entering through Ca_v channels and released from intracellular stores, but they are not tightly coupled to Ca^{2+} channels. SK channels use calmodulin constitutively bound to the C terminus of each α-subunit as a high-affinity Ca^{2+} sensor (163, 164). The high-conductance K^+ [calcium-activated big K^+ channels (BK)] channels represent the other family of K_{Ca} channels, only distantly related to SK and intermediate-conductance channels. BK channels are composed of four pore-forming K_{Ca}1.1 α-subunits that share the 6TM topology of Kv channels but contain an additional TM segment at the N terminus, termed S0 (Fig. 6). Alternative splicing of the RNA produces numerous transcripts, resulting in channels exhibiting distinct functional properties. BK channels are activated by voltage, and their open probability is modulated by Ca^{2+}. In contrast to SK channels, BK channels form macromolecular complexes with Ca_v channels and establish a prototypic Ca^{2+} nanodomain. This provides an effective mechanism for control of activity of these channels by Ca^{2+} influx through Ca_v channels. Calcium activation of BK channels is not dependent on calmodulin but is mediated by cation binding sites in the C termini of channel α-subunits (165). The gating properties of these channels are influenced by auxiliary β-subunits. BK channels are blocked by charybdotoxin, iberiotoxin, and paxilline (164, 166).

There are several reasons why K_{Ca} channels are incorporated into the Ca^{2+} signaling pathway. The colocalization of BK and Ca_v channels facilitates spike repolarization, which limits AP-driven Ca^{2+} influx. BK channel activation can also influence the frequency of AP-driven [Ca^{2+}], transients by slowing the pacemaker depolarization. Activation of these channels may relieve the steady inactivation of Na_v and Ca_v channels, which stimulates or enhances AP generation in some cells. BK channels may also play a role in the generation of the pseudo-plateau bursting type of electrical activity in pituitary cells (see Section IV.D). Such diverse effects on AP firing probably depend on the type of K_{Ca} channels expressed and the context of other channels (167). Activation of K_{Ca} channels and the resulting membrane hyperpolarization may also serve to synchronize electrical activity and secretion in cell networks and electrical activity and Ca^{2+} release through IP_3 receptors (IP_3Rs) (168). Both BK and SK channels are operative in endocrine pituitary cells. SK currents were initially identified in GH cells and had magnitude of less than 20% that of Ca_v current (169, 170) and about 25% of the hyperpolarizing current triggered by TRH (171). The expression of SK channels is also well documented in rat (172, 173), mouse (174), and ovine gonadotrophs (72), and the level of their expression is dependent on estradiol (71). SK currents are mainly responsible for the oscillatory hyperpolarization triggered by activation of GnRH receptors, leading to periodic Ca^{2+} release from the ER (discussed in Section VIII.A). In these cells, the function of SK channels is facilitated by PKC (175). Corticotrophs also express SK channels that are activated by AVP (176). An additional Ca^{2+}-activated current was observed in rat gonadotrophs and masked by SK channels when the recording was done.
in cells held at −40 mV (177). Other Ca²⁺-sensitive channels, such as Cl⁻ channels, which are expressed in AtT-20 cells (178) and native lactotrophs (179), may also be masked. It has also been suggested that SK channels contribute to the after-spike hyperpolarization in GH cells (96). In GH₃ cells, SK channel activation requires high-frequency firing, prolongation of APs by voltage-dependent K⁺ channel inhibitors, or release of Ca²⁺ from intracellular Ca²⁺ stores (169). In native pituitary cells, VGCI does not activate SK channels (70). We speculated that the SK channels in pituitary cells are in close proximity to intracellular Ca²⁺ release sites and can be activated only by Ca²⁺-mobilizing receptors, sustained VGCI, and/or Ca²⁺-induced Ca²⁺ release (180). In GnRH-secreting neurons, agonist-induced Ca²⁺ mobilization and the concomitant increase in firing frequency are needed to activate SK channels (181).

Single-channel recordings have shown that BK channels are expressed in melanotrophs (182) and lactotrophs (183). Whole cell current recordings confirmed the presence of BK current in rat somatotrophs and lactotrophs, but not in gonadotrophs (70). The relatively high levels of BK channel expression in somatotrophs and lactotrophs should limit AP-driven Ca²⁺ influx compared with that in gonadotrophs, which exhibit the lowest levels of BK channel expression. However, duration of the AP waveform is longer in somatotrophs and lactotrophs (100–500 msec) than in gonadotrophs (10–100 msec) (69, 116, 149, 184). In addition, both the amplitude and duration of the spontaneous, extracellular Ca²⁺-dependent Ca²⁺ transients are greater in somatotrophs and lactotrophs than in gonadotrophs (184–186). It is unlikely that the prolonged duration of AP-driven Ca²⁺ entry in somatotrophs and lactotrophs is due to the inability of AP firing to activate BK channels because short Ca²⁺ influx steps (<25 msec) were sufficient to activate BK channels in both cell types. An atypical role of BK channels in regulating the pattern of spontaneous AP firing and Ca²⁺ signaling in anterior pituitary cells is discussed in detail in Section IV.D. The role of GPCRs in regulation of BK channel activity in endocrine pituitary cells has not been systematically investigated. In one study, the role of G(i/o), coupled ET₄ and dopamine D₂ receptors in activation of these channels in lactotrophs was addressed (183).

The BK-type K⁺ channels are expressed in GH₃ and AtT-20 pituitary cell lines (169, 187–189), and activation of PKC inhibits these channels, which could account for the sustained excitability of pituitary cells during sustained activation of Gαs-coupled GPCRs (190). In GH₃ cells, BK channels contribute to AP repolarization (169), and these cells have been used frequently to study modulation of native BK channels by different compounds (191–193). The mslo transcripts encoding the pore-forming α-subunit of BK channels are robustly expressed in AtT-20 cells, and native channels are not functionally coupled to β-subunits (194). In these cells, glucocorticoids rapidly activate BK channels via a nongenomic mechanism (195). Glucocorticoids applied for 2 h also promote de novo BK mRNA and protein synthesis and antagonize PKA- and PKC-dependent inhibition of BK channels (188, 196). Further studies revealed that glucocorticoid regulation of BK channels in these cells is mediated by serine/threonine protein phosphatase (197) and that alternative splicing determines sensitivity of BK channels to glucocorticoids and switches their sensitivity to protein phosphorylation (198, 199). The functional effect of such regulation is thought to be to facilitate ACTH release in response to CRH through inhibition of BK channels (188).

Alternative splicing of the BK channel pore-forming α-subunit also occurs in the adrenal and pituitary gland (200–202). Hypophysectomy of rats causes changes in alternative splicing of the α-subunit in the adrenal medulla, which can be reversed by ACTH treatment (202). This splice decision is also regulated in both adrenal and pituitary tissues in stress situations (200, 203). The role of glucocorticoids in the regulation of BK α-subunit alternative splicing in these tissues was confirmed in vitro (200, 201), and the action of glucocorticoids is mediated by both mineralocorticoid and glucocorticoid receptors (200). Female mice genetically deficient in the pore-forming BK subunit respond to restraint stress with reduced ACTH and corticosterone release. It appears that both CRH expression in the paraventricular nucleus and ACTH peptide content in the pituitary were reduced in mice deficient in BK channels (204). Gonadal testosterone also plays a role in the regulation of Slo α-subunit alternative splicing in pituitary cells (205). Stress influences not only the splice decision but also the level of mRNA expression of α-, β₂-, and β₄-subunits (167). The β₂- and β₄-subunits of BK channels are regulated by steroid hormones (206).

6. Cyclic nucleotide-modulated channels

These channels are directly activated by cyclic nucleotides, in contrast to other channels regulated by their protein kinases. Thus, they translate changes in concentrations of cyclic nucleotides to changes in membrane potential. These channels belong to two families: the hyperpolarization-activated and cyclic nucleotide-modulated channels (HCN channels, with resulting current Iₜ, where “h” stands for hyperpolarization) and CNG channels (Fig. 7). As their name indicates, HCN channels are activated by voltage and cyclic nucleotides, whereas CNG channels are virtually voltage-independent and activated by cyclic nucleotide binding. Structurally, these channels belong to the superfamily of K⁺ channels. However, HCN
and CNG channels functionally dissociate from other 6TM domain K+ channels. Their activation does not dampen excitation, but it increases the firing of APs. Such a paradoxical role for channels that structurally belong to the K+-channel family comes from their permeability properties: HCN channels are weakly K+-selective channels, and CNG are practically nonselective cation channels (207).

In mammals, the HCN channel family comprises four subunit isoforms, encoded by four genes, HCN1-4 (Fig. 7). When expressed alone, each subunit forms functional channels, but the native channels are probably organized as heterotetramers. HCN channels are activated by hyperpolarization beyond −60 mV, do not inactivate, and conduct Na+ and K+. In cells expressing these channels, their activation leads to slow depolarization, an action consistent with their equilibrium potential of about −30 mV. HCN channels were identified first in cardiac sinoatrial node cells, and subsequently in a variety of peripheral and central neurons. Their voltage sensitivity is modulated by cAMP. HCN channels serve three principal functions in excitable cells: 1) they contribute to the resting potential; 2) they generate or contribute to the pacemaker depolarization that controls rhythmic activity in spontaneously firing cells; and 3) they compensate for inhibitory postsynaptic potentials. A small fraction of HCN channels are tonically activated at rest, producing the first two functions of these channels (208).

Qualitative RT-PCR analysis suggests that AtT-20 cells express mRNA transcripts for HCN1 (209). GH3 cells express mRNA transcripts for HCN2, HCN3, and HCN4, but not for HCN1 (210). Consistent with these data, electrophysiological experiments confirmed the presence of Ih in GH3 cells (210, 211), AtT-20 cells (209), melanotrophs (150), somatotrophs (211), and lactotrophs (212). The biophysical and pharmacological properties of this current are similar to the Ih current described in neuronal and cardiac cells. This includes the sensitivity to both ZD7288 and Cs+ (209–211) as well as to tramadol (213). In contrast to neuronal and cardiac cells, experiments with GH3 cells showed no effect of elevated cyclic nucleotides on the channel activity in resting cells. Specifically, application of TRH, forskolin, and 8-Br-cAMP does not affect the channel activity in GH3 cells (210, 211). However, inhibition of the basal cAMP production significantly attenuates the Ih current, which fully recovers by the application of 8-Br-cAMP (210). In AtT-20 cells, current is also robustly inhibited by a cAMP antagonist (209). These results suggest that in pituitary cells, Ih is under tonic activation by basal levels of cAMP.

This current is unlikely to play a major role in pacemaking or setting the resting membrane potential in pituitary cells in vitro. First, in GH3 cells these channels operate in the range of potentials negative to −60 mV, with small activation at the resting membrane potential. Second, although the extracellular application of Cs+ and ZD7288 almost completely blocks Ih, it does not stop spontaneous electrical activity or influence the resting membrane potential. However, Ih may limit the excessive hyperpolarization in response to hyperpolarizing stimuli (209–211). Finally, in normal and immortalized GH3 pituitary cells, ZD7288 has Ih-independent effects. This compound induced a rapid increase in the frequency of spontaneous APs and Ca2+ transients in a fraction of cells, which was accompanied by a transient and dose-dependent increase in PRL release in perfused pituitary cells, indicating that channels other than Ih could also be affected by this compound (210, 212). Further studies should be focused on characterization of HCN channels in intact pituitary tissue, including the effects of dopamine and somatostatin on cAMP production, and the role of Ih in pacemaking activity.

In vertebrates, there are six CNG subunits: CNGA1, CNGA2, CNGA3, CNGA4, CNGB1, and CNGB3 (Fig. 7). As with other channels, differential splicing of primary transcripts yields channels of altered structure and behavior. CNGA1-3 subunits can form homomeric channels in heterologous expression systems, and other subunits can coassemble to form functional heteromeric channels. These channels are expressed in olfactory neurons and outer segments of rod and cone photoreceptors, where they play a critical role in sensory transduction. Photoreceptors have a strong preference for cGMP, whereas the olfactory channel is almost equally sensitive to both ligands. The channels are permeable to Na+, K+, and Ca2+, but not to Cl− and other anions. Low levels of mRNA transcripts for these channels are also found in brain, testes, kidney, and heart (207). The mRNA transcripts for rod CNG were also detected in rat pituitary cells by RT-PCR analysis (214) and RNA blot hybridization (215). The zebrafish-specific CNGA5 mRNA and protein transcripts are also expressed in the pituitary (216). Stimulation of cGMP production by nitric oxide donors did not change the pattern of spontaneous VGCl in rat lactotrophs (217), and application of a cell permeable 8-Br-cGMP was also ineffective (218), arguing against the relevance of CNG channels in signaling and secretion. Further studies are required to clarify their expression at the protein level and their potential role in electrical activity and Ca2+ signaling in other endocrine and/or non-secretory pituitary cells.

7. Transient receptor potential channels

TRP channels were initially found in Drosophila, where they contribute to phototransduction. Six protein
families comprise the mammalian TRP superfamily: the “canonical” receptors (TRPCs), the vanilloid receptors (TRPVs), the melastatin receptors (TRPMs), the poly-sistins (TRPPs), the mucolipins (TRPMLs), and the ankyril TM protein 1 (TRPA1). These channels resemble K+ channels in overall structure. However, they show limited conservation of the S4-positive charges and P loop sequences. Assembly of channel subunits as homo- and heterotetramers results in the formation of cation-selective channels. Two members of this superfamily are Ca2+-selective (TRPV5 and TRPV6), and two are monovalent cation selective (TRPM4b and TRPM5); all other channels are relatively nonselective (219). These channels have been studied extensively in numerous tissues, but not in the pituitary gland. The TRPM3 channel mRNA transcripts are present in pituitary cells (220) as well as unidentified member(s) of the TRPC family of channels that are activated by phospholipase C (PLC) (221). The mRNA transcripts for TRPC1, TRPC3, TRPC5, and TRPC7 have also been identified in human pituitary cells (222). There are at least two types of currents present in pituitary cells whose nature is unknown and could be mediated by TRP channels: Ca2+-activated nonselective cationic currents in GH3 cells (223) and gonadotrophs (177), and TTX-insensitive Na+ conductance present in all endocrine pituitary cells (80, 224). It is reasonable to suggest that future electrophysiological investigations in pituitary cells should be focused on this superfamily of channels, especially on members of the TRPC and TRPM families.

B. Chloride channels and transporters

Anion channels are proteins forming pores in biological membranes that allow the passive diffusion of negatively charged ions along their electrochemical gradients. Because all of these channels conduct Cl−, the most abundant anion in organisms, they are often called chloride channels. However, some of these channels may be better conductors of ions other than Cl−. As with cation channels, the most logical classification of Cl− channels is based on their molecular structure, but entire gene families of anion channels remain to be discovered. The most common division of these channels is based on molecular structure and biophysical characteristics and include: voltage-gated chloride channels, ligand-gated (GABA and glycine) chloride channels, calcium-activated chloride (CaCC) channels, high (maxi) conductance channels, the cystic fibrosis TM conductance receptor, and volume-regulated channels (225).

GABA and glycine channels have been studied in pituitary cells and are described in Section VI.A.3. The expression and role of voltage-gated chloride channels in pituitary cells have not been studied. The CLIC6, a member of the intracellular Cl− channel family, was identified in the posterior pituitary (226). Most studies on pituitary cells were focused on CaCC channels. The first report about the Ca2+-dependent Cl− conductance was done in GH3 cells (227). These channels are also present in AtT-20 cells and contribute to the control of APs and VGCI (178). In native lactotrophs, TRH activates CaCC channels in addition to BK and SK channels (228). Depolarization-induced Ca2+ influx and Ca2+ ionophore application also trigger activation of these channels in lactotrophs (179). A large conductance Ca2+-sensitive chloride channel is present in lactotrophs and takes part in the background regulation of the intracellular chloride concentration (229). Hypotonicity also activates CaCC channels in GH4/C1 cells (230).

The common characteristic of K+ and Cl− ions in neurons is their negative equilibrium potential. Activation of channels conducting these ions draws the membrane potential closer to their equilibrium potentials and farther from the threshold for firing. Channels conducting these ions tend to stabilize the membrane potential by setting the resting potential, repolarize and hyperpolarize cells after a depolarizing event, and control the interspike interval. In the majority of cells from adult animals, intracellular chloride concentration ([Cl−]) is low, which permits chloride channels to stabilize the membrane potential of excitable cells. Experiments with AtT-20 cells, however, found that [Cl−]i was between 40 and 50 mM and that activation of CaCC channels by Ca2+ influx during APs tends to maintain the membrane potential at a depolarized level and to enhance VGCI (178). In lactotrophs, [Cl−]i was estimated to be around 60 mM (231). Electroneutral ion transporters, such as the Cl− extruding K+-Cl− cotransporter KCC2 and the Cl− accumulating Na+-K+-2Cl− cotransporter NKCC1, participate actively in maintaining a high [Cl−]i, in the endocrine pituitary cell (232). For further discussion on this subject, see Section VI.A.3.

Elevated basal [Cl−]i may explain the finding that Cl− channel blockers inhibit ACTH release from corticotrophs (233) and provides a rationale for the complex pattern of interactions between Ca2+ and Cl− movements in pituitary lactotrophs (179). It has also been shown that PRL secretion is an osmotically driven process depending on [Cl−]i (234). Additionally, granule fusion recorded by the patch clamp technique is facilitated when the intrapituitary Cl− is elevated (235, 236). Experiments with substitution of Cl− with other ions also confirmed the specific role for this anion in stimulus-secretion coupling (237).

C. Channels expressed in and controlled by the endoplasmic reticulum

The expression of ion channels is not limited to the plasma membrane. Two families of Ca2+ release channels, IP3Rs and ryanodine receptors (RyRs), are predominantly
expressed in the ER/sarcoplasmic reticulum membrane. These channels are structurally and functionally similar. IP$_3$Rs are activated by two classes of plasma-membrane receptors known as Ca$^{2+}$-mobilizing receptors, whereas RyRs provide an effective mechanism for intracellular transduction and translation of electrical signals. The activity of both types of channels is regulated by Ca$^{2+}$, and the variety of Ca$^{2+}$ signaling patterns, including Ca$^{2+}$ sparks and puffs and oscillatory Ca$^{2+}$ waves, depends critically on the [Ca$^{2+}$], dependence of these two families of channels. Activation of IP$_3$Rs leads to stimulation of voltage-insensitive Ca$^{2+}$ channels expressed on the plasma membrane, a process known as capacitative Ca$^{2+}$ entry (238, 239).

1. IP$_3$ receptors

IP$_3$Rs are found in the ER and nuclear membranes of almost all cells. These receptors are composed of four similar subunits that are noncovalently associated to form a four-leaf clover-like structure, the center of which makes the Ca$^{2+}$-selective channel. Each subunit contains approximately 2700 amino acids, with the cytoplasmic N terminus comprising approximately 85% of the protein mass, a hydrophobic region predicted to contain six membrane-spanning helices, and a short cytoplasmic C terminus. Physiologically, activation of IP$_3$Rs is triggered by GPCRs and the plasma membrane receptor tyrosine kinases. Calcium-mobilizing GPCRs activate PLC-β, whereas receptor tyrosine kinases activate PLC-γ. Both enzymes hydrolyze the membrane-associated PIP$_2$ to increase the production of IP$_3$ and DAG. IP$_3$ rapidly diffuses into the cytosol to activate IP$_3$Rs. In addition to IP$_3$, Ca$^{2+}$ plays an important role in the control of permeability of these channels. There are three subtypes of IP$_3$Rs, which exhibit the isoform-specific properties in terms of their sensitivity to IP$_3$ and Ca$^{2+}$. Further diversity of IP$_3$R expression is created by alternative splicing. Most cells express multiple isoforms of IP$_3$Rs, indicating that they have different functions (240).

IP$_3$Rs are commonly expressed in pituitary cells, as indicated by the ability of numerous Ca$^{2+}$-mobilizing agonists, including GnRH (241), TRH (242), AVP and oxytocin (54, 243), angiotensin II (244), ET-1 (245), neurotensin (246), and ATP (247), to trigger an extracellular transduction and translation of electrical signals. The activity of both types of channels is regulated by Ca$^{2+}$, and the variety of Ca$^{2+}$ signaling patterns, including Ca$^{2+}$ sparks and puffs and oscillatory Ca$^{2+}$ waves, depends critically on the [Ca$^{2+}$], dependence of these two families of channels. Activation of IP$_3$Rs leads to stimulation of voltage-insensitive Ca$^{2+}$ channels expressed on the plasma membrane, a process known as capacitative Ca$^{2+}$ entry (238, 239).

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Among pituitary cells, the oscillatory pattern of Ca$^{2+}$ release through IP$_3$Rs is a unique characteristic of gonadotrophs and is not species specific (174, 253). Other native pituitary cells typically release Ca$^{2+}$ from the ER in a nonoscillatory manner (254). Immortalized αT3-1 and LB72 gonadotrophs also release Ca$^{2+}$ in a nonoscillatory manner (42, 255), indicating that the oscillatory response is not established at an early phase of pituitary development. However, it is already established at birth because neonatal gonadotrophs respond to GnRH application with oscillatory Ca$^{2+}$ release (256). The cell-type specificity in the pattern of Ca$^{2+}$ release could be related to the expression of IP$_3$R subtypes, which has not been studied in native pituitary cells. αT3-1 gonadotrophs express all three subtypes of receptors, but the expression of IP$_3$R1 dominates (257). Furthermore, in these cells the sustained desensitization of GnRH action is due to uncoupling of IP$_3$ generation and Ca$^{2+}$ mobilization (258) and down-regulation of IP$_3$R1 (257). Ubiquitination and proteasomal degradation account for down-regulation of endogenous and exogenous IP$_3$R1 in αT3-1 gonadotrophs (259), and this action is triggered by the concerted action of IP$_3$ and Ca$^{2+}$ binding to this receptor (260).

2. Ryanodine receptors

RyRs are the largest known ion channels. Mammalian tissues express three isoforms: RyR1 is expressed predominantly in skeletal muscle; RyR2 is expressed in cardiac muscle; and RyR3 has a wide tissue distribution, including the nonexcitable cells. RyRs are tetramers, with a large N-terminal region forming the head and a C-terminal region that forms the Ca$^{2+}$-selective channel. Intracellular Ca$^{2+}$ is a major regulator of RyRs. The ability of Ca$^{2+}$ to stimulate Ca$^{2+}$ release from the endoplasmic/sarcoplasmic reticulum via RyRs is known as Ca$^{2+}$-induced Ca$^{2+}$ release. This process is of fundamental importance for coordinating the elementary Ca$^{2+}$-release events into Ca$^{2+}$ spikes and waves. Unlike IP$_3$Rs, RyRs can release Ca$^{2+}$ in response to an increase in [Ca$^{2+}$], with no other change in the concentration of second messengers. This is crucial for excitation-contraction coupling. For example, in cardiac cells, Ca$^{2+}$ entry through dihydropyridine-sensitive Ca$_v$ channels activates RyRs to induce a further increase in [Ca$^{2+}$]. In skeletal muscle cells, the dihydropyridine receptors act primarily as voltage sensors to directly activate RyRs in response to membrane depolarization. These
receptors are susceptible to many different modulators. Ryanodine and dantrolene activate and inhibit RyRs, depending on their concentrations, and caffeine is a standard pharmacological tool for the activation of RyRs (261, 262).

Ryanodine-sensitive Ca\(^{2+}\) stores are present in goldfish pituitaries and contribute to agonist-induced Ca\(^{2+}\) signaling and secretion (263). In tilapia fish pituitary cells, however, ryanodine-sensitive Ca\(^{2+}\) stores do not contribute to hypoosmotically-induced PRL release (264). The mRNA transcripts for RyR\(_1\) and RyR\(_3\) are present in AtT-20 cells, and caffeine stimulates ACTH release, presumably by activating these receptors (265). Furthermore, it has been suggested that cADP-ribose is a second messenger in these cells that regulates ACTH secretion by a mechanism dependent on activation of RyRs by extracellular Ca\(^{2+}\) (266). In GH\(_3\) cells, caffeine stimulates Ca\(^{2+}\) release from intracellular pools but also inhibits Ca\(^{2+}\) influx through L-type Ca\(_v\) channels (267). The mRNA transcripts for RyR\(_2\) and RyR\(_3\), but not RyR\(_1\), are present in rat pituitary cells, and ruthenium red treatment, which should block these channels, inhibits GnRH-induced LH release from gonadotrophs (268). In contrast, the ryanodine treatment does not affect GnRH-induced Ca\(^{2+}\) oscillations, suggesting that a ryanodine-sensitive pool does not contribute significantly to IP\(_3\)-dependent Ca\(^{2+}\) oscillations in rat gonadotrophs (249, 269). It is also unlikely that Ca\(^{2+}\) influx through Ca\(_v\) channels is coupled to Ca\(^{2+}\)-induced Ca\(^{2+}\) release in rat somatotrophs (214), lactotrophs, and GH\(_3\) cells (270). A detailed characterization of the expression and function of RyRs in endocrine and nonendocrine pituitary cells is needed.

3. **STIM-controlled Orai channels**

The term “capacitative Ca\(^{2+}\) entry,” by analogy with a capacitor in an electrical circuit, implies that intracellular Ca\(^{2+}\) stores prevent entry when they are charged (filled by Ca\(^{2+}\)) but promote entry as soon as the stored Ca\(^{2+}\) is discharged (released). The similarities in the properties of this entry within different cell types, including excitable cells, suggest a common mechanism. In addition to Ca\(^{2+}\)-mobilizing agonists, capacitative Ca\(^{2+}\) entry can be activated by injection of IP\(_3\) or its nonmetabolized forms into the cell, inhibition of the ER-Ca\(^{2+}\) pumps [sarcoplasmic-ER Ca\(^{2+}\) ATPase (SERCA)] by thapsigargin, discharge of the intracellular content by Ca\(^{2+}\) ionophores, or prolonged incubation of cells in Ca\(^{2+}\)-deficient medium. When heparin, an IP\(_3\)R inhibitor, was injected into the cells, it completely blocked agonist and IP\(_3\)-induced Ca\(^{2+}\) mobilization and capacitative Ca\(^{2+}\) entry. Because depletion of the ER-Ca\(^{2+}\) stores is followed by the influx of Ca\(^{2+}\) into the cell, the channels involved in such influx were termed store-operated Ca\(^{2+}\)-selective plasma-membrane channels. Two proteins have been identified as critical for this pathway: stromal-interacting molecule (STIM) and Orai (271).

Capacitative Ca\(^{2+}\) entry is probably operative in immortalized pituitary cells. It has been suggested that capacitative Ca\(^{2+}\) entry cooperates with Ca\(_v\) channels to generate spontaneous Ca\(^{2+}\) oscillations and the sustained rise in [Ca\(^{2+}\)]\(_i\) in TRH-stimulated GH\(_3\) cells (272). In these cells, thapsigargin and thymol treatment is also accompanied with capacitative Ca\(^{2+}\) influx, which is inhibited by La\(^{3+}\) and SKF 96365 (273–275). In GH\(_4\)C\(_1\) cells, the contribution of capacitative Ca\(^{2+}\) entry to [Ca\(^{2+}\)]\(_i\) is modest compared with the robust Ca\(^{2+}\) influx through Ca\(_v\) channels (276, 277) and is facilitated by loperamide (278). Furthermore, some of the effects could be related to the expression of TRP channels in pituitary cells (222). Thapsigargin also triggers Ca\(^{2+}\) influx in αT3-1 cells, but in these cells GnRH stimulates Ca\(^{2+}\) influx predominantly through L-type Ca\(_v\) channels (279, 280).

At the present time, there is no information about the contribution of capacitative Ca\(^{2+}\) entry in normal pituitary cells. Two lines of evidence, however, argue against their activation upon ER-Ca\(^{2+}\) store depletion. In lactotrophs, activation of \(\alpha\)ET\(_A\) receptors leads to stimulation of Ca\(^{2+}\) release through \(G_{\alpha11}\) coupling accompanied with a sustained inhibition of Ca\(^{2+}\) influx, in contrast to the expected rise in [Ca\(^{2+}\)]\(_i\) (281, 282), suggesting that depletion of the ER-Ca\(^{2+}\) pool does not trigger capacitative Ca\(^{2+}\) influx in these cells. In gonadotrophs, the duration of GnRH-induced Ca\(^{2+}\) oscillations depends on the membrane potential (269). Hyperpolarization of the cell membrane should facilitate capacitative Ca\(^{2+}\) influx, but in gonadotrophs Ca\(^{2+}\) signaling is terminated after depletion of the ER-Ca\(^{2+}\) pool. Furthermore, there was a rapid recovery of Ca\(^{2+}\) oscillations when cells were depolarized to facilitate VGCl, suggesting that Ca\(_v\) channels in these cells provide the major pathway for Ca\(^{2+}\) influx after the ER-Ca\(^{2+}\) depletion (283). It has also been shown that in rat gonadotrophs, GnRH-stimulated LH release is not mediated by store-dependent Ca\(^{2+}\) influx (279). Further studies in this field should focus on the expression of STIM-Orai in secretory and nonsecretory pituitary cells and, if expressed, on the mechanism of their activation and blockade by GPCRs.

4. **Spontaneous Electrical Activity**

A. **Spiking and bursting**

Electrically excitable cells have been defined as those with voltage-sensitive ion permeability that show regenerative and propagated electrical activity spontaneously or in response to stimulation. Thirty-five years ago it was
shown for the first time that endocrine pituitary cells, like neurons, generate APs (1). Initially, it was believed that only lactotrophs and GH pituitary cells are excitable. With time, it became obvious that all secretory pituitary cell types of vertebrates fall into this category. The membrane potential of isolated pituitary cells in vitro is not stable but oscillates from resting potentials of −60 to −50 mV, reflecting the balance between the activity of depolarizing and hyperpolarizing channels. When membrane potential oscillations reach the threshold level, cells generate APs. In vitro, firing of APs has been observed in frog (284), mouse (285), porcine (286), ovine (72), and bovine (287) endocrine pituitary cells. Firing of APs in cultured cells is not an in vitro artifact; it has also been observed in situ in rat pituitary slices (288).

However, the pattern of electrical activity varies among cells. Gonadotrophs obtained from male rats are typically quiescent (69), whereas about half of those from female rats exhibit spontaneous spiking (184, 289), but this difference could reflect the method of recording (whole cell recording in male gonadotrophs and perforated cell recording in female gonadotrophs). The spiking frequency is typically approximately 0.7 Hz, and the APs are tall and narrow, with amplitude of more than 60 mV (from initiation to peak) and half-width of less than 50 msec (185). Ovine gonadotrophs also fire single APs spontaneously (72). In one study, approximately 80% of lactotrophs and somatotrophs from female rats exhibited spontaneous activity (184). The pattern of activity can be similar to female gonadotrophs, with large and narrow spikes (80, 218, 290), but more often a bursting pattern is produced. These bursts consist of periodic depolarized potentials with superimposed small-amplitude spikes (185, 290, 291). The bursts have a much longer duration (several seconds) than gonadotroph APs, and the burst frequency is significantly lower (~0.3 Hz). The membrane potential rarely goes above −10 mV during a plateau burst, and the spikes are quite small, with amplitude of 10 mV or less (185). This was originally termed “plateau bursting,” but it has recently been renamed “pseudo-plateau bursting” to distinguish it from the type of bursting produced in agonist-stimulated gonadotrophs and pancreatic islets, where the spikes are larger and the bursting pattern is longer and more regular (292). Corticotrophs also exhibit both spontaneous large-amplitude spiking and pseudo-plateau bursting (293, 294), as do melanotrophs (284–286) and GH cell lines (96, 295, 296). Little is known about the spontaneous electrical activity of thyrotrophs.

B. Pacemaking mechanisms

What drives the spontaneous activity of anterior pituitary cells? Although there is still uncertainty about which subthreshold ionic currents are most responsible for depolarizing the cell to the AP threshold, much has been learned in recent years about the candidate currents. The resting membrane potential of −50 to −60 mV in pituitary cells suggests that in addition to resting K⁺ conductance, there are also depolarizing conductances due to other ions. The resting membrane potential rapidly reaches about −85 mV, a value close to equilibrium potential for K⁺, when extracellular Na⁺ is substituted with large organic cations, suggesting the constitutive activity of a Na⁺-conducting channel. Such prominent hyperpolarization of the plasma membrane in the absence of bath Na⁺ causes abolition of spontaneous firing of APs in gonadotrophs, lactotrophs, somatotrophs, and GH₃ cells. In contrast, blockade of Na⁺ channels by TTX does not affect resting membrane potential in a majority of pituitary cells. These observations indicate that constitutive activity of TTX-insensitive Na⁺-conducting channels, termed the background Na⁺ (Naᵦᵦ) channels, contributes to the control of resting membrane potential and may account for the pacemaking depolarization (80, 96, 224, 297). The nature of Naᵦ channels has not been clarified. There are several channel candidates, which could potentially account for the existence of this conductance in pituitary cells. It has been shown recently that the neuronal channel Na⁺ leak channel, nonselective contributes resting Na⁺ permeability (298). Because pituitary cells express several TRP channels (see Section III.A.7), it is also possible that the background activity of these channels could contribute to the resting membrane potential.

The channels that mediate subthreshold TTX-insensitive Na⁺ currents are frequently activated by cyclic nucleotides, either directly or indirectly. Evidence for this hypothesis comes from data showing that inhibition of phosphodiesterases (PDEs) led to an increase in the frequency of bursting in somatotrophs, suggestive of increased excitation (291). Stimulation of pituitary cells with forskolin also initiates firing of APs in quiescent lactotrophs and increases the frequency of firing in spontaneously active lactotrophs (218). Both treatments increase levels of cAMP and cGMP (218, 299), suggesting that elevation in their intracellular concentrations accounts for changes in the pattern of electrical activity.

In general, cAMP can modulate channel activities indirectly, by PKA-mediated phosphorylation of channels (89, 300) or directly by activating HCN and CNG channels (207). Some years ago, Kato et al. (301) showed that GHRH stimulates a rise in [Ca²⁺], and GH secretion by a mechanism involving cAMP/PKA. They also reported that these effects were dependent on bath Na⁺ but were not abolished by TTX in concentrations that block Na₁,4 and Na₆-7 channels. Elevation in TTX concentrations had a partial inhibitory effect, suggesting that TTX-insensitive Na₅ and Nav8-9 channels could account for the
GHRH-stimulation of VGCI and GH secretion (301). Hille’s group (302) also reported a TTX-insensitive Na⁺ current that is up-regulated by PKA phosphorylation and was proposed to be important for GHRH-stimulated pacemaking activity in somatotrophs. The synthetic peptide, GHRP-6, elevates the intracellular Na⁺ concentration in somatotrophs by facilitating Na⁺ influx, which in turns facilitates VGCI (303). This agonist, however, does not elevate cAMP production but operates as a Ca²⁺-mobilizing agonist (304). Recently, Chen’s group (87) reported a stimulatory effect of GHRH on TTX-resistant Na⁺ channels in somatotrophs from GH-green fluorescent protein transgenic mice, but suggested that PKC mediates the action of GHRH. None of these studies suggested that TTX-insensitive Na⁺ channels account for the Na⁺ conductance.

The HCN channels are permeable to both Na⁺ and K⁺, and the current mediated by the channels (Ih) has a reversal potential of about −30 mV (305). Thus, activation of the currents depolarizes the cell. The HCN channels activate at hyperpolarized voltages, typically negative to −50 or −60 mV, and deactivate upon depolarization. Their activity is up-regulated by cAMP, and to a much lesser extent by cGMP. The cyclic nucleotides bind directly to HCN channels and shift the voltage dependence of activation toward more depolarized potentials (306). HCN current is present in several pituitary cell types (see Section III.A.6), but it appears to contribute little to pacemaking. In both AtT20 and GH₃ cell types, spontaneous activity continued when Ih was blocked, although Ih was maximally activated at basal levels of cAMP (209, 210). Such fully activated HCN channels in turn may contribute to resting Na⁺ permeability. These currents may also play the role of a brake of membrane hyperpolarization or in the fast recovery from inhibition that follows activation of Gi/Go-coupled receptors (209). Pituitary cells also express mRNA transcripts for the rod type of CNG channels (see Section III.A.7). However, if functional CNG channels are expressed in pituitary cells, it is unlikely that they contribute to spontaneous firing of APs. Neither application of cGMP-permeable analogs nor stimulation/inhibition of soluble guanylyl cyclase (sGC) activity had any effect on the spontaneous firing of APs in lactotrophs (217, 218). Also, spontaneous firing persisted in lactotrophs when adenyl cyclase (AC) activity was blocked (218).

One subthreshold current, commonly found in neurons, is the T-type Ca²⁺ current. Voltage-clamp studies have shown that this current is present in anterior pituitary cells and is most prominent in somatotrophs (see Section III.A.2). The T-type current activates at lower voltages than other types of Ca⁺⁺ channels, but inactivates within approximately 10 msec. Thus, it provides transient depolarization that can help bring a repolarized cell to the AP threshold. It has been suggested that T-type Ca²⁺ current acts as the pacemaker current for the firing of APs in gonadotrophs (307) or contributes to the control of pacemaking because the frequency of AP decreases in cells with blocked T-type Ca⁺⁺ channels (291). Because PKA does not phosphorylate the T-type channel (91), it is unlikely that it accounts for forskolin-stimulated electrical activity in pituitary cells. The L-type Ca²⁺ channels are stimulated by PKA-dependent phosphorylation of their α-subunits (89), which in turn could change the steady-state Ca²⁺ influx (308) and facilitate pacemaking.

C. Channels involved in spike depolarization

In neurons, TTX-sensitive Na⁺ channels are critical for the development of the depolarizing phase of APs (131). As previously discussed, all pituitary cells express TTX-sensitive Na⁺ channels. However, in the majority of rat anterior pituitary cells, inhibition of these channels does not affect the pattern of spontaneous electrical activity, whereas removal of extracellular Ca²⁺ or blockade of L-type Ca⁺⁺ channels by dihydropyridines abolishes electrical activity without affecting the resting membrane potential, indicating that these channels are critical for spike depolarization (184, 291). In contrast, in a fraction of ovine gonadotrophs (72) and bovine lactotrophs (287), Na⁺ channels are responsible for AP generation. Also, two lactotroph subpopulations have been identified that differ with respect to their level of Na⁺ channel expression; only in lactotrophs expressing high levels of Na⁺ channels did TTX application abolish basal hormone secretion (79). Furthermore, TTX-sensitive Na⁺ channels may contribute to the firing of APs and the accompanied VGCI in frog and rat melanotrophs (65, 66). Involvement of TTX-sensitive Na⁺ channels in PACAP-induced GH secretion was also reported (309).

These differences could reflect the impact of culturing conditions on cell behavior in vitro and the status of TTX-sensitive Na⁺ channels at the resting potential. The lack of TTX-sensitive Na⁺ channel involvement in controlling membrane excitability and secretion in many rat pituitary cells is most likely due to the inactivation of a large proportion (above 90%) of these channels at the resting membrane potential in these cells in vitro (70). Consistent with this, GnRH-induced transient membrane hyperpolarization in rat gonadotrophs is required to remove the steady-state inactivation of TTX-sensitive Na⁺ channels before they can contribute to AP firing (69). In melanotrophs, at typical resting potential of −50 mV, approximately 60–70% of the channels are in the inactivated state, which may explain the presence of a TTX-sensitive component in spike depolarization. However, the firing of TTX-sensitive APs is not an essential requirement for hormone release from these cells (64). These observations raised the
question about the physiological importance of the resting membrane potential on the contribution of Na, and L-type Ca, channels in the spike depolarization. In GnRH-secreting GT1 neurons, a shift in the firing of APs from TTX + dihydropyridine sensitive to exclusively dihydropyridine-sensitive APs was consistently observed in receptor and nonreceptor depolarized cells, which limits the participation of both channels in firing but facilitates AP-driven Ca2+ influx (310, 311). Further studies are needed to clarify whether the same mechanism is operative in endocrine pituitary cells.

D. A mechanism for bursting

Why is the spontaneous activity of gonadotrophs characterized by tall AP spiking, whereas that of other cell types is often characterized by bursting? The simplest explanation could be that there is a cell-specific expression of channels, leading to different patterns of spiking. In a study of ion channel distribution in cells from randomly cycling female rats, it was shown that lactotrophs and somatotrophs express lower levels of TTX-sensitive Na, current than do gonadotrophs (70). As mentioned above, however, blockage of these channels with TTX was shown to have no impact on the frequency of spontaneous electrical activity in these cells (184). The T-type Ca, current is more abundant in somatotrophs than in lactotrophs and gonadotrophs (70), but this could not explain why bursting is observed in both somatotrophs and lactotrophs, but not gonadotrophs.

One type of current that is larger in somatotrophs and lactotrophs than in gonadotrophs is the BK current. These channels activate rapidly upon membrane depolarization, most likely due to colocalization of the BK channels with Ca2+ channels (70, 185). The BK current acts in conjunction with the delayed rectifying K+ current to repolarize the cell membrane during the downstroke of an AP (312). There is evidence that this inhibitory current is the key to bursting behavior in a fraction of somatotrophs (185). First, the membrane-permeable Ca2+ chelator BAPTA-AM was used in somatotrophs and converted spontaneous bursting to large-amplitude spiking. By rapidly chelating Ca2+, BAPTA is thought to greatly attenuate the Ca2+ nanodomain that forms at the mouth of an open Ca, channel, and thus reduce the degree of activation of BK channels. Second, the BK channel blockers iberiotoxin and paxilline both convert bursting to large-amplitude spiking in somatotrophs. Apamin, a blocker of SK channels has little effect on KCa current in somatotrophs. Third, other agents including GHRH and KCl did not convert the bursting to spiking, but only increased the burst frequency and produced baseline depolarization. Taken together, these data suggest that BK channels are a key element in the production of bursting, and that their greater expression in somatotrophs is responsible for the different activity patterns of somatotrophs/lactotrophs and gonadotrophs (185).

Mathematical modeling was used to understand how an inhibitory current could have a stimulatory action by converting spiking to bursting (185). Results from a similar model are shown in Fig. 8 to demonstrate the behavior. When a hyperpolarizing voltage-independent current is added to a spiking model cell, it simply reduces the spike frequency and produces some baseline hyperpolarization (Fig. 8A). Increasing the magnitude of the current accentuates this effect and eventually brings the model cell to a low-voltage steady state (data not shown). If instead a hyperpolarizing BK-like current is added to a spiking cell, the spiking is converted to bursting (Fig. 8B). The burst frequency and size of the spikes is decreased when more BK-like current is added. The explanation is that the fast activation of the BK current reduces the amplitude of an AP. As a result, less delayed rectifying current is activated, so the downstroke of the spike is less extreme, reaching less negative voltages. Thus, the spikes of the burst ride on a depolarized plateau. It is thought that the burst ends when Ca2+ has built up sufficiently to activate BK channels that are more distant from the Ca2+ channels (185). When this accumulated Ca2+ is removed by Ca2+-ATPases, a new burst is initiated. A mathematical analysis of this type of bursting has been performed (291, 313), and the bursting is named “pseudo-plateau bursting.” The resetting properties of this type of bursting oscillation are quite different from those of “plateau bursting” typically exhibited by neurons (292).

Consistent with this observation, stimulation of BK channels prolongs the duration of APs, whereas their inhibition potentiates the firing of APs in dorsal root ganglion neurons (314). Participation of these channels in broadening of APs was also observed in rat amygdala cells.
However, in some pituitary cells exhibiting bursting, blockade of BK channels does not lead to single spiking. In GH3 cells, BK channels act primarily to end the APs (169, 312). Mathematical modeling suggests that the A-type K+ current, like the BK current, can help convert a spiking cell to a bursting cell (313). The mechanism for this is similar to that for the BK current, in that the A-current limits the amplitude of the voltage spike and thus reduces the activation of the delayed-rectifying K+ current. It has been suggested recently that such diverse effects on AP firing probably depend on the type of KCa channels, gating properties, and the context of other channels (167). In rat chromaffin cells, inactivating and noninactivating BK channels contribute differentially to AP firing behavior (316). Also, in the somatotroph cell model the localized Ca2+ rather than bulk Ca2+ accounts for the burst-promoting effect of BK channels. A small distance between Ca, and BK channels was also proposed for rat supraoptic neurons (317) and Xenopus motor nerve terminals (318). Additional experiments are needed using several different pituitary cell types exhibiting plateau bursting and single spiking, preferably in intact tissue, to clarify the specific roles of BK channels in electrical activity and the alternative mechanisms for generating plateau bursting.

E. Functional roles of spontaneous spiking

In their article published in 1996, Mollard and Schlegel (319) addressed the question of why endocrine pituitary cells are excitable. Since then there has been significant progress in understanding the role of excitability in pituitary cell functions. It appears that both changes in the membrane potential and the accompanied changes in Ca2+ influx have functional roles in endocrine pituitary cells. In this section, we review the functional role of spontaneous excitability in isolated pituitary cells at resting conditions. In the following sections, the focus is on modulation of spontaneous electrical activity by gap junction coupling (Section V) and activation of receptor channels (Section VI) and GPCRs (Sections VII and VIII) that are endogenously expressed in pituitary cells.

1. AP-driven Ca2+ signals

The high-voltage activated Ca, channels in pituitary cells not only give rise to APs in the same way as Na, channels, but also provide an effective pathway for Ca2+ influx during the transient depolarization, which acts as an intracellular messenger controlling a variety of cellular functions. The patterns of spontaneous electrical activity in the different cell types have a large impact on the intracellular Ca2+ dynamics and overall Ca2+ levels. Simultaneous measurements of membrane potential and [Ca2+]i showed that the bulk Ca2+ levels are low in spontaneously spiking gonadotrophs (20 to 70 nM), whereas they are much higher (300 to 1200 nM) and clearly oscillatory in spontaneously bursting lactotrophs and somatotrophs (184, 291) and GH3B6 cell line (296). Others also observed high-amplitude spontaneous Ca2+ transients in somatotrophs (186), lactotrophs (94), corticotrophs (320), and GH and AtT-20 cell lines (96, 125, 321–323). Rhythmic bursts of Ca2+ transients were also observed in acute anterior pituitary slices (324). The pattern of Ca2+ signaling also varies among cells of the same origin. For example, the light fraction of lactotrophs was found to have higher basal PRL release and this is correlated with high [Ca2+], and the presence of spontaneous Ca2+ transients, as well as a depolarized (~−45 mV) resting potential and spontaneous electrical activity. In contrast, the heavy fraction of lactotrophs has a more hyperpolarized resting potential (~−65 mV), and the cells are generally silent with lower [Ca2+], levels (325). A similar heterogeneity was found in porcine somatotrophs, with the low-density somatotrophs exhibiting higher basal [Ca2+], than the high-density somatotrophs (326).

The difference in the patterns of Ca2+ transients between cells firing single APs and those exhibiting pseudoplateau bursting is reflected in the dynamics of Ca2+ channel activation and in the spatial distribution of Ca2+ within the cell. Both large-amplitude spikes and bursts depolarize the membrane sufficiently to activate the various types of Ca2+ channels expressed in pituitary cells (70, 104). However, Ca, channels are open for a short time during the short duration of a gonadotroph AP, and as a consequence the elevated Ca2+ concentration is localized to nanodomains that form at the inner mouth of open channels. With the longer duration and smaller amplitude of somatotroph/lactotroph bursts, channels stay open longer and significant Ca2+ entry occurs throughout the burst, which lasts several seconds. A global Ca2+ signal is produced because individual Ca2+ nanodomains overlap, producing a global signal that is easily resolved with fluorescent Ca2+ dyes such as fura-2, as shown by confocal measurements in pituitary somatotrophs (214). Thus, the Ca2+ influx summed over time is much greater during bursting than during large-amplitude spiking (180).

It is unlikely that Ca2+-induced Ca2+ release through RyRs contribute to the generation of such global Ca2+ signals in mammalian lactotrophs and somatotrophs (214, 270), but their contribution in GH3C1 pituitary cell types should not be excluded at the present time (323). The most complex pattern of spontaneous Ca2+ oscillations was observed in frog melanotrophs. These cells also exhibit spontaneous Ca2+ transients, which are dependent on Ca2+ influx through Ca, channels (327, 328). The rise in Ca2+ , however, occurs in a stepwise manner (329), and
The generation of Ca\(^{2+}\)/H1\(100\) transient is abolished in cells in which the ER Ca\(^{2+}\)/H1\(100\) pump is blocked by thapsigargin (330). It appears that in these cells spontaneous VGCI is coupled to Ca\(^{2+}\)/H1\(100\)-induced Ca\(^{2+}\)/H1\(100\) release, presumably through IP\(_3\)Rs (331).

2. Dependence of the cyclic nucleotide signaling pathway on electrical activity

Anterior pituitary cells not only fire APs spontaneously, but also generate cyclic nucleotides in resting conditions. Two lines of evidence support the conclusion that basal AC activity accounts for cAMP production in unstimulated cells. First, in pituitary cells in vitro several inhibitors of ACs decrease basal cAMP production in cells with inhibited PDEs, a family of enzymes that metabolize cyclic nucleotides (218). Second, basal cAMP production in pituitary cells in vitro was also inhibited by activation of two G\(_{i/o}\)-coupled receptors, dopamine D\(_2\) and ET\(_A\) (218, 281). Unstimulated pituitary cells also generate cGMP due to basal activity of the nitric oxide synthase-sGC signaling pathway, as indicated by a concentration-dependent decrease in cGMP production in cells treated with variable nitric oxide synthase inhibitors (332–335). Intracellular cyclic nucleotide levels in pituitary cells in resting conditions are controlled by PDEs (336) and multidrug resistance proteins, which in pituitary cells operate as cyclic nucleotide efflux pumps (224, 332). The relevance of PDEs in the control of intracellular cyclic nucleotides in pituitary cells at rest was indicated in experiments with 3-isobutyl-1-methylxanthine, a general inhibitor of these enzymes (218, 334), whereas the relevance of multidrug resistance proteins in the control of cyclic nucleotide intracellular levels at rest was suggested based on experiments with probenecid, an inhibitor of this pump (224, 332).

In many cell types, Ca\(^{2+}\)/H1\(100\) and cyclic nucleotide signaling pathways are tightly interconnected at the level of intracellular messenger generation and at the level of their intracellular effectors. That is also the case with pituitary cells. Figure 9 summarizes the effects of VGCI on the cyclic nucleotide signaling pathway in pituitary cells. Both normal and immortalized GH pituitary cells express Ca\(^{2+}\)/H1\(100\)-inhibitable AC, as indicated by the ability of elevated Ca\(^{2+}\)/H1\(100\) to inhibit cAMP production in rat broken cells and cell membranes. In these experiments, concentrations of Ca\(^{2+}\) required for inhibition of AC activity were in the range observed in intact pituitary cells, suggesting that spontaneous electrical activity may influence cAMP production. In intact pituitary cells, VGCI also attenuates intrinsic AC activity independently of the status of PDEs (218). In GH\(_3\) cells, there is an intimate colocalization of ACs with L-type Cav channels and capacitative Ca\(^{2+}\) entry channels (276). RT-PCR and Western blot analysis confirmed the expression of Ca\(^{2+}\)/H1\(100\)-inhibitable AC3, AC5/6, and AC9 in pituitary cells (218). AC9 is also expressed in AtT-20 cells, and its activation leads to stimulation of VGCI, which inhibits the enzyme. The negative feedback effect of Ca\(^{2+}\)/H1\(100\) on the enzyme activity is mediated by calcineurin (337).

The nitric oxide synthase-sGC signaling pathway is also modulated by Ca\(^{2+}\)/H1\(100\) influx through Ca\(_v\) channels. Calcium stimulates the activity of several adenylyl cyclases, PDE1, and nitric oxide synthase (NOS) in a calmodulin (CaM)-dependent manner (continuous lines). It also inhibits some adenylyl cyclase isozymes and sGS directly (dotted lines). Changes in the resting membrane potential affect the multidrug resistance protein (MRP)-mediated cyclic nucleotide efflux. PKG, Protein kinase G; LPS + IFN, lipopolysaccharide and interferon.
also Ca\(^{2+}\)-independent and cAMP/PKA-dependent activation of sGC (299, 333). In contrast, facilitation of VGCI in high K\(^+\)-depolarized pituitary cells leads to inhibition of sGC activity, suggesting that Ca\(^{2+}\) also serves as a negative feedback to break the stimulatory action of nitric oxide on enzyme activity in intact pituitary cells (335). It is also well established that Ca\(^{2+}\) plays important roles in the control of PDE-1 activity in other cell types. Roles of these enzymes in the control of cyclic nucleotide signaling have not been systematically investigated in pituitary cells.

The excitability of pituitary cells may also influence multidrug resistance protein-mediated cyclic nucleotide efflux. In normal and GH\(_1\) pituitary cells, abolition of Na\(_9\), conductance by complete or partial replacement of extracellular Na\(^+\) with organic cations or sucrose not only induces a rapid and reversible hyperpolarization of cell membranes and inhibition of AP firing, but also rapidly inhibits cyclic nucleotide efflux. Valinomycin-induced hyperpolarization of the plasma membranes also inhibits cyclic nucleotide efflux, whereas depolarization of the cell membrane facilitates cyclic nucleotide efflux. In contrast to AC and sGC, AP-driven Ca\(^{2+}\) influx is not coupled to the control of the cyclic nucleotide efflux pump activity. It appears that changes in the resting membrane potential not only influence steady-state Ca\(^{2+}\) influx through Ca\(_9\) channels and switches the pattern of firing between TTX-sensitive and dihydropyridine-sensitive channels, but also represents the signal for changes in cyclic nucleotide pump activity (224).

### 3. AP-secretion coupling

Neurotransmitter and hormone secretion is a process of synthesis and release of proteins out of the cell. The path of a protein destined for secretion has its origins in the rough ER, and the protein then proceeds through the many compartments of the Golgi apparatus before ending up in small secretory vesicles containing neurotransmitters (neurons) and large dense-core vesicles (also known as secretory granules) containing hormones (neuroendocrine and endocrine cells). Biogenesis of both types of secretory vesicles (formation of immature vesicles and their remodeling to form mature secretory vesicles) was followed over a significant distance, using actin- and the microtubule-based cytoskeletons along with several motor proteins. Vesicles in the reserve pool are loosely tethered to the plasma membrane, whereas those that are docked are held within a bilayer’s distance from the plasma membrane (<5–10 nm for synaptic vesicles). Stable docking probably represents several distinct, molecular states: the molecular interactions underlying the close and tight association of a vesicle with its target may include the molecular rearrangements needed to trigger bilayer fusion. Tethering and docking of a vesicle at the target membrane precedes the formation of a tight core SNARE complex, a step called priming. Vesicle fusion is driven by SNARE proteins, which when triggered cause the vesicle membrane to merge with the plasma membrane, releasing the neurotransmitters/hormones into the synaptic cleft (for neurons) or extracellular space (for hormones). The trafficking of secretory vesicles toward the plasma membrane and their fusion with the plasma membrane is termed exocytosis (338).

In general, this process can occur in the absence of stimuli (constitutive exocytosis), or in response to stimuli (regulated exocytosis). The main difference between constitutive and regulated exocytosis is in the last two steps of exocytosis. In neuronal and endocrine exocytosis, there is priming of secretory vesicles, including all of the molecular rearrangements and ATP-dependent protein and lipid modifications taking place after the initial docking of a synaptic vesicle but before fusion, and a rise in \([\text{Ca}^{2+}]_i\) is needed to trigger nearly instantaneous neurotransmitter release. In other cell types, whose secretion is constitutive (i.e., continuous, Ca\(^{2+}\)-independent, nontriggered), there is no priming and no need for elevation in \([\text{Ca}^{2+}]_i\) to complete fusion of secretory vesicles. In cells secreting by regulated exocytosis, not only VGCI but also GPCR-mediated Ca\(^{2+}\) mobilization from the ER can initiate vesicle release. Other signaling molecules triggered by activation of GPCRs also contribute to the control of hormone release by exocytosis, suggesting that the term stimulus-secretion coupling is more appropriate for regulated exocytosis, independent of the pathways involved (338–340). Here we will use the term AP secretion coupling to focus on the role of spontaneous electrical activity and the accompanying VGCI in hormone secretion by endocrine pituitary cells.

Pituitary cells secrete hormones in a Ca\(^{2+}\)-regulated manner. Calcium plays several roles in this process, including priming of secretory granules, and triggering of granule exocytosis (101, 341–343). The last step has a low Ca\(^{2+}\) affinity and requires an elevation of intracellular Ca\(^{2+}\) that results from the opening of Ca\(^{2+}\) channels or release of Ca\(^{2+}\) from intracellular stores (which does not occur in unstimulated pituitary cells). Early experiments by Douglas and Shibuya (344) showed that removal of Ca\(^{2+}\) and blockade of L-type Ca\(_9\) channels by dihydropyridines diminish \(\alpha\)-MSH release, whereas facilitation of VGCI by high K\(^+\)-induced depolarization of cells facilitated hormone release. Our early experiments also showed that removal of bath Ca\(^{2+}\) and addition of nifedipine reduced \([\text{Ca}^{2+}]_i\) and diminished GH and PRL release (345). However, we were unable to observe any significant changes in basal release of LH, FSH, TSH, and ACTH. In contrast, high K\(^+\)-induced depolarization and the conse-
quent Ca\(^{2+}\) influx stimulated secretion of all six hormones in a mixed population of pituitary cells (345). These observations could suggest that only melanotrophs, lactotrophs, and somatotrophs exhibit spontaneous firing of APs.

However, as discussed above, all endocrine pituitary cells are excitable. More recently, we showed that lactotrophs, somatotrophs, and gonadotrophs from the same preparation exhibit spontaneous firing of APs, but only in somatotrophs and lactotrophs is the spontaneous electrical activity coupled to hormone secretion (184). Specifically, in perifused pituitary cells, the level of basal secretion of PRL from lactotrophs and GH from somatotrophs is high, whereas basal secretion of LH from gonadotrophs is negligible, although these cells also exhibit spontaneous activity. The near absence of secretion from gonadotrophs, even in the presence of electrical activity, is in contrast with neurotransmitter release from synapses, where single APs are typically effective at evoking release. In the case of the synapse, there is spatial colocalization of Ca\(^{2+}\) channels and secretory vesicles, so that single Ca\(^{2+}\) nanodomains are capable of evoking release (346). Such extreme colocalization is not present in endocrine and neuroendocrine cells, so secretion is evoked by the overlap of many Ca\(^{2+}\) nanodomains. This is well illustrated in experiments with melanotrophs, in which all classes of Ca\(_i\) channels couple with equal efficiency to exocytosis (347).

The difference in basal secretion between gonadotrophs and lactotrophs/somatotrophs is likely due to the very different basal levels of [Ca\(^{2+}\)]\(_i\), and the location of release sites relative to Ca\(^{2+}\) channels. In the presence of Bay K 8644, the duration of single APs in gonadotrophs is prolonged, resulting in larger [Ca\(^{2+}\)]\(_i\) transients and initiation of LH release (184). On the other hand, in somatotrophs, conversion of pseudo-plateau bursts to single APs by BK channel blockade reduces spontaneous Ca\(^{2+}\) influx (185). Similarly, single APs evoke only a small amount of secretion from chromaffin cells, whereas prolonged depolarization induces massive secretion (348). So the much higher basal level of [Ca\(^{2+}\)]\(_i\) in lactotrophs and somatotrophs compared with gonadotrophs leads to the higher basal hormone secretion in these cells. Indeed, blockage of Ca\(^{2+}\) channels with nifedipine reduces basal secretion from lactotrophs and somatotrophs to levels similar to the basal secretion level from gonadotrophs (184). The dependence of basal GH secretion on the amplitude of spontaneous Ca\(^{2+}\) transients is nicely shown in measurements of [Ca\(^{2+}\)]\(_i\) by imaging microscopy and GH secretion by plaque assay in the same cells (186). The same study also showed that the amount of GH released correlates with both the amplitude and the frequency of Ca\(^{2+}\) transients. The L-type Ca\(_v\) channels and K\(_p\) channels play a critical role in the frequency control of Ca\(^{2+}\) transients and PRL release (125). Single-cell recordings of secretion in lactotrophs using single green fluorescent atrial natriuretic peptide-labeled secretory vesicles and FM 4-64 revealed that basal hormone release, also known as spontaneous secretion, is slow compared with stimulated exocytosis, which occurs rapidly. The authors also observed differences between two secretion modes in lactotrophs, both in terms of kinetics and in the rates of loading and discharge of the two probes (349).

**V. Signaling by Gap Junction Channels**

Pituitary cells are not randomly distributed throughout the gland but are highly organized in three-dimensional network structures. Folliculostellate cells make the most impressive network (350). In rodents, this network starts to develop 10 d after birth and is fully developed by the peripubertal period (351). The GH-producing cells also form a network shortly after GH-expressing cells are formed (embryonic d 16), and this network undergoes profound changes, especially during puberty (352). Different contact and signaling molecules could contribute to the formation of these networks, including cadherins (353). Such networks provide the basis for coordinating the activities of different members of the endocrine pituitary population. There are two mechanisms for communication between cells: electrical and chemical. The first requires cells to be coupled through low resistance pathways such as gap junctions. The second requires the release of chemical transmitters, which act as agonists for receptors expressed in electrically interconnected (autocrine mode of regulation) and neighboring cells (paracrine mode of regulation). In this section, we discuss the expression of gap junction proteins in pituitary cells, and in Sections VI–VIII, we focus on receptor channels and GPCRs.

The cytoplasmic compartments of neighboring cells are frequently connected by gap junctions, which are clusters of intercellular channels that form a cytoplasmic bridge between adjacent cells to allow for the cell-to-cell transfer of ions, metabolites, and small messenger molecules, including Ca\(^{2+}\), ATP, cAMP, cADP ribose, and IP\(_3\). Thus, gap junction channels provide an effective mechanism for electrical, Ca\(^{2+}\), and metabolic coupling, depending on the size of the pore. Vertebrate intercellular channels are made up of a multigene family of conserved proteins called connexins. The invertebrate gap junction channels, called innexins, have no detectable sequence homology with vertebrate gap junctions, although they exhibit similar functions and membrane topology. Recently, another family of junctional coupling proteins, called pannexins, has been identified in mammals. These channels have low sequence homology, but general structure similarity, to a family of...
innexins. The gap junction proteins show identical membrane topology: four TM domains connected by two extracellular loops and one intracellular loop with both N and C termini in the cytosol. Such structure is essential for the formation of hexameric pore complexes termed hemichannels, which are large, nonselective ion channels expressed in the plasma membrane before their assembly into gap junctions (354).

A. Connexins

Mammalian connexins are encoded by a gene family of 20 members. Six connexin subunits assemble in a circle to form hemichannels known as connexons in the plasma membrane that can dock to another hemichannel in the plasma membrane of an adjacent cell to form an intercellular channel that spans the gap between the two cells. Hemichannels can contain a single type of connexin (homomeric), or multiple connexins (heteromeric) to form the hemichannel pore, and two identical connexons or different connexons can join to form either homotypic or heterotypic intercellular channels, respectively. The presence of heteromeric connexins and heterotypic intercellular channels can produce a diverse group of structurally different intercellular channels, with different permeabilities and/or function. A variety of other factors, including membrane potential, Ca$^{2+}$, pH, and phosphorylation of channels, can modulate gap junction channels. Several neurotransmitters and hormones, such as dopamine, acetylcholine, GABA, and estrogens, have also been found to alter intercellular channel activity. Because of the large size of the channel pore, several diffusible second-messenger molecules are potential candidates for mediating the propagation of intercellular Ca$^{2+}$ waves via gap junctions, including IP$_3$ and Ca$^{2+}$ itself (355, 356).

Gap junctions in the anterior pituitary were initially shown by Fletcher et al. (5). Subsequent studies have revealed that gap junctions are formed between folliculostellate cells and that the number of gap junctions increases with the developmental increase in the number of these cells. Other physiological and experimental conditions also influence the gap junction connections of the folliculostellate cells (56, 357, 358). Electrical coupling between some, but not all, folliculostellate cells was observed (359). The network of these cells participates in the long distance conduction of information in intact anterior pituitary cells, which involves Ca$^{2+}$ (350). It has also been suggested that the gap junction-mediated network of folliculostellate cells provides messages necessary for the hormone release by anterior pituitary cells (357).

Northern blot analysis and immunostaining studies indicated the expression of connexins 26, 32, and 43 in pituitary cells (360), and localization of connexin 43 in folliculostellate cells and pituicytes (361). In mink anterior pituitary, changes in connexin 43 expression in folliculostellate cells are associated with seasonal changes in PRL content (362). TGFβ3 may act on folliculostellate cells to increase gap junction communication, resulting in stimulation of fibroblast growth factor by these cells (363). Locally produced adenosine stimulates connexin 43 expression and gap junctional communication in folliculostellate cells (364). In a folliculostellate cell line, proinflammatory cytokines also modulate the level of connexin 43 expression (365), and TNF-α causes cell uncoupling mediated by connexin 43 dephosphorylation (366). The recent development of the S100b-GFP transgenic rat with expressed green fluorescent protein specifically in folliculostellate cells in anterior pituitary will facilitate further work on the relevance of this network in pituitary cell functions (367).

It has also been suggested that the endocrine anterior pituitary could be coupled by gap junctions. In a hypothalamo-pituitary slice preparation from the tilapia fish, electrotonic coupling between neighboring cells was detected, as well as diffusion of Lucifer Yellow between cells. Such coupling was observed in about one third of the cells (368). Diffusion of Lucifer Yellow was also observed in intact anterior pituitary cells up to 300 µm apart from its site of induction. In addition to folliculostellate cells, coupling was also observed between lactotrophs and somatotrophs (369). This conclusion was confirmed using [Ca$^{2+}$]$_i$ measurements by real-time confocal imaging in pituitary slices and halothane, a gap junction blocker. It appears that somatotrophs in pituitary slices are either single units or arranged in synchronized gap junction-coupled assemblies scattered throughout the anterior lobe (370).

B. Pannexins

Pannexins are a three-member family of channels. Unlike connexins and innexins, homomeric pannexin 1 hexamers do not form gap junctions when expressed in mammalian cells but operate as hemichannels (371). They are activated by membrane depolarization, mechanical stress, and in a receptor-dependent manner. The channel pore is permeable to ions, small molecules, and metabolites up to 1 kDa, including ATP, ADP, nicotinamide adenine dinucleotide, cyclic nucleotides, and IP$_3$. Such wide permeability probably accounts for their numerous nonjunctional functions in variable cell types (372, 373). ATP-gated P2X$_7$ receptor channels are the potential partners in Panx1-mediated signaling (374, 375). However, the details regarding the association of pannexin 1 with purinergic receptors and their modes of interaction have not been clarified. This includes a lack of information regarding the specificity of physical associations between purinergic receptors and pannexins. Recently, it has been shown that rat anterior pituitary cells expressed pannex-
VI. Signaling by Receptor Channels

The pattern of electrical activity in single pituitary cells is also modulated by chemical signals. These signals act as ligands or agonists for specific plasma membrane receptors expressed in pituitary cells. Many of these agonists are delivered by hypothalamic neurons and are released into the posterior lobe or into the hypophyseal portal systems. Other agonists are secreted by pituitary cells and act in autocrine and paracrine manners. Agonists can also reach the pituitary cells through the general blood circulation. There are several classes of stimuli, including neurotransmitters, hormones, eicosanoids (metabolites of arachidonic acid), growth factors, and chemokines. Four groups of receptors recognize these agonists: extracellular ligand-gated ion channels (receptor channels), GPCRs, enzyme-linked receptors (receptor tyrosine kinases, natriuretic peptide receptors, cytokine receptors, and intracellular enzyme-containing receptors), and intracellular steroid receptors. There are other channels that are activated or modulated by ligands, such as IP₃Rs and RyRs that are expressed in the ER membrane (see Section III.C) and cyclic nucleotide-regulated channels of the plasma membrane (see Section III.A.6), and they are known as intracellular ligand-gated ion channels. In this section, we review the literature on the role of receptor channels in electrical activity, whereas the role of GPCRs in electrical signaling is summarized in Sections VII and VIII. Pituitary cells also express the enzyme-linked receptors and the intracellular steroid receptors, but their roles in electrical activity and Ca²⁺ signaling have not been systematically investigated and will not be reviewed here.

Receptor channels contain two functional domains: an extracellular domain that binds an agonist, and a TM domain that forms an ion channel. Because these proteins combine transmitter binding and channel functions into a single molecular entity, they are also called ionotropic receptors. The agonists for these channels are acetylcholine, GABA, glycine, 5-HT, glutamate, and ATP. Based on ion conductivity, receptor channels are divided into two classes: the excitatory cation-selective channels, operated by acetylcholine, glutamate, 5-HT, and ATP; and the anion-selective channels, activated by GABA and glycine, which are usually inhibitory. With the exception of glycine, these agonists can also activate so-called metabotropic receptors, which belong to the family of GPCRs. These receptors do not have an ion channel as a part of their structure, but they can affect channel activity through one or several metabolic steps (see Sections VII and VIII).

From a structural point of view, receptor channels belong to three families of evolutionary related proteins (377, 378). The acetylcholine, 5-HT, GABA, and glycine-activated receptor channels are grouped as one family, known as ligand-gated ion channels of the Cys-loop family. These channels are composed of five subunits (pentamers), each of which contributes to the ionic pore. All subunits have a large extracellular N-terminal region followed by four hydrophobic TM segments and an extracellular C terminus (379). The second family represents glutamate-activated receptor channels, which are also composed of four TM segments, but their TM2 segment forms a pore-loop structure, entering and exiting the cell membrane from the intracellular side. Thus, the N terminus is extracellularly located, whereas the C terminus is intracellularly located and is regulated by signaling molecules, including the kinases. A detailed analysis of the intrasubunit interactions that govern glutamate-receptor assembly indicates that these channels are dimers of dimers (380). The third family is known as P2XR channels. Members of this family have only two TM domains, with the N and C termini facing the cytoplasm. As with acetylcholine and glutamate channels, the functional diversity of P2XR channels is generated by subunit multimerization. The functional channels are composed of three subunits (381).

A. Cys-loop family of receptor channels

1. Nicotinic acetylcholine receptor channels

Acetylcholine is an agonist for two classes of membrane receptors: muscarinic and nicotinic acetylcholine receptors. Muscarinic receptors belong to the GPCR superfamily of receptors. There are five subtypes of these receptors, termed M₁–M₅; the M₁, M₃, and M₅ receptors signal predominantly through the Gₛ/₁₁ pathway, whereas M₂ and M₄ receptors are coupled to the G₁₆ signaling pathway (382). Nicotinic acetylcholine receptors (nAChRs) are a family of acetylcholine-gated channels. The nAChRs are more diverse, with genes encoding a total of 17 identified subunits that can assemble into a variety of pharmacologically distinct receptor subtypes. Muscle types of nAChRs are located postsynaptically at the neuromuscular junctions, where they mediate fast synaptic transmission of electrical signals from motor neurons. Neuronal types of nAChRs are expressed in the central and peripheral nervous system and are distributed post-, pre-, and perisyn-
aptically. The pore of activated channels is permeable to Na$^+$ and K$^+$ and, for some neuronal subtypes, to Ca$^{2+}$ (383, 384).

The role of acetylcholine as a putative autocrine factor has been relatively well established in intermediate pituitary functions. Acetylcholine is released from frog melanotrophs onto M1 receptors (385) and stimulates electrical activity and α-MSH release (386, 387). Functional nAChRs are described in porcine intermediate pituitary cells at both the whole-cell and single-channel levels (388). These channels are depolarizing, and their activation leads to facilitation of Ca$^{2+}$ influx directly through the pore of the channel and indirectly by activating Ca$^+$ channels (389). The possibility that nAChRs are cross-coupled to the PLC signaling pathway has also been proposed (390), whereas the role of these channels in secretion has not been studied. Denef's laboratory (391) also suggested that acetylcholine acts as a paracrine factor in the anterior pituitary. This group observed that immunoreactivity for choline acetyltransferase, the enzyme catalyzing the biosynthesis of acetylcholine, was present in anterior pituitary cells and that acetylcholine was released by high K$^+$ depolarized pituitary cells. They further showed that acetylcholine stimulates secretion by corticotrophs and the corticotroph cell line AtT-20 through activation of nAChRs (4). The structure of nAChRs, their biophysical and electrophysiological properties, and Ca$^{2+}$-signaling function have not been studied in the anterior pituitary cells.

2. 5-Hydroxytryptamine receptor (5-HT$_3$R) channels

The neurotransmitter 5-HT is a native agonist for seven receptors. Six of these are heteromeric GPCRs, whereas the 5-HT$_3$R operates as a receptor channel (392). The 5-HT$_3$R exists as a pentamer of four TM subunits that form a cationic-selective channel. Three 5-HT$_3$ subunits (5-HT$_{3A}$, 5-HT$_{3B}$, and 5-HT$_{3C}$) have been cloned, but only homomeric 5-HT$_{3A}$ and heteromeric 5-HT$_{3A}+3B$ form functional receptors when expressed in heterologous systems. A short form of the 5HT$_{3A}$ subunit was also identified, but this splice form does not differ physiologically from the full-size channel. Homomeric and heteromeric channels mediate a rapidly activating, desensitizing, inward current that predominantly carries Na$^+$ and K$^+$. Some forms are also permeable to Ca$^{2+}$ (393). 5-HT$_3$Rs are expressed throughout the central and peripheral nervous systems, where they mediate a variety of physiological functions. The receptors are also involved in information transfer in the gastrointestinal tract, and in the enteric nervous system they regulate gut motility and peristalsis (394). The hypothalamic actions of 5-HT and its receptors, including the control of PRL, gonadotropin, CRH, AVP, and oxytocin release, are well characterized (reviewed in Refs. 54 and 395). In addition, 5-HT$_3$Rs are likely to be expressed in the fish pituitary and to play important roles in signaling and secretion (396). It has also been suggested that functional 5-HT$_3$Rs are expressed in mammalian pituitary cells in culture and LBT3 cell lines. The evidence includes RT-PCR analysis and pharmacological studies on basal and agonist-stimulated LH and ACTH release, but not the electrophysiological characterization of 5-HT$_3$R current and its role in signaling (397–400).

3. GABA receptor channels

GABA acts as a neurotransmitter through three structurally and pharmacologically distinct classes of receptors: G protein-coupled GABA$_B$ receptors and ligand-gated GABA$_A$ and GABA$_C$ chloride channels. There are two GABA$_B$ subunits, and functional receptors are probably heterodimers; the specific agonist for these receptors is baclofen (401). To date, 16 different GABA$_A$ subunits (α1-6, β1-3, γ1-3, δ, ε, π, and θ) have been cloned and sequenced from the mammalian nervous system. Additional variants arise through alternative splicing (402). The GABA$_A$ receptor is a pentameric assembly derived from a combination of various subunits. The preferred combination includes two α-, two γ-, and one β-subunit. However, the colocalization of these three types of subunits is not an absolute requirement for the formation of functional channels. The great diversity of receptor subunits leads to profound differences in tissue distribution, ontogeny, pharmacology, and regulation of GABA$_A$ receptors. These receptors are targets for many drugs in wide clinical use, including benzodiazepines, barbiturates, neurosteroids, ethanol, and general anesthetics, which increase the conductance through the pore of the channels. A specific agonist is muscimol, and a specific blocker is bicuculline (403, 404). The molecular components of GABA$_C$ receptors are p1-3 subunits, which form functional channels without assembling with GABA$_A$-α and -β subunits. These receptors are specifically activated by (+)-cis-2-aminomethylcyclopropane carboxylic acid (405).

GABA$_{AC}$ channels are chloride ion channels, and the nature of their actions depends on the [Cl$^-$]. In the majority of adult neurons, [Cl$^-$] is low and activation of GABA$_{AC}$ channels leads to hyperpolarization of the cell membrane and silencing of electrical activity. In developing neurons, however, [Cl$^-$] is relatively high and GABA channels are depolarizing, leading to facilitation of electrical activity and VGCI. Chloride homeostasis in most brain cells is controlled by two electrically neutral cation/chloride cotransporters, called NKCC1 and KCC2. The ubiquitously expressed NKCC1 derives energy from the electrochemical gradient for Na$^+$ to take up Cl$^-$, whereas KCC2 uses the K$^+$ gradient to facilitate Cl$^-$ extrusion.
and that bicuculline and picrotoxin block the action of GABA. It has also been shown that muscimol inhibits PRL release from pituitary cells (409, 412). The RT-PCR analysis indicated the presence of α1, α4, β1, β2, β3, and γ2-subunits in a mixed population of anterior pituitary cells (409, 412). More recently, mRNA transcripts for α2, α3, α5, α6, γ1, γ3, δ, and ε-subunits were found in anterior pituitary cells, and the expression of α3, α5, and α6 mRNAs was somewhat lower. Immunocytochemical studies further showed the expression of α1- and β1-subunit proteins in all secretory anterior pituitary cells (414). Immunohistochemical labeling revealed that frog melanotrophs in situ and in cell culture were intensely stained with α2, α3, γ3, and β2/β3-subunits (415). Thus, the whole repertoire of mRNAs for γA3 receptor subunits is present in the pituitary gland, providing the possibility for variable and cell type-specific combinations of subunits into pentamers.

There have been contradictory reports about the nature of GABA actions in pituitary cells. Earlier studies suggested that GABA inhibits PRL release in vitro. It has also been reported that muscimol inhibits PRL release in vitro and that bicuculline and picrotoxin block the action of GABA and muscimol, suggesting the presence of hyperpolarizing γA3 receptors in these cells and their potential inhibitory role in VGCI and secretion (for references, see Ref. 414). Inhibitory effects of GABA on α-MSH secretion from the intermediate lobe and on AVP and oxytocin from the posterior pituitary were also reported by several laboratories (for references, see Ref. 416). Contrary to these findings, GABA and muscimol were found to stimulate, rather than inhibit, secretion of ACTH, GH, LH, and TSH from pituitary cells in vitro (417–419). It has also been shown that GABA increases [Ca2+]i in a majority of the anterior pituitary cells, including lactotrophs, gonadotrophs, and αT3-1 cells. This effect is mimicked by muscimol, antagonized by picrotoxin, and abolished by removal of extracellular Ca2+ (414, 420, 421). In frog melanotrophs, GABA also stimulates Ca2+ influx and α-MSH release (422). Augmentation of exocytosis and the depolarizing effect of GABA at high [Cl−]i has also been documented in melanotrophs from mouse pituitary tissue slices (236) and frogs (284, 423). Stimulation of γA3 receptors in somatotrophs also increases [Ca2+]i (424), further supporting the view that GABA receptors in the majority of pituitary cells are depolarizing, leading to stimulation of VGCI.

Electrophysiological measurements provided more conclusive evidence for the expression of γA3 channels. The GABA-induced activation of a chloride current with pharmacological properties of γA3 receptors was initially shown in bovine lactotrophs (425), frog melanotrophs (426), and neonatal rat anterior pituitary cells (427). GABA-induced currents were also detected in posterior pituitary nerve terminals (416, 428). It has also been suggested that the activity of γA3 channels depends on the status of sGC activity in frog melanotrophs (429). To clarify whether GABA-induced current is depolarizing or hyperpolarizing, it is essential to preserve [Cl−]i, which was done in intermediate and anterior pituitary cells using gramicidin-perforated patch clamp recordings. These experiments showed that the reversal potential of GABA current is positive to the resting membrane potential, indicating that [Cl−]i in the majority of pituitary cells from adult animals is elevated and that activation of γA3 channels leads to Cl− efflux causing depolarization. GABA-induced depolarization of pituitary cells was associated with either an increase in the frequency of APs in spontaneously firing cells or a sustained depolarization (284, 414).

In accordance with these results, the expression of NKCC1 in postpubertal anterior pituitary cells is high, whereas mRNA expression for KCC2 (if present) is low (414), and imaging studies suggested that [Cl−]i in lactotrophs is around 50 mM (231). In posterior pituitary nerve endings, however, [Cl−]i was estimated to be around 20 mM, and the GABA current is hyperpolarizing (416). The contradiction in the field about the nature of GABA actions in anterior pituitary cells might be explained in part by the presence of γA3 receptors, because their activation leads to inhibition of spontaneous electrical activity and basal AC activity (402), and both Ca2+ and cAMP regulate exocytosis in these cells (343, 430–432).

Although the majority of cells in the anterior and intermediate lobes express GABA receptors and their in vitro activation triggers substantial Cl− influx and alters the pattern of electrical activity, Ca2+ signaling, and hormone secretion, the in vivo operation and physiological relevance of this signaling pathway has not been clarified (433). In search of a PRL inhibitory factor, Schally et al. (434) isolated GABA from the hypothalamus. Subsequent studies showed that GABA is released from tuberoinfundibular and other hypothalamic regions, that concentrations of GABA in portal blood are higher than in peripheral blood, and that electrical stimulation of the median eminence induces a several-fold increase in the rate of GABA release (435), which could indicate that GABA acts in the pituitary as a hypothalamic neurohormone. However, it has also been reported that GABA is synthesized and released from intermediate and anterior pituitary...
lobes (435–438), which could suggest that GABA acts as a paracrine factor. At the present time, little information exists about the mechanism of GABA release in the portal blood and pituitary. It has been suggested that substance P modifies hypothalamic GABA release (439) and that injection of estradiol leads to a several-fold increase in the intrapituitary GABA concentration (440).

4. Glycine receptor (GlyR) channels

GlyRs are pentameric proteins composed of three α-subunits and two β-subunits. In contrast to other members of this group of receptor channels, GlyRs do not have a counterpart in the GPCR receptor family. There are four isoforms of α-subunits, which have highly homologous sequences but different pharmacological and functional properties, and alternative splicing of α-subunits further increases GlyR heterogeneity. The α-subunit contains the ligand-binding site and is sufficient to form a functional homomeric channel, whereas the β-subunit modulates the pharmacological and conductance properties of the GlyRs. There are many similarities between GABA and GlyRs, including ion selectivity, which arise from their close and conservative evolutionary relationship. As with GABA receptors, the direction of the flux depends on the electrochemical gradient for Cl−. In contrast to GABA receptors, GlyRs are not expressed in the anterior pituitary. However, there are reports on the expression of these receptors in the nerve endings in posterior pituitary cells and their activation by taurine, a GABA-like amino acid that is released by pituicytes (441, 442).

B. Glutamate receptor channels

L-Glutamate is the major excitatory neurotransmitter in the CNS, acting as an agonist for eight members of GPCRs (443) and receptor channels, each encoded by 18 genes that assemble to form four major subtypes: AMPA (α-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid); kainate; NMDA (N-methyl-D-aspartate); and δ receptor channels. Molecular cloning has revealed several subunits for each receptor group. Four AMPA receptor genes (GluR1–GluR4) denote the AMPA-sensitive family, whereas five kainate receptor genes (GluR5–GluR7, KA1, and KA2) make up the kainate subclass. For NMDA-receptor channels, seven subunits (NR1, NR2A–NR2D, NR3A, and NR3B) have been established. In addition, two δ-subunits exist, belonging to the GluR type, but the function of this particular subunit is unknown. Finally, the molecular diversity of NR1 and GluRs is further increased by variants created by alternative splicing and RNA editing. In addition to the specific structure and pharmacology, NMDA channels exhibit a different excitation behavior than the other channel types. These channels are both ligand- and voltage-gated. Full activation of the NMDA receptor requires application of two ligands, L-glutamate and glycine. The NMDA receptors only become fully activated by glutamate after their Mg2+ block has been relieved by membrane depolarization. Kainate and AMPA-receptor subunits do not form mixed channel complexes, but both types of receptors can be expressed in the same neuron. Native AMPA receptors are either homomeric or heteromeric oligomers composed of these multiple subunits (444).

Glutamate has numerous well-established indirect effects on pituitary hormone secretion by modulating hypothalamic functions (445). In addition, several reports have suggested that glutamate directly affects cells of the anterior and intermediate lobes. However, there are some contradictions in these reports. Initially, it was reported that glutamate stimulates PRL release in perfused pituitary cells, and this effect is abolished by a selective non-competitive NMDA receptor antagonist (446). Further work in this field suggested a dual effect of glutamate on PRL release, consisting of a stimulatory effect mediated via receptor channels and an inhibitory effect via GPCRs (447). Others reported inhibitory effects of kainate and NMDA on PRL release in static cultures (448). A stimulatory effect of glutamate through non-NMDA receptors on LH release was also reported (449), and kainate-2 mRNA was detected in embryonic rat pituitary tissue (450). Double immunohistochemistry suggested that only a fraction (4–11%) of all secretory anterior pituitary cells express NMDA receptors (451). Single-cell Ca2+ measurements showed a glutamate-induced rise in [Ca2+]i in TRH- and GHRH-responsive rat cells (452). In tilapia PRL cells, glutamate also induced a rise in [Ca2+]i, due to cell depolarization and activation of Ca2+ channels (453). No data on the electrophysiological characterization of glutamate receptor channels in anterior pituitary cells are available. In contrast, the expression of these receptor channels in cells from the intermediate lobe was demonstrated by patch clamp recording of glutamate-induced current, and pharmacological characterization of these responses was consistent with the presence of AMPA-type glutamate channels in these cells (454). Single-cell secretory data confirmed that activation of these channels is sufficient to trigger α-MSH release (455), whereas agonists specific for glutamate GPCRs were unable to trigger release of this hormone (456).

C. Purinergic receptor channels

ATP is released by excitable and nonexcitable cells and acts as an extracellular messenger for two families of purinergic receptors: seven-TM domain P2Y receptors (P2YRs) and two-TM domain P2XR channels. The agonist actions of ATP are terminated by several enzymes,
These experiments reveal functional P2XRs in all secreting cells, establishing the role of P2XRs in anterior pituitary cells. ATP also triggers other signaling pathways (458–460). Through homo- and heteropolymerization, the P2X7R generated currents. They can form ion permeable pores after the initial depolarization of cells by P2XR-generated IP3 signals. UTP (uridine-5'-diphosphate), and UDP glucose are also native ligands that hydrolyze ATP to ADP, AMP, and adenosine. ADP and adenosine also act as extracellular messengers; ADP activates a few P2YRs but not P2XR, whereas adenosine acts as an agonist for four G protein-coupled adenosine receptors (Fig. 10). UTP (uridine-5'-triphosphate), UDP (uridine-5'-diphosphate), and UDP glucose are also native agonists for P2YRs. The purinergic signaling pathway is operative in the hypothalmo-posterior pituitary system as well as in the intermediate and anterior pituitary lobes (457).

P2XRs comprise a family of ATP-gated cation channels, which are expressed in numerous excitable and non-excitable cells and play important roles in a variety of physiological processes. Seven mammalian P2XR subunits, termed P2X1–7, and several nonmammalian subunits have been identified. Each subunit is composed of cytoplasmic N and C termini, two TM domains, and a large extracellular domain; and three subunits are required for formation of a functional receptor. P2XRs differ with respect to their ligand-selectivity profiles, antagonist sensitivity, and cation selectivity. Their activation leads to an increase in \([Ca^{2+}]_{i}\), with \(Ca^{2+}\) influx occurring through the pores of these channels and through \(Ca_{v}\) channels after the initial depolarization of cells by P2XR-generated currents. They can form ion permeable pores through homomeric and heteropolymerization. The P2X7R also triggers other signaling pathways (458–460).

Single-cell \(Ca^{2+}\) measurements were instrumental in establishing the role of P2XRs in anterior pituitary cells. These experiments reveal functional P2XRs in all secretory anterior pituitary cell types and raise the possibility that several subtypes of these channels are expressed in a cell type-specific manner (reviewed in Ref. 461). However, this method is of limited use for identifying the receptor subtypes expressed because the rapidly desensitizing homomeric and heteromeric P2XRs are not able to generate global \(Ca^{2+}\) signals (462). In more recent studies, molecular biology techniques combined with electrophysiology were used to better understand the structure of P2XRs expressed in anterior pituitary cells and their downstream signaling pathways. Quantitative RT-PCR analysis revealed that secretory cells abundantly express P2X2R and P2X4R, with less expression of other subunits. Western blot analysis showed the expression of P2X2R, P2X4R, and P2X7R at the protein level. Cloning experiments showed that rat anterior pituitary cells express two functional splice forms of the P2X2 subunit, termed P2X2a and P2X2b (463), and that mouse pituitary cells express three functional forms of the P2X2R subunit, termed P2X2a, P2X2b, and P2X2e. The P2X2b and P2X2e subunits are missing 69 and 90 residues, respectively, in their C termini (464). When expressed as homomeric channels, three splice forms of P2X2R differ in the rate of receptor desensitization; P2X2eR desensitizes most rapidly, at a rate comparable to that observed in cells expressing P2X1R and P2X3R, whereas the rate of P2X2bR desensitization is faster than P2X2a but slower than P2X2e receptors (464). Deletions in the C terminal of P2X2aR also effectively reduced the peak amplitude and duration of \(Ca^{2+}\) signals, indicating a role of Arg371-Pro376 (P2X2R numbering) in receptor desensitization (465, 466). The physiological relevance of these splice forms is in the formation of functional heteromers, which desensitize faster than full-size receptors but slower than the homomeric splice receptors. This in turn limits excessive ion influx but does not terminate signaling during prolonged agonist stimulation (467).

Functional P2X2Rs are expressed in gonadotrophs and somatotrophs, but not in other pituitary cell types (463). In gonadotrophs, their activation leads to firing of APs along with modulation of the frequency of firing in spontaneously active cells, accompanied by elevation in \([Ca^{2+}]_{i}\), that reflects \(Ca^{2+}\) influx through both P2X2R channel pores and \(Ca_{v}\) channels (468). The ATP-induced rise in \([Ca^{2+}]_{i}\) is sufficient to trigger LH release (468, 469). ATP also influences GnRH-induced current and membrane potential oscillations in an extracellular \(Ca^{2+}\)-dependent manner. These IP3-dependent oscillations are facilitated, slowed, or stopped depending on ATP concentrations and the time of ATP application (468). Thus, P2X2R could contribute to the pacemaking and modulation of GPCR-controlled electrical activity (468). Mice deficient in
P2X2R subunit are available and are fertile (470), but changes in pituitary function in such mice have not yet been studied. The biophysical and pharmacological properties of recombinant rat P2X4R cloned from the pituitary gland have also been characterized. This receptor desensitizes with a rate comparable to that observed in cells expressing P2X2bR. In contrast to the P2X2, suramin, pyridoxal-phosphate-6-azophenyl-2',4'-disulfonylic acid tetrasodium salt (PPADS), and reactive blue 2 are not effective antagonists of P2X4R, but these receptors are sensitive to ivermectin, a high molecular weight lipophilic compound used as an antiparasitic agent in human and veterinary medicine. Ivermectin increases sensitivity of P2X4R to ATP, amplifies peak current amplitude in response to supramaximal agonist concentrations, and delays receptor deactivation (471). These allosteric actions of ivermectin on P2X4R were successfully used in structural and functional characterization of recombinant receptors (471–473).

The PPADS insensitivity of P2XRs in TRH-responsive cells suggests that lactotrophs and/or thyrotrophs from the anterior lobe express functional P2X4Rs (474). This was confirmed recently by electrophysiological characterization of P2XR current in TRH-responsive cells. Activation of these channels leads to stimulation of electrical activity and promotion of voltage-gated and voltage-insensitive Ca\(^{2+}\) influx in these cells. In the presence of ivermectin, the peak amplitude of the current increases, as well as the sensitivity of the receptors to ATP, whereas the receptor deactivation slows. The activation of these receptors causes depolarization, which is sufficient to increase the frequency of APs and to initiate spiking in quiescent cells, as well as to facilitate Ca\(^{2+}\) influx through Ca\(_{\text{V}}\) channels. Calcium influx through the pore of P2X4Rs also contributes to ATP-induced Ca\(^{2+}\) signaling. Ivermectin also enhances ATP-induced PRL release, indicating that these receptors are expressed in lactotrophs (475). Further studies should clarify whether thyrotrophs also express these channels and whether there is any change in the status of electrical activity and Ca\(^{2+}\) signaling in pituitary cells from P2X7\(^{-/-}\) animals, which are available (476).

The physiological sources of ATP required for activation of purinergic receptors in the pituitary gland remain largely uncharacterized. The magnocellular neurons of the hypothalamus with nerve endings in posterior pituitary also contain ATP, and the specific pattern of APs originating from the cell bodies of these neurons has been suggested to control the release of this nucleotide (477). The extracellular ATP concentration in posterior pituitary can reach 4–40 \(\mu\text{M}\), a concentration range sufficient to activate the majority of P2XRs (478). In addition to its action on nerve terminals of vasopressinergic neurons in the posterior pituitary, released ATP probably acts on pituicytes (478, 479).

In the anterior pituitary, ATP probably acts as an autocrine/paracrine factor. Normal and immortalized GH\(_3\) and αT3-1 pituitary cells release ATP at resting conditions, and such basal ATP release is enhanced in cells treated with ARL67156, an inhibitor of ectonucleotidases (480). GnRH-induced stimulation of gonadotropin release is accompanied by elevation in basal ATP release, suggesting that ATP is stored in the secretory vesicles of these cells (469). This is consistent with an earlier study showing Ca\(^{2+}\)-dependent ATP release (247) and modulation of ATP release by PRL secretagogues (481). However, we did not observe ATP release when PRL release was evoked by depolarization of perifused pituitary cells (480), and the amplitude of GnRH-induced ATP release in perifused pituitary cells is too low to activate endogenous P2XRs. In other tissues, ABC-binding cassette transporters, pannexins, and P2X7R have been suggested to participate in nonvesicular ATP release (482). Interestingly, anterior pituitary cells express functional multidrug resistance proteins (224, 332) and P2X7R (475, 483), which could contribute to ATP release.

The action of ATP as an autocrine/paracrine factor is critically dependent on its rapid metabolism by ectonucleotidases, which include members of the ectonucleotide triphosphate diphosphohydrolase family of enzymes (E-NTPDase) and ecto-5’-nucleotidase, among others. Four of eight known E-NTPDases are expressed in the plasma membrane. These enzymes not only hydrolyze extracellular ATP and/or ADP to AMP but also metabolize other nucleotide tri- and diphosphates, including UTP and UDP, whereas cAMP is hydrolyzed by the ecto-5’-nucleotidase family of enzymes (484, 485). These enzymes are also expressed on the plasma membrane of pituicytes and neurosecretory posterior pituitary terminals (486). Extracellularly applied ATP is rapidly hydrolyzed by the isolated posterior pituitary accompanied by accumulation of adenosine, suggesting that these enzymes provide a pathway for the activation of adenosine receptors in this tissue and termination of ATP-induced vasopressin release (487). The mRNA transcripts for plasma membrane-located E-NTPDases 1, 2, and 3 are also expressed in pituitary tissues, cultured pituitary cells, and αT3-1, AtT-20, and GH\(_3\) cell lines, and normal and immortalized pituitary cells rapidly metabolize ATP (480). Ecto-5’-nucleotidases (CD73), which generate adenosine from AMP, were found by immunocytochemistry in a fraction of anterior pituitary cells (364). Because anterior pituitary cells express the ADP-activated P2Y1 receptor and several adenosine-activated receptors, it is reasonable to suggest a role for
ectonucleotideases in the sequential activation of purinergic receptors in the anterior pituitary (Fig. 10).

VII. Role of GPCRs in the Regulation of Electrical Activity

GPCRs are a very large and diverse superfamily of receptors that help define cellular responsiveness to extracellular signals. They share a common structure of seven α-helix TM domains with an extracellularly located N terminus and an intracellularly located C terminus and are coupled to heterotrimeric G proteins. These G proteins have α- and βγ-subunits, which function as transducers to relay information to different signaling pathways, such as the PLC and AC signaling pathways, which operate as amplifiers by producing intracellular messengers. These messengers carry information to intracellular sensors and effectors. Downstream effectors produce one or more actions, including membrane depolarization/hyperpolarization, activation/inactivation of membrane Ca\(^{2+}\) channels, mobilization of Ca\(^{2+}\) channels from the ER, and sensitization/desensitization of the exocytotic machinery. The specific action initiated by the hormone agonist is determined by the nature of the GPCR and the cell type (488, 489). In this section and the following section, we discuss the actions of GPCRs linked to the G\(_{411}\), G\(_{s}\), and G\(_{i/o/z}\) signaling pathways and their impact on the electrical activity, cytosolic Ca\(^{2+}\) dynamics, and secretion from pituitary cells. The focus in our review will be on the role of G proteins, Ca\(^{2+}\), cyclic nucleotides, PKC, and PKA in control of channel activity, Ca\(^{2+}\) signaling, and secretion. These receptors have numerous other functions, including activation of the MAPK cascade, which are not discussed here because of their limited role in electrical signaling and calcium mobilization.

A. Stimulation of electrical activity by GPCRs

cAMP is a ubiquitous intracellular messenger generated by the AC family of enzymes that regulates numerous cellular responses, including electrical activity and VGCI. There are nine plasma membrane isoforms of these enzymes, each with two 6TM regions and two cytosolic domains, C1 and C2, which contain the catalytic region that converts ATP into cAMP (Fig. 11). The intrinsic activity of these enzymes is up-regulated by GPCRs linked to heterotrimeric G\(_{s}\) proteins and down-regulated by GPCRs linked to heterotrimeric G\(_{i/o/z}\) proteins. There are several cAMP signaling effectors: PKA, PDEs, CNG and HCN channels, and the exchange proteins guanine nucleotide exchange factors (GEFs) that activate small GTP binding protein Rap1. PKA is composed of two regulatory subunits (R) and two catalytic subunits (C) (Fig. 11). Binding of cAMP to the R subunits enables the C subunits to phosphorylate different substrates, including the plasma membrane channels. A family of PKA-anchoring proteins determines the cellular localization of PKA. The cross talk between cAMP and Ca\(^{2+}\) is important in the control of cellular functions. In general, activation of receptors linked to G\(_{s}\) increases [Ca\(^{2+}\)], by up-regulating electrical activity. In cells expressing HCN and/or CNG channels, this occurs through their activation and the subsequent depolarization of the plasma membrane and facilitation of Ca\(_{\text{L}}\) channel activity. PKA also phosphorylates numerous plasma membrane channels, including Ca\(_{\text{L}}\) channels, leading to facilitation of excitability of cells. Phosphorylation of other channels, such as Na\(_{\text{L}}\), down-
regulates excitability or changes excitability from Na\(^+\) to Ca\(^{2+}\) spikes (2).

GPCRs linked to the G\(_s\)-signaling pathways are operative in endocrine pituitary cells. The G\(_s\)-signaling pathway in corticotrophs is triggered by hypothalamic CRH (490). Somatotrophs express two receptors coupled to the G\(_s\)-signaling pathway, GHRH (491) and VIP/PACAP (47). VIP/PACAP receptors are also present in mammalian melanotrophs (492), lactotrophs (54), and folliculostellate cells (493). Some eicosanoids may also signal through this pathway in pituitary cells (494). The lack of expression of typical G\(_s\)-coupled receptors in other pituitary cell types does not mean that the cAMP signaling pathway is not operative. These cells also express ACs, and their activities are regulated by other mechanisms. This could include the cross-coupling of other GPCRs to the G\(_s\) signaling pathway, as suggested by stimulation of AC by GnRH receptors in other cell types (495, 496). VGCI also affects cAMP signaling in pituitary cells by modulating AC activity (218, 497). Activation of G\(_s\)-linked GPCRs in pituitary cells stimulates electrical activity and facilitates VGCI. The type of Ca\(^{2+}\) response typically obtained through this pathway is a plateau elevation of [Ca\(^{2+}\)], or an increase in the frequency and/or amplitude of Ca\(^{2+}\) transients. The cross talk between Ca\(^{2+}\) and cAMP also exists at effector levels, including the control of exocytosis (see Section VII.B.4).

1. **CRH-induced calcium influx in corticotrophs**

CRH, also known as CRF, is the main regulator of ACTH release in normal and immortalized corticotrophs. It acts on CRH receptors known as CRF-R\(_1\) receptors coupled to the G\(_s\) signaling pathway, leading to stimulation of cAMP production (19, 490). One of the main functions of CRH in corticotrophs is to modulate spontaneous electrical activity and facilitate Ca\(^{2+}\) influx; in the absence of extracellular Ca\(^{2+}\), _de novo_ production of cAMP is not affected, but CRH-induced ACTH release is completely blocked. This does not exclude the Ca\(^{2+}\)-independent effects of the cAMP signaling pathway on exocytosis, but it demonstrates that the modulatory role of this signaling pathway could be manifested only in the presence of elevated Ca\(^{2+}\). The relevance of this cation in cAMP effects on secretion is also demonstrated in experiments with AVP and CRH. AVP stimulates Ca\(^{2+}\) mobilization from the ER in corticotrophs, but alone it is not a very potent secretagogue. However, in the presence of CRH, secretion is greatly enhanced by AVP (498).

A key element in the control of spontaneous firing of APs in pituitary cells, including corticotrophs, is control of the resting membrane potential and slow membrane depolarization, called the pacemaking depolarization. CRH changes the resting membrane potential and the rate of the pacemaking depolarization, leading to an increase in the firing rate of spontaneously active cells and causing silent cells to become active (293, 320, 499). The slow membrane depolarization is caused in large part by a reduction in a background K\(^+\) conductance mediated by a member of the K\(_w\) channel family (294, 500). Depolarization alone is not sufficient to induce spiking in quiescent cells (499), and the firing frequency of cells depolarized by CRH is higher than for corticotrophs depolarized to the same voltage level by blocking the K\(_w\) current. This indicates that a small component of the depolarization might be mediated by the reduction of another type of background K\(^+\) conductance or facilitation of an inward current (294). A mathematical model of the corticotroph confirms that depolarization induced by blocking a background K\(^+\) current facilitates spiking but cannot by itself trigger AP firing (501). On the other hand, facilitation of the L-type Ca\(_v\) current is sufficient for the model to generate spikes or bursts (501, 761). However, there is no evidence that CRH receptor activation leads to phosphorylation of these channels by PKA, as in other cell types. The inward current is not generated by HCN channels, which are expressed in AtT-20 cells, but are fully activated by the resting cAMP levels (209). The background Na\(^+\) conductance discovered by Simasko and colleagues (80, 297), which is present in all endocrine pituitary cells (224), could play a major role in CRH- and GHRH-induced electrical activity and secretion (discussed in Section VII.A.2).

The inhibition of K\(_w\) and associated depolarization and increase in spike frequency last up to 15 min after removal of CRH from the physiological solution, suggesting that phosphorylation of K\(_w\) channels could account for this memory (294). These effects of CRH and their time courses are mimicked by application of forskolin, an activator of AC, and by membrane-permeant analogs of cAMP (294, 500). No CRH-induced depolarization is observed in the presence of intracellular Rp-cAMPS, a blocker of PKA (500), confirming that the effects of CRH on the pacemaking depolarization are mediated through cAMP activation of PKA. However, some effects of CRH were resistant to the PKA inhibitor H-89, raising the possibility that CRH might act through an additional G protein pathway (499, 502).

Ca\(_v\) channels can be activated with sufficiently strong pacemaking depolarization, and their opening produces the upstroke of the AP spike and a robust Ca\(^{2+}\) influx. It appears that the main channel involved is the L-type Ca\(_v\) channel, but P-type Ca\(_v\) channels also play a role in the regulation of spike frequency, and another high-voltage-activated and toxin-resistant Ca\(_v\) channel may as well (293). Rapid PKA-mediated phosphorylation of L-type Ca\(_v\) channels has not been studied but should not be excluded. However, cAMP stimulates the expression of the
L-type Ca\(_2\)\(^{2+}\) channels in AtT-20 cells at the mRNA and protein levels, presumably through PKA (503). Corticotrophs also express T-type Ca\(_2\)\(^{2+}\) channels, but their role in CRH action has not been studied. The recovery of membrane potential in corticotrophs depends on at least two K\(^{+}\) channels, the delayed rectifier and BK channels. In AtT-20 cells, CRH inhibits BK channels through activation of PKA (188). Thus, whereas the intracellular signaling pathway activated by CRH may be the same in corticotrophs and AtT-20 cells, the PKA targeted ion channels appear to be different.

2. GHRH-induced calcium influx in somatotrophs

The GHRH receptor is expressed predominantly in the pituitary gland and is coupled to the G\(_{s}\) signaling pathway. In rats, there are two splice forms of this receptor showing similar sensitivity to GHRH, but only the short receptor isoform stimulates cAMP production (491). In a fraction of porcine pituitary cells, GHRH could also trigger Ca\(^{2+}\) mobilization in an IP\(_3\)-dependent manner, which could indicate the cross-coupling of GHRH receptors to the G\(_{q/11}\) signaling pathway (504). Like CRH in corticotrophs, GHRH facilitates electrical activity and Ca\(^{2+}\) influx through L-type Ca\(_2\)\(^{2+}\) channels of silent or already active somatotrophs (214, 291, 302, 505–509). Forskolin also increases Ca\(^{2+}\) influx in somatotrophs (510, 511), and inhibition of PDEs increases the electrical activity of somatotrophs (291), indicating the relevance of cAMP in GHRH action. An increase in electrical activity after GHRH application is also observed in pituitary slices (288). This response can be detected for up to 90 min after removal of GHRH (505). Such long-lasting effects of G\(_s\)-linked receptors on the electrical status of somatotrophs suggests that the time course of the phosphorylation-dephosphorylation cycle, rather than the direct effect of cAMP, could account for prolonged effects of GHRH on electrical activity in these cells. Consistent with this, although somatotrophs and GH\(_3\) lactosomatotrophs also express HCN channels (210, 211), these channels are unlikely to contribute to the GHRH-stimulated electrical activity because basal AC activity is sufficient to fully activate them.

It is possible that GHRH decreases the intrinsic activity of a K\(_s\) channel in somatotrophs, as CRF does in corticotrophs. This was shown in experiments with blockade of these channels in spontaneously firing cells and model simulations (291). Several findings suggested that a background Na\(^{+}\) conductance could mediate the action of GHRH on electrical activity in these cells. First, extracellular Na\(^{+}\) is essential for GHRH-induced and cAMP-induced GH release. Removal of Na\(^{+}\) does not affect GHRH-stimulated cAMP production, further indicating that this messenger alone is not sufficient to trigger exocytosis. Second, blockade of Na\(_v\) channels by TTX does not mimic the effects on GH release of replacing bath Na\(^{+}\) with organic cations. Finally, Li\(^{+}\) can substitute for Na\(^{+}\) in GHRH actions (301, 512). Isolation of GHRH-stimulated current revealed that the channel is permeable to Na\(^{+}\), Li\(^{+}\), and K\(^{+}\) but not to organic cations, and that the stimulatory action of GHRH is mimicked by a cAMP analog (302) and blocked by inhibition of PKA (513). The Na\(^{+}\)-dependent inward conductance involved in the GHRH-induced depolarization could be the background conductance that mediates spontaneous oscillations of membrane potential. This conductance is inhibited by large (10 mM) extracellular concentrations of Ca\(^{2+}\), Mg\(^{2+}\), and Sr\(^{2+}\) and is activated at low Ca\(^{2+}\) levels (214).

It appears that the action of GHRH on electrical activity also includes other ion channels. In parallel to the effects of cAMP-PKA on cardiac Ca\(_2\)\(^{2+}\) channels, GHRH was shown to increase L- and T-type Ca\(^{2+}\) conductances in ovine somatotrophs and human adenoma GH cells (514–516). A role for delayed rectifier and A-type K\(^{+}\) conductances in GHRH-induced depolarization in ovine somatotrophs, human adenoma GH cells (514, 517), and GH\(_4\)C\(_1\) cells (143) has also been suggested. Although the increase in the Ca\(_v\) currents was mediated through cAMP/PKA (514, 516), the decrease in the K\(^{+}\) currents was not blocked by PKA inhibitors, but was abolished by PKC inhibitors and mimicked by a PKC activator (143, 514, 517). The synthesized GH-releasing peptide GHRP-2 also acts through the cAMP/PKA pathway to increase L- and T-type currents (518). Other GH-releasing peptides that have been synthesized also depolarize the somatotrophs membrane, allowing Ca\(^{2+}\) influx, but they may act through different pathways (519).

3. VIP/PACAP-induced calcium influx in pituitary cells

The high degree of sequence homology between VIP and PACAP would be consistent with the presence of a common receptor for the two agonists. However, three distinct types of receptors exist in vertebrates. The type I (PAC\(_{1}\)) receptor exists in six splice forms, a short form and five variants having inserts in the third intracellular loop of the receptor. Two of these variants are linked to activation of both AC and PLC equipotently; two forms exhibit 10-fold preference for coupling to the PLC signaling pathway; two forms have characteristics intermediate between the two groups. Type II and III (VPAC\(_{1}\) and VPAC\(_{2}\)) receptors exhibit equal potency for VIP and PACAP and signal exclusively through the AC pathway. Pituitary cells express PAC\(_{1}\) and VPAC\(_{2}\) receptors; gonadotrophs express the PAC\(_{1}\) receptor linked to the PLC signaling pathway; and somatotrophs, lactotrophs, and melanotrophs express VPAC\(_{2}\) receptors coupled to the G\(_q\) signaling pathway (47, 520).

PACAP activates Ca\(^{2+}\) mobilization through the PLC/IP\(_3\) pathway mediated by G\(_{q/11}\) in both gonadotrophs and
αT3-1 cells (521). In αT3-1 cells, PACAP also stimulates cAMP production and facilitates extracellular Ca\(^{2+}\) influx through dihydropyridine-sensitive Ca\(_v\) channels, an effect blocked by the PKA antagonist H-89 and mimicked by forskolin and 8-Br-cAMP. This finding is consistent with the expression of both PAC\(_1\) and VPAC\(_3\) receptors in αT3-1 cells. A higher concentration of PACAP (≥ 1 nM) produces a pulse-decay-plateau response, the pulse being mediated by IP\(_3\) and the plateau by cAMP/PKA, a response similar to the TRH-evoked stimulation of the intracellular Ca\(^{2+}\) level in lactotrophs (47, 522).

PACAP stimulates cAMP production and α-MSH release from melanotrophs and ACTH release from ArT-20 cells (523). In melanotrophs, PACAP stimulates Ca\(^{2+}\) influx through L-type Ca\(_v\) channels but does not trigger Ca\(^{2+}\) mobilization from the ER. The rise in [Ca\(^{2+}\)]\(_i\) is mimicked by activation of PKA and inhibited by blockade of this enzyme. Electrophysiological experiments also revealed that PACAP stimulation of melanotrophs causes an inward nonselective cation current, which depolarizes the cells and stimulates VG61 (492). In somatotrophs, PACAP also stimulates Ca\(^{2+}\) influx through Ca\(_v\) channels (524, 525). These responses are similar to the GHRH-evoked Ca\(^{2+}\) plateaus and transients. They are blocked by PKA antagonists and mimicked by forskolin and by the cAMP analog 8-Br-cAMP (526).

PACAP also causes extracellular Ca\(^{2+}\) influx in lactotrophs (527). In GH\(_3\) cells, VIP evokes a modest Ca\(^{2+}\) influx via an increase in cAMP (528), and both VIP and PACAP increase cAMP levels in GH\(_4\)C\(_1\) cells (529). There are also numerous reports of effects of VIP on PRL release from dissociated pituitary cells (reviewed in Ref. 54). There is another mechanism by which the cAMP signaling pathway could contribute to the control of electrical activity in lactotrophs. In vivo lactotrophs are tonically inhibited by dopamine, which decreases cAMP levels and opens K\(^+\) channels, decreasing [Ca\(^{2+}\)]\(_i\) to its basal level (29, 51). The main stimulatory signal for PRL release is generated by inhibition of dopamine secretion and results in an increase in cAMP levels and stimulation of electrical activity and Ca\(^{2+}\) influx (530). Consistent with this, cAMP acts as a potent modulator of electrical activity, Ca\(^{2+}\) influx, and PRL release in lactotrophs in vitro (218).

4. cAMP signaling pathway and secretion
Forskolin, an activator of AC, and cell-permeable cAMP stimulate GH (531), PRL (218), LH (532, 533), and ACTH release (534). In single lactotrophs, cAMP-induced facilitation of exocytosis, measured by changes in the plasma membrane capacitance, was also detected (430). In these cells, forskolin increases the number of granule-to-granule fusion events without altering the number of granule-to-plasma membrane fusion events (535). In single rat melanotrophs, cAMP also stimulates the fusion of larger granules with the plasma membrane (431). In general, facilitation of hormone release reflects dual actions of the cAMP signaling pathway on exocytosis, indirectly by facilitating VG61 (Section VII.A.1–3) and directly on the exocytotic pathway. In some cell types, the calcium-independent effect of cAMP on exocytosis is mediated by PKA (536). In neurons, PKA-dependent facilitation of synaptic transmission includes recruitment of synaptic vesicles from the reserve pool to the readily releasable pool of vesicles (537) and phosphorylation of the secretory vesicle-associated synapsin proteins (538). A more recent study suggested that cAMP facilitates Ca\(^{2+}\)-dependent exocytosis in melanotrophs through both PKA and Epac2 (432). Further studies are needed to identify the PKA and Epac2-sensitive steps in the exocytotic pathway in this and other pituitary cell types.

B. Inhibition of electrical activity by GPCRs
GPCRs linked to the G\(_{i/o}\)-signaling pathways are also operative in endocrine pituitary cells, and their activation leads to inhibition of electrical activity and hormone secretion. Somatostatin (539, 540) and dopamine (51, 541) are two major hypothalamic factors that inhibit pituitary hormone secretion via G\(_{i/o}\)-coupled receptors. Pituitary cells also express several other GPCRs linked to this signaling pathway, including receptors activated by adenosine (542), ET-1 (543), GABA (544), melatonin (545), neuropeptide Y (546), and 5-HT (25) (Fig. 11). Inhibition of AC activity by these receptors represents one of the mechanisms by which spontaneous electrical activity and hormone secretion are inhibited. The 

1. Somatostatin inhibits electrical activity, calcium influx, and hormone secretion
Somatostatin or somatotropin release-inhibiting factor was initially discovered in hypothalamic extracts and was found to inhibit GH secretion from cultured anterior pituitaries. Subsequently, it was found that somatostatin also inhibits TSH and PRL release from normal pituitary cells, GH and PRL release from adenomatous glands in humans and from GH\(_3\)C\(_1\) cells, and ACTH release from human and mouse ACTH-producing tumors. Somatostatin was found in other CNS regions and in peripheral tissues, including the pancreas, the gut, and the thyroid gland. The actions of somatostatin are mediated by five receptors, termed sst\(_1\), sst\(_2\), sst\(_3\), sst\(_4\), and sst\(_5\), all linked to the G\(_{i/o}\) signaling pathway. The effector molecules include AC, K\(^+\) channels, Ca\(^{2+}\) channels, Na\(^+\)/H\(^+\) exchanger, and
cGMP-dependent protein kinase. Pituitary cells express predominantly sst₁, sst₂, and sst₅ receptors (539, 547).

In somatotrophs, somatostatin inhibits spontaneous and GHRH-stimulated electrical activity, VGCI, and GH secretion. In spontaneously firing somatotrophs and GH cell lines, somatostatin hyperpolarizes the plasma membrane, leading to inhibition of electrical activity and basal VGCI (116, 291, 505, 508, 509, 548–551). Somatostatin also inhibits basal and forskolin-stimulated [Ca^{2+}]i in human TSH-secreting adenoma cells (552). Because somatostatin inhibits cAMP production (553), it should antagonize the effects of GHRH mediated through cAMP/PKA. Indeed, somatostatin reverses the stimulatory effects of GHRH and cell permeable cAMP on VGCI (506, 508, 509) and on the background, TTX-independent Na⁺ conductance (554) in somatotrophs and tumoral GH-secreting cell lines. In one study, a low concentration of somatostatin only abolished the early phase of GHRH-induced Ca^{2+} influx, whereas at higher concentration it abolished the early and late phases of the response, suggesting that somatostatin operates on multiple targets (508).

Two channel families modulated by GHRH in somatotrophs (Kᵢᵣ and Cₐᵦ) are also modulated by somatostatin, but in the opposite direction (555); there is evidence that somatostatin activates Kᵢ₃ channels (116, 552) and inhibits Cₐᵦ channels (549, 556–558). The latter is also observed in ACTH-secreting AtT-20 cells (559). It appears that the L-type Cₐᵦ channels are negatively coupled to somatostatin receptors (560, 561) and that withdrawal of somatostatin augments this current in rat somatotrophs (562) and cells from human somatotroph adenomas (563). Whereas GHRH-stimulated and PKA-mediated phosphorylation accounts for facilitation of Cₐᵦ currents in somatotrophs, somatostatin inhibits these channels in a cAMP/PKA-independent manner (556, 559, 564). Control of activity of Kᵢᵣ and Cₐᵦ channels by somatostatin is not unique to pituitary cells, but was also observed in other cell types expressing these receptors (540).

It has also been suggested that somatostatin stimulates BK-type Kᵣᵢᵣ channels through protein dephosphorylation (557), as well as the A-type and delayed rectifier K⁺ channels (145, 549). Inhibition of the background Na⁺ conductance by somatostatin has also been postulated; the somatotroph model of Tsaneva-Atanasova et al. (291) reproduces the effect of somatostatin on membrane potential and intracellular Ca^{2+} by decreasing the background Na⁺-dependent conductance and increasing a Kᵢᵣ conductance. In contrast, activation of BK channels might increase the duration of spontaneous electrical events and therefore may not have an inhibitory effect on Ca^{2+} influx (see Section IV.D).

The effects of somatostatin on ion channels and electrical activity are antagonized by preincubation of the cells with pertussis toxin, confirming that the somatostatin receptor is coupled to the Gᵣₒ family of G proteins (558, 561, 565). There were numerous studies focused on the subtypes of G proteins involved in the coupling of somatostatin receptors to channels. Treatment of ovine somatotrophs with antibodies and antisera to various Gα-subunits has suggested that Gᵣ₂ mediates the reduction of Cₐᵦ currents (566), whereas Gᵣ₃ mediates the increase in Kᵢᵣ currents (567). In GH₃ cells, down-regulation of Gᵣₒαᵢ₂β₁γ₃ protein expression eliminates the inhibitory effect of somatostatin on Cₐᵦ channels (568). It is reasonable to postulate that Gᵣᵢγ complexes released from Gᵣ₁ and Gᵣₒ proteins mediate the action of somatostatin receptors on Kᵢᵣ and Cₐᵦ channels. In GH₃ cells, αᵢ₂ specifically mediates inhibition of AC (569). The ability of somatostatin to stimulate inositol phosphate turnover, Ca^{2+} mobilization, and GH secretion in a fraction of porcine somatotrophs could suggest that the Gᵣᵢγ dimer of Gᵣₒ also stimulates PLC. However, somatostatin in this subpopulation of somatotrophs also triggers elevation in cAMP production (570), suggesting that further studies are needed to clarify the mechanism of activation of these pathways.

2. Dopamine modulation of calcium influx in lactotrophs and melanotrophs

Among catecholamines, dopamine plays the major role in the control of pituitary cell functions. It is secreted from hypophyseal hypothalamic neurons and acts as a principal inhibitory regulator of PRL release by lactotrophs (51, 54) and α-MSH by melanotrophs (571). In low concentrations, dopamine also stimulates PRL release (572). There are five subtypes of dopamine receptors, called D₁, D₂, D₃, D₄, and D₅/D₁₅. By using the radioligand binding assay, it was shown in the late 1970s that the dopamine D₃ subtype of receptors mediates the tonic inhibitory control of hypothalamic dopamine on PRL release in these cells (541). Later investigations showed that two subtypes of D₃ receptors, termed D₃₂ and D₃₁, are generated by alternative splicing in lactotrophs and melanotrophs (571, 573, 574). Lactotrophs express varying ratios of these two receptor subtypes, depending on the level of gonadal steroids (575). Consistent with these findings, the knockout D₂ mice showed chronic hyperprolactinemia, pituitary hyperplasia, and a moderate decrease in α-MSH content (576).

The pituitary dopamine receptors are functionally associated with pertussis toxin-sensitive Gᵣₒ proteins (577–579). Dopamine-induced inhibition of PRL release is also affected by pertussis toxin treatment (572, 578, 580). Two intracellular messengers that play major roles in controlling the fusion of secretory vesicles with the plasma membrane to release hormones in endocrine cells (581), cAMP and Ca^{2+}, are affected by activation of D₂ receptors in
pituitary cells. Early studies suggested that inhibition of cAMP production contributes to inhibition of PRL release (578, 580). However, the relevance of cAMP in dopamine actions on PRL release was questioned by the finding that dopamine inhibits PRL secretion in cells with activated ACs by forskolin (582). We also observed that the dopamine agonist-induced inhibition of basal release of pre-stored PRL was preserved when cAMP levels were elevated by forskolin treatment (583). These experiments certainly do not exclude the modulatory role of cAMP/PKA in Ca\(^{2+}\)-controlled exocytosis, but suggest that control of Ca\(^{2+}\) influx represents the major pathway by which dopamine controls PRL release.

In lactotrophs, dopamine blocks spontaneous and stimulated VGCI in a similar way to Ca\(^{2+}\) channel blockers or removal of extracellular Ca\(^{2+}\) (583–587). Dopamine also hyperpolarizes the membrane and suppresses APs and bursts, which explains the decrease in VGCI (583, 587, 588). A similar effect of dopamine was observed in melanotrophs (589). The role of K\(_a\) channels in dopamine-induced hyperpolarization has been suggested (114, 123, 589, 590). It has also been reported that dopamine increases K\(_a\) conductance (588) and the BK-type K\(_{ca}\) conductance (183). Dopamine was also reported to inhibit Ca\(_a\) channels in lactotrophs (591–593), a conclusion questioned by others (594). Inhibition of high voltage-activated Ca\(_a\) currents was also observed in melanotrophs from neonatal rats (22, 589, 595).

In general, the action of dopamine on electrical activity could be mediated through cAMP-dependent and -independent mechanisms. Dopamine does decrease cAMP in lactotrophs (583) and an elevation of the cAMP level increases electrical activity and Ca\(^{2+}\) influx in these cells (218). However, only a minority of lactotrophs exhibits decreased electrical activity when cAMP is reduced from its basal level (583). Furthermore, whereas the effects of dopamine on VGCI are pertussis toxin-sensitive, they persist in cells with elevated cAMP (123, 583, 587, 588, 596, 597). Finally, the activation of voltage-independent K\(^+\) channels by dopamine is observed in excised outside-out patch (596), demonstrating that no second messenger is required to mediate this action. This suggests that coupling between the G protein and K\(_a\) channels is mediated by the βγ dimer (123).

In physiological conditions in vivo, dopamine tonically inhibits lactotrophs, and a transient release from such inhibition constitutes a stimulatory signal for PRL secretion (530). Even a brief removal of dopamine can potentiate the subsequent PRL-releasing action of TRH, presumably through a cAMP/PKA increase that leads to a long-lasting phosphorylation of Ca\(_a\) channels (530). In support of this hypothesis, dopamine application has been shown to inhibit Ca\(_a\) currents after a short (1–10 min) and a prolonged (over 24 h) application in GH\(_4\)C\(_1\) cells transfected with D\(_2\) receptors. After such treatments, washout of dopamine for 10–40 min doubled Ca\(_a\) currents, and the current further increased 24 h after dopamine removal (590). Withdrawal of dopamine after a short application also evokes a rapid rebound increase of basal PRL secretion above the level observed before agonist application (598), and such rebound is blocked by preventing VGCI (599, 600). Given that both rapidly and slowly inactivating Ca\(_a\) currents are potentiated after a hyperpolarizing conditioning potential, this suggests that recruitment of inactivated channels by dopamine-induced hyperpolarization contributes to the rebound effect on PRL release.

It has also been reported that picomolar doses of dopamine (about 1000 times less than the inhibitory range of concentration) can stimulate PRL release (601). This stimulatory effect is mediated, at least in part, by a rapid increase in [Ca\(^{2+}\)]\(_i\), but not cAMP (572). There is evidence that this stimulatory effect is mediated by D\(_2\) receptors (572, 602) and/or the D\(_5\) receptor (603). There is also contradictory evidence regarding the implication of the G\(_i\) pathway in the stimulatory action of dopamine on PRL release (572, 604). If it is assumed that the effects of a low dose of dopamine encompass a subset of the effects of inhibitory doses, then a possible mechanism is that dopamine activates either BK or A-type K\(^+\) channels. As discussed above, the BK channel can promote bursting and increase the amplitude of Ca\(^{2+}\) oscillations in pituitary somatotrophs. Models of pituitary cells also show that BK and A-type K\(^+\) currents can promote bursting, due to their fast voltage-dependent activation, which prevents full-blown spikes and rapid membrane repolarization (291, 313).

Two additional transduction mechanisms have also been reported for D\(_2\) dopamine receptors in target tissues. First, the D\(_2\) receptors can exert their actions independently of G proteins by promoting the formation of a signaling protein complex composed of β-arrestin, Akt, and protein phosphatase-2A (605). In chromaffin cells, Akt-induced phosphorylation of cysteine string protein plays a role in late stages of exocytosis (606). Akt also regulates the PRL-promoter activity (607), whereas the contribution of this signaling pathway to dopamine-controlled PRL release has not been observed (583). Second, dopamine D\(_{2S}\) and D\(_{2L}\) receptors couple to the same extent to the pertussis toxin-sensitive Gi/0 protein and to the pertussis toxin-insensitive G\(_i\) proteins in vitro (608) and in vivo (609). Other subtypes of dopamine receptors also couple to G\(_i\) proteins (608, 610). Indirect evidence has recently been presented that D\(_2\) receptors in pituitary cells are also linked to the G\(_i\) signaling pathway, and that such coupling...
provides an additional mechanism for inhibition of PRL release downstream of VGCI (583).

Dopamine actions in lactotrophs depend also on the gonadal steroid milieu. Estradiol decreases the expression of D2 receptors (575) and the number of Gi/o immunoreactive lactotrophs (611) but does not affect the expression ratio of the long and short D2 receptor subtypes, in contrast to progesterone and testosterone (575). Also, the stimulatory and inhibitory actions of dopamine on PRL release vary throughout the estrous cycle and in ovariectomized animals vary with steroid replacement therapy (611). The rebound PRL release after dopamine withdrawal also appears to be steroid dependent (612). Dopamine-induced activation of Kcs channels was observed in most lactotrophs from proestrous females, but not in cells from estrous or diestrous rats (612), indicating that effects of estradiol on dopamine response are not limited to the control of expression of D2 receptors. Furthermore, bath application of estradiol can quickly reverse the inhibitory effect of dopamine on electrical activity, indicating a nongenomic action of this steroid hormone on electrical activity (613). Estradiol can also affect the percentage of light and heavy fractions of lactotrophs, which respond specifically to dopamine and TRH (50, 100, 325).

3. ET inhibition of VGCI in pituitary cells

The ET family of peptides, originally discovered for their vasoconstrictive effects on vascular tissue, is composed of three endogenous isoforms (ET-1, ET-2, and ET-3), which are encoded by different genes (614, 615). The peptides are differentially expressed in tissues of the periphery and CNS and have profound effects on neuroregulatory and endocrine functions, in addition to effects on cardiovascular functions (616, 617). In mammals, there are two plasma membrane ET receptor subtypes, ETA (618) and ETB (619). These receptors are GPCRs that signal through variable G proteins, depending on the cell type in which they are expressed (620). The ETA receptor is selective for ET-1 and ET-2 over ET-3, whereas the ETB receptor is activated equally by these peptides (621). The ETc receptor cloned from Xenopus laevis dermal melanophores is ET-3 specific (622); however, the mammalian homolog for the ETc receptor does not exist. ET receptors arise through divergent intron-containing genes, and mRNAs arising from alternative splicing have been reported (621). Some splice isoforms of rat ET receptors are functional (623–625). The human ETA receptor gene has also been proposed to give rise to several alternative splice isoforms (626–628).

Functional ET receptors are expressed in all five major secretory cell types (52, 629, 630), and ETs are produced by pituitary cells (631), suggesting autocrine or paracrine modes of action. Stimulation of these receptors in gonadotrophs leads to activation of the Gq/11 signaling pathway accompanied with the oscillatory Ca2+ release from intracellular pools and gonadotropin secretion (245). The stimulatory action of these receptors on Ca2+ signaling and secretion in gonadotrophs is transient due to their rapid desensitization and internalization (37). In somatotrophs and lactotrophs, ETs also activate the Ca2+-mobilization pathway and transiently stimulate GH and PRL release. In contrast to gonadotrophs, the stimulatory effect of ET is followed by inhibition of PRL and GH release below the basal levels (48, 282, 632, 633). In lactotrophs, the inhibitory phase lasts for several hours (52, 53), arguing against rapid desensitization of these receptors. Such a difference in the actions of ETs in gonadotrophs vs. somatotrophs/lactotrophs would be consistent with the expression of both subtypes of these receptors, but pituitary cells express only ETA receptors (37, 634, 635), which are most likely a combination of the full-size and spliced forms of these receptors (628).

In general, activation of Ca2+-mobilizing receptors leads to sustained Ca2+ influx. In nonexcitable cells, Ca2+ influx occurs through Orai channels (see Section III.C.3), and in excitable cells, Ca2+-mobilizing receptors frequently facilitate VGCI. This is also the case with several Ca2+-mobilizing agonists in pituitary cells, including TRH and angiotensin II (282). In contrast, ET-1-induced Ca2+ mobilization in lactotrophs and somatotrophs is followed by a return of [Ca2+]i levels to baseline for quiescent cells, or to below control levels for spontaneously active cells (282). Such sustained inhibition of VGCI is not affected by raising intracellular cAMP (115), ruling out down-regulation of the cAMP/PKA pathway as the mechanism for ET-1-induced inhibition of Ca2+ influx.

The sustained inhibitory action of ET-1 on [Ca2+]i levels is replaced by stimulation of Ca2+ influx through Ca2+ channels when the cells are pretreated with pertussis toxin, suggesting that activation of the Gi/o pathway inhibits Ca2+ influx. Also, ET-1 inhibits spontaneous and Bay K 8644 (an activator of L-type Ca2+ channels) stimulated Ca2+ transients, but it does not inhibit Ca2+ influx stimulated by high K+, suggesting that the inhibition is not mediated by directly closing Ca2+ channels. Rather, ET-1 increases a cesium-sensitive Kcs current in both somatotrophs and lactotrophs (48, 115). In parallel to somatostatin and dopamine actions, facilitation of Kcs currents by ET-1 hyperpolarizes the membrane, suppressing electrical activity and the resulting Ca2+ transients. Physiologically, the suppression of VGCI is sufficient to block secretion. In cells in which the Gi/o signaling pathway is blocked, ET-1 still inhibits AC activity and PRL release, indicating that there is cross-coupling of ET-A receptors to
impairment may also replace dopamine as the primary inhibiting factor. Electro-
physiological experiments revealed that adenosine inhibits electrical activity-driven calcium transients in GH cell lines (645). In frog melanotrophs, adenosine also inhibits spontaneous electrical activity (646), presumably reflecting the inhibitory action on Ca channels (647), facilitation of A-type K channels (151), and/or potentiation of delayed rectifier K channels (150).

**GABA receptors** are GABA- and baclofen-sensitive G subtypes, and their activation induces late hyperpolarization, attenuation of Ca currents, facilitation of Kv3 channels, and inhibition of AC activity (648). Pituitary melanotrophs express all three subtypes of these receptors [GABA(B1a), GABA(B1b), and GABA(B2)], and functional receptors were identified in postnatal and adult melanotrophs (649). Activation of these receptors leads to inhibition of AC activity (423) and spontaneous calcium oscillations (650), and inhibition in the calcium-dependent basal a-MSH release (328). These receptors are also expressed in anterior pituitary cells and contribute to the control of PRL and gonadotropin secretion in an age-dependent manner (544, 651, 652). In female rats that received estradiol implants for 5 wk, pituitary GABA receptor mRNA was significantly decreased compared with proestrous rats, and the baclofen-induced decrease in [Ca2+]i in pituitary cells was abolished (652). No details about the effects of these receptors on the electrical activity of anterior pituitary cells have been reported.

Pituitary cells from neonatal animals express the functional MT1 subtype of melatonin receptors that signal through pertussis toxin-sensitive G proteins. Their activation by melatonin leads to a decrease in cAMP production and PKA activity and attenuation of GnRH-induced gonadotropin secretion (545). Single-cell calcium and electrophysiological recordings revealed that the reduction in gonadotropin release results from melatonin-induced inhibition of both components of GnRH-induced calcium signaling in gonadotrophs, calcium influx through Ca channels, and IP3-mediated calcium release from intracellular stores (653–655). Inhibition of calcium influx by melatonin results in a delay of GnRH-induced calcium signaling. On the other hand, attenuation in GnRH-induced calcium release affects the amplitude of the calcium signals. The potent inhibition of GnRH-induced calcium signaling and gonadotropin secretion by melatonin provides an effective mechanism to protect premature initiation of pubertal changes that are dependent on gonadotropin plasma levels. During development, the tonic inhibitory effects of melatonin on GnRH action gradually attenuate, due to a decline in expression of functional MT1 receptors and changes in the GnRH receptor signaling pathways (656–658). In adult

4. **Other pituitary receptors linked to the G G signaling pathway**

Cloning of 5-HT receptors led to the recognition of several types of 5-HT-activated G proteins. All 5-HT1 receptors are negatively coupled to AC via G, whereas 5-HT4, 5-HT6, and 5-HT7 receptors stimulate AC through Gs (636). In addition to dopamine D2 receptors, porcine pituitary melanotrophs also express 5-HT1A and 5-HT1C receptors, and their activation leads to inhibition of L-type Ca channels (637). Inhibition of L-type and Q-type Ca channels mediated by 5-HT also occurs in rat melanotrophs (25). In both cell types, inhibition of Ca currents was abolished in cells treated with pertussis toxin, indicating the coupling of 5-HT receptors to the G signaling pathway. There are contradictory conclusions about direct actions of 5-HT on PRL release (638, 639).

Adenosine is a potent inhibitor of -MSH release from frog melanotrophs (640). The structure of these receptors has not been identified, but pharmacological, electrophysiological, and secretory data indicate the expression of adenosine receptors of the A subtype, which is negatively coupled to the AC signaling pathway through pertussis toxin-sensitive G proteins (461). Two reports have also indicated the operation of adenosine receptors in pituitary lactotrophs (641, 642), but further studies are required to clarify their structure, coupling, and effects (stimulatory or inhibitory) on PRL secretion. GH and GHGHc cell lines also express A1 receptors, and their activation causes inhibition of PRL and GH secretion (643, 644).
animals, melatonin does not affect pituitary functions directly, whereas the coupling between melatonin release and hypothalamic functions, including GnRH release, are preserved and are critically important in synchronizing the external photoperiods and reproductive functions through mechanisms that are not well characterized (545).

Neuropeptide Y is a 36-amino acid peptide mainly localized in the nervous system that exerts its biological actions through five receptors, called Y1 to Y5 (659). Pituitary lactotrophs, somatotrophs, and gonadotrophs probably express the Y1 receptor subtype, and the expression is regulated by estrogens in a cell-specific manner (660). In gonadotrophs, neuropeptide Y inhibits GnRH-induced Ca^{2+} signaling and LH release in a pertussis toxin-sensitive manner (661). The effects of neuropeptide Y on electrical activity in anterior pituitary cells have not been studied. Neuropeptide Y also inhibits spontaneous Ca^{2+} transients and the accompanying α-MSH release in melanotrophs (662). Electrophysiological experiments revealed that neuropeptide Y inhibits spontaneous electrical activity and Ca currents in these cells (663).

Galanin is produced by pituitary cells and acts as a paracrine factor (4). Pituitary cells express the GalR2 receptor subtype (664–666). In general, this receptor couples to the G_{q11} signaling pathway (667). However, the cloned receptor also couples to the G_{o} signaling pathway (668). The coupling of GalR2 receptors in pituitary cells has not been studied. Based on observations that galanin stimulates PRL release (669) and inhibits gonadotropin secretion (670), it is reasonable to suggest differential coupling of these receptors in pituitary cells. At the present time, there is no information about the effects of galanin on electrical activity and Ca^{2+} signaling in these cells.

VIII. Calcium-Mobilizing Receptors and Electrical Activity

GPCRs linked to the G_{q11} signaling pathway are activated by agonists in all anterior pituitary cell types and include: acetylcholine M1 and M3 receptors, angiotensin receptor AT1b, ATP-activated P2Y1 and P2Y2 receptors, ETA receptors, galanin receptor GalR2, ghrelin receptor GHS-R1a, GnRH receptor, serotonin 5-HT2A and 5-HT2B receptors, substance P receptor NK1, TRH receptor, AVP/oxytocin V1b and OT receptors, and VIP/PACAP receptor PAC1b (Fig. 12). In gonadotrophs, this signaling pathway is activated by GnRH (33, 671), which is the main agonist for these cells, as well as by ETs (52, 245), PACAP (36), substance P (39, 672), and AVP/oxytocin (38, 673). In thyrotrophs, the G_{q11} signaling pathway is activated by TRH, the main agonist for these cells (242, 671), and ETs (617). Lactotrophs express numerous Ca^{2+}-mobilizing receptors, activated by: acetylcholine (54), angiotensin II (244), ATP (474), ETs (52, 53), oxytocin (674), 5-HT (54), substance P (672, 675), TRH (242), and galanin (676–678). Mammalian melanotrophs express muscarinic receptors (679), and frog melanotrophs express Ca^{2+}-mobilizing receptors for TRH and neuropeptide Y (26), in addition to muscarinic receptors (27). In corticotrophs, the Ca^{2+}-mobilizing pathway is activated by AVP (176, 680, 681), norepinephrine (682), and potentially by 5-HT (398). Somatotrophs express Ca^{2+}-mobilizing ghrelin (46) and ETA (48) receptors. Several Ca^{2+}-mobilizing receptor tyrosine kinases are also expressed in pituitary cells (683–689), but their effects on Ca^{2+} signaling and electrical activity have not been studied in pituitary cells.

The activated G_{q11} protein leads to phosphoinositide hydrolysis and the production of IP_3 and DAG (690). IP_3 binds to IP_3Rs in the ER membrane and along with Ca^{2+} is required for their activation. Activation binding sites for both IP_3 and Ca^{2+} are on the cytoplasmic side of the membrane. As stated in Section III.C.1, there are three closely
related subtypes of the mammalian IP$_3$R, and the functional receptor forms a tetramer (691). It appears likely that each subunit must be bound by IP$_3$ and Ca$^{2+}$ for activation of the receptor (692). Once activated, the IP$_3$R functions as a Ca$^{2+}$ channel, allowing Ca$^{2+}$ to flow down its concentration gradient from the ER into the cytosol. The Ca$^{2+}$ flux can be terminated by inactivation of the receptor, which occurs through binding of Ca$^{2+}$ to an inactivation site on each subunit on the cytoplasmic side of the receptor. Thus, activation of one branch of the G$_q/11$ pathway leads to Ca$^{2+}$ mobilization from the ER store. The other branch follows the production of DAG, which together with Ca$^{2+}$ activates PKC (693).

A. The dynamics of Ca$^{2+}$ release

The ER is the primary storehouse for Ca$^{2+}$ in most cells, including pituitary cells, with a resting Ca$^{2+}$ concentration ([Ca$^{2+}]_{ER}$) of a few hundred micromolar (694, 695). This is in contrast to the resting level of [Ca$^{2+}]_{i}$, which is approximately 0.1 μM. The high [Ca$^{2+}]_{ER}$ is maintained by SERCA pumps. Efflux of Ca$^{2+}$ from the ER is through passive leakage and through IP$_3$Rs and/or RyRs (696). Because of the large concentration difference, the activation of IP$_3$Rs by a G$_q/11$ agonist results in a large and sudden increase in [Ca$^{2+}]_{i}$ (172, 173). After this initial Ca$^{2+}$ pulse, one of two behaviors is typically observed, depending on the cell type and in some cases on agonist. One behavior involves oscillations, whereas the other does not. In lactotrophs, somatotrophs, thyrotrophs, and cells from the GH cell lines, the pulse is typically followed by a slow decline to a plateau in [Ca$^{2+}]_{i}$, although some cells may only have a pulse or a plateau, and in a fraction of cells oscillations are observed (242, 282, 697, 698). PRL secretion from lactotrophs is increased during both the [Ca$^{2+}]_{i}$ pulse and the subsequent decay and plateau phases (282). In mammalian gonadotrophs, the pulse is typically followed by large [Ca$^{2+}]_{i}$ oscillations (172, 173, 245, 521, 699). Fish gonadotrophs also show an oscillatory Ca$^{2+}$ response to application of GnRH (700). However, αT3-1 (255) and LβT2 (42) mice gonadotrophs show nonoscillatory amplitude-modulated Ca$^{2+}$ signals in response to GnRH application. Corticotrophs respond to norepinephrine with extracellular Ca$^{2+}$-independent Ca$^{2+}$ oscillations (682). In contrast, stimulation of these cells with AVP results in the pulse-decay-plateau type of response (176, 680).

1. Pulse-decay-plateau Ca$^{2+}$ release

The pulse-decay-plateau Ca$^{2+}$ response requires only that the IP$_3$Rs open and remain open during agonist application. That is, the IP$_3$R is passive. Such response is illustrated with a mathematical model (701) in Fig. 13. The top panel shows [Ca$^{2+}]_{i}$ in response to agonist activation of the G$_q/11$ pathway and subsequent production of IP$_3$. The bottom panel shows [Ca$^{2+}]_{ER}$, a quantity that is difficult to measure experimentally. The initial rapid increase in [Ca$^{2+}]_{i}$, is followed by a slow decline, reflecting the removal of Ca$^{2+}$ from the cell by plasma membrane ATPase pumps and a Na$^+$-dependent Ca$^{2+}$ efflux. The decline in [Ca$^{2+}]_{i}$, is mirrored by a decline in [Ca$^{2+}]_{ER}$, although [Ca$^{2+}]_{ER}$ is much larger. As [Ca$^{2+}]_{ER}$ declines to a sufficiently low level, a Ca$^{2+}$ entry pathway is activated, bringing additional Ca$^{2+}$ into the cell and producing an elevated plateau in [Ca$^{2+}]_{i}$, that is evident near the end of the agonist application. When the agonist is removed, [Ca$^{2+}]_{i}$ initially declines to a subbasal level and slowly climbs back to a basal level. This drop and slow rise are due to the increased Ca$^{2+}$ flux into the ER that occurs as [Ca$^{2+}]_{ER}$ slowly returns to its basal level.

Notice a relatively rapid depletion of the ER Ca$^{2+}$ store in the presence of agonist in Fig. 13. In cells bathed in Ca$^{2+}$-deficient medium or with blocked VGCl, [Ca$^{2+}]_{ER}$, drops to basal levels within a few minutes, indicating that sustained Ca$^{2+}$ signaling by Ca$^{2+}$-mobilizing GPCRs is critically dependent on Ca$^{2+}$ influx. This is well illustrated in TRH-stimulated lactotrophs in cells bathed in Ca$^{2+}$-deficient medium and by cells stimulated with ET (a Ca$^{2+}$-mobilizing agonist that inhibits VGCl; see Section VII.B.3) in the presence of Ca$^{2+}$ (281, 282).

Although the [Ca$^{2+}]_{ER}$ of pituitary cells is not typically measured, Fig. 13 suggests that the time course of [Ca$^{2+}]_{ER}$ during agonist application is reflected in the [Ca$^{2+}]_{i}$ time course. That is, the rate of decline in [Ca$^{2+}]_{i}$, after the initial peak is determined largely by the time dynamics of [Ca$^{2+}]_{ER}$. If the decline of [Ca$^{2+}]_{ER}$ is more
rapid, then so too will be the decline in $[\text{Ca}^{2+}]_i$. If factors such as the total IP$_3$R conductance activated by agonist or the Ca$^{2+}$ leakage rate are larger in one cell than a second cell, then the decline in $[\text{Ca}^{2+}]_i$ and $[\text{Ca}^{2+}]$ will be more rapid in the first cell. In contrast, if the SERCA pumping rate is larger in one cell than another, then the decline in $[\text{Ca}^{2+}]_i$ and $[\text{Ca}^{2+}]$, will be slower in the first cell. Thus, the decline in the readily measured $[\text{Ca}^{2+}]_i$, is an assay for the typically unmeasured $[\text{Ca}^{2+}]_i$ that could also be used in further work with pituitary cells.

The Ca$^{2+}$ response to a $G_{q/11}$-activating agonist can have a large impact on the plasma membrane potential. The pulse of Ca$^{2+}$ that follows agonist application activates SK type K$_{Ca}$ channels in the plasma membrane in rat somatotrophs, lactotrophs, corticotrophs, and GH$_3$ cells (171, 680, 704). The K$_{Ca}$ current hyperpolarizes the membrane, terminating any spontaneous electrical activity that was present before agonist application (228, 282). Some time after the initial Ca$^{2+}$ pulse, the membrane typically depolarizes, due to the modulation of a still-unidentified current, presumably the down-regulation of an M (153) or an erg current (155, 158, 161). This depolarization activates Ca$_v$ channels, further depolarizing the cell and initiating electrical activity such as spiking or bursting (282). This electrical activity would then be reflected in the $[\text{Ca}^{2+}]$, time course as small oscillations on top of the plateau in Fig. 13A and would contribute to the plateau. In studies done in the absence of extracellular Ca$^{2+}$, the $[\text{Ca}^{2+}]$, declines to below its basal level even while the agonist (TRH) is present (698). In this case, neither capacitative Ca$^{2+}$ entry nor Ca$_v$ currents provide the Ca$^{2+}$ influx required for the $[\text{Ca}^{2+}]$, plateau or small oscillations after the initial surge in $[\text{Ca}^{2+}]$.

Thus, VCGI represents the major pathway for the plateau Ca$^{2+}$ response during the sustained agonist stimulation and for recovery of the ER calcium pool after removal of agonist, but other pathways could also be operative. Although there is some evidence supporting the presence of store-operated Ca$^{2+}$ entry in pituitary cells (272, 698), the data are inconclusive. In other cell types, it is also well established that Ca$^{2+}$-mobilizing receptors can activate the TRP family of channels, which conduct Ca$^{2+}$ and also depolarize the cells, leading to facilitation of VCGI (219). Further studies are needed to clarify whether this pathway is activated by Ca$^{2+}$-mobilizing receptors in pituitary cells.

Sustained activation of Ca$^{2+}$-mobilizing GPCRs also causes changes in the gating properties of plasma membrane channels. In GH$_3$ cells, the Ca$^{2+}$ released from the ER initially inactivates a Ca$_v$ current, but this phase is followed by stimulation of the Ca$_v$ current, which can contribute to the plateau (705). Both $G_i$ ($\alpha_2$ and $\alpha_3$) and PKC are required for this TRH-induced stimulation of a Ca$_v$ current (706). In thyrotrophs and lactotrophs, a return to tonic spiking at higher frequency was accompanied with lower spike amplitude in the presence of TRH due to inhibition of L-type Ca$_v$ channels (707).

**2. IP$_3$R-mediated Ca$^{2+}$ oscillations**

Unlike the spontaneous Ca$^{2+}$ oscillations that often occur in pituitary cells, those induced by GnRH in gonadotrophs or norepinephrine in corticotrophs persist when the bath Ca$^{2+}$ is removed as well as in cells bathed in the presence of Ca$^{2+}$ but clamped at potentials that silence Ca$^{2+}$ influx through Ca$_v$ channels (283, 682). This illustrates that the oscillation is intrinsic to the Ca$^{2+}$ handling properties within the cell. There are differences in the oscillatory Ca$^{2+}$ mobilization between these two cell types. In gonadotrophs, the oscillatory Ca$^{2+}$ release is activated not only by GnRH but also by ET-1, PACAP, and AVP (172, 173, 245, 521, 699). In contrast, baseline Ca$^{2+}$ oscillations are triggered by $\alpha$-adrenergic stimulation of corticotrophs but not AVP application (682). Furthermore, the frequency of Ca$^{2+}$ oscillations in gonadotrophs is determined by agonist concentration and varies between three and 20 pulses per minute (253, 708), whereas norepinephrine generates Ca$^{2+}$ oscillations with a frequency of about one per minute (682).

In gonadotrophs, oscillations in IP$_3$ are not required to generate oscillatory Ca$^{2+}$ release as documented by injection of nonmetabolized IP$_3$ analogs. Furthermore, the concentration of IP$_3$ underlines the frequency of Ca$^{2+}$ spiking (249). The $[\text{Ca}^{2+}]$, influences IP$_3$-dependent Ca$^{2+}$ release in these cells. The rapid stimulatory effect of Ca$^{2+}$ on IP$_3$-dependent Ca$^{2+}$ release in gonadotrophs is nicely demonstrated by phase resetting of GnRH-induced oscillations by a brief pulse of Ca$^{2+}$ entry (250). The inhibitory effect of high $[\text{Ca}^{2+}]$, on GnRH-induced Ca$^{2+}$ oscillations is also shown (251). Finally, inhibition of SERCA pumps causes a transition from the oscillatory to the nonoscillatory mode of Ca$^{2+}$ release in GnRH-stimulated gonadotrophs (709, 710). At the present time, it is unknown whether nonoscillatory elevation in intracellular IP$_3$ levels could generate Ca$^{2+}$ oscillations in corticotrophs.

**3. Mathematical models of IP$_3$-induced Ca$^{2+}$ oscillations**

A series of mathematical models were developed to help understand the mechanism for $G_{q/11}$ agonist-induced Ca$^{2+}$ oscillations in gonadotrophs and how these depend on Ca$^{2+}$ flux across the plasma membrane and between the cytosol and the ER (252, 711–714). These models all assume that the key player in the Ca$^{2+}$ oscillations is an active IP$_3$R (called class I models in Ref. 715). These models also assume no role for mitochondria in the oscillations, which is contrary to some data (but the models can
easily be adapted, as discussed in Section VIII.A.4). For simplicity, we use models described in Ref. 716 to illustrate the mechanism for Ca\textsuperscript{2+} oscillations generated by an active IP\textsubscript{3}R.

There are two essential ingredients to agonist-induced Ca\textsuperscript{2+} oscillations mediated by an active IP\textsubscript{3}R. One is a Ca\textsuperscript{2+} concentration difference between the cytosol and ER, which is maintained by SERCA pumps through ATP hydrolysis. The other is an IP\textsubscript{3}R that is rapidly coactivated by IP\textsubscript{3} and cytosolic Ca\textsuperscript{2+} and more slowly inactivated by cytosolic Ca\textsuperscript{2+}. In the equilibrium state, this latter feature leads to a bell-shaped dependence on the [Ca\textsuperscript{2+}], (717, 718). More importantly for Ca\textsuperscript{2+} oscillations, there is a substantial time scale difference between Ca\textsuperscript{2+} activation and Ca\textsuperscript{2+} inactivation of the IP\textsubscript{3}R. The former provides positive feedback and is responsible for the upstroke of each Ca\textsuperscript{2+} spike. The latter provides delayed negative feedback and is responsible for the downstroke. This combination of rapid positive feedback and delayed negative feedback is similar to what occurs during the production of APs, where the source of the feedback is the membrane potential acting through Na\textsubscript{v} and K\textsubscript{v} channels. This analogy led to a significant simplification in a model of the IP\textsubscript{3}R dynamics and facilitates understanding of the oscillation mechanism (713).

We illustrate the basic components of the Ca\textsuperscript{2+} oscillations with a closed-cell model, neglecting any flux of Ca\textsuperscript{2+} across the plasma membrane (intracellular Ca\textsuperscript{2+} is conserved). This allows us to focus solely on the IP\textsubscript{3}R-mediated dynamics. Figure 14A shows a closed-cell model simulation of Ca\textsuperscript{2+} oscillations that occur only when IP\textsubscript{3} is present. The variable $h$ (dashed curve) is the fraction of IP\textsubscript{3}Rs not inactivated; $b = 1$ means that no IP\textsubscript{3}Rs are inactivated. At the beginning of the simulation, the $G_{q/11}$ pathway has not been activated, so IP\textsubscript{3} is below the threshold level needed for activation of IP\textsubscript{3}Rs. In this unstimulated case, $b$ is approximately 1 and [Ca\textsuperscript{2+}] is low and steady. When IP\textsubscript{3} is present at a critical level, the system produces a periodic train of Ca\textsuperscript{2+} spikes. The upstroke of each spike is driven by fast Ca\textsuperscript{2+} activation of IP\textsubscript{3}Rs, and during this time the variable $h$ declines as some of the IP\textsubscript{3}Rs become inactivated. As $h$ declines, so too does the total flux through the population of IP\textsubscript{3}Rs, resulting in a reduction in the net Ca\textsuperscript{2+} flux out of the ER. Eventually, the net flux becomes negative because the flux into the ER through SERCA pumps exceeds the efflux through IP\textsubscript{3}Rs. This produces a decline in [Ca\textsuperscript{2+}], which is the downstroke of the Ca\textsuperscript{2+} spike. The cytosolic Ca\textsuperscript{2+} level then returns to a low level, allowing the IP\textsubscript{3}Rs to slowly recover from inactivation. This deactivation process is reflected in Fig. 14A as a slow increase in $h$ after each spike. When the receptors recover sufficiently, a new spike is initiated, and the process repeats as long as IP\textsubscript{3} is present. The spiking in [Ca\textsuperscript{2+}], would be reflected in small-amplitude oscillations in the (much larger) [Ca\textsuperscript{2+}]\textsubscript{ER}, as has been measured in gonadotrophs (719). The production of each Ca\textsuperscript{2+} spike depends critically on the time scale difference between activation (fast) and inactivation (slow). If the inactivation is made too fast in the model, no oscillations are produced. Also, the period of the Ca\textsuperscript{2+} oscillations is determined by the time required for the IP\textsubscript{3}Rs to recover from inactivation. Thus, the time scale for the inactivation process determines whether or not oscillations are produced and the period of the oscillations.

Although IP\textsubscript{3}R-mediated Ca\textsuperscript{2+} oscillations can be generated in a closed cell, in actual cells there is influx of Ca\textsuperscript{2+} through Ca\textsubscript{v}, and other channels, as well as efflux of Ca\textsuperscript{2+} through plasma membrane pumps. The effect that this Ca\textsuperscript{2+} flux has on IP\textsubscript{3}R-mediated oscillations is illustrated in Fig. 14B. In this figure, produced by an open-cell model, there is a constant Ca\textsuperscript{2+} flux into the cell and efflux that depends on [Ca\textsuperscript{2+}], (efflux is greater when [Ca\textsuperscript{2+}] is greater). When the $G_{q/11}$ pathway is activated (Fig. 14B, gray bar), the Ca\textsuperscript{2+} oscillations start and, after an initial large spike, continue with a constant amplitude and frequency. However, when Ca\textsuperscript{2+} influx is stopped ($J_{in} = 0$), simulating the removal of extracellular Ca\textsuperscript{2+}, the oscillation amplitude and frequency get progressively smaller. If continued further in time, the oscillations would terminate altogether. The reason for this behavior is that each Ca\textsuperscript{2+} spike transfers Ca\textsuperscript{2+} from the ER to the cytosol, from where some fraction is pumped out of the cell. When this is accompanied by Ca\textsuperscript{2+} influx through plasma membrane channels ($J_{in} > 0$), the ER refills between spikes, so at the

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**FIG. 14.** Model simulation of one mechanism for Ca\textsuperscript{2+} oscillations that can be produced by $G_{q/11}$ activation. A. Oscillations are due to the fast activation and slow inactivation of IP\textsubscript{3}Rs. The variable “$h$” is the fraction of receptors that are not inactivated. B. Ca\textsuperscript{2+} oscillations persist as long as there is sufficient Ca\textsuperscript{2+} influx into the cell. When influx is eliminated ($J_{in} = 0$), the oscillation amplitude and frequency decline, and eventually oscillations cease.
start of the next spike the driving force will be the same as it was for the prior spike. However, when the influx through the plasma membrane is eliminated \( (I_{in} = 0) \), there is a progressive loss of \( \mathrm{Ca}^{2+} \) from the cell, and therefore the ER does not refill completely between spikes. For this reason, the driving force is smaller for each subsequent spike, resulting in a slow decline in the spike amplitude. Also, because of the progressive decline in the driving force, the fraction of \( \mathrm{IP}_3 \)-Rs that must recover from inactivation for the start of a new spike gets progressively larger. That is, \( b \) must rise to progressively larger values to trigger spikes, so the time between spikes increases (frequency decreases). These behaviors, a slow decline in spike amplitude and spike frequency, are observed in GnRH-stimulated gonadotrophs in a \( \mathrm{Ca}^{2+} \)-deficient medium (252, 289).

Although an active \( \mathrm{IP}_3 \)R provides a good explanation for agonist-induced \( \mathrm{Ca}^{2+} \) oscillations in gonadotrophs (and other cells in which similar oscillations are produced), \( \mathrm{Ca}^{2+} \) oscillations may also be produced through the stimulatory feedback of \( \mathrm{Ca}^{2+} \) onto PLC (720), which produces \( \mathrm{IP}_3 \), or onto 3-kinase (721), which converts \( \mathrm{IP}_3 \) to \( \mathrm{IP}_4 \). With this mechanism (class 2 models) (715), oscillation in the \( \mathrm{IP}_3 \) concentration is a key element, unlike the case described above where oscillations in \( \mathrm{Ca}^{2+} \) are produced for a constant \( \mathrm{IP}_3 \) concentration (class 1 models). These two classes of models for \( \mathrm{G}_{\mathrm{q/11}} \)-agonist-induced oscillations have been recognized for many years (722, 723). It is difficult to determine from inspection only which mechanism is responsible for oscillations in a given cell, and there is evidence that the oscillation class is determined by the type of GPCR activated (724). Consistent with this, norepinephrine generates \( \mathrm{Ca}^{2+} \) oscillations in corticotrophs, whereas AVP does not. The slow frequency of \( \mathrm{Ca}^{2+} \) oscillations in corticotrophs may suggest that these oscillations were mediated by oscillatory \( \mathrm{IP}_3 \) production.

4. Role of mitochondria in \( \mathrm{IP}_3 \)-induced \( \mathrm{Ca}^{2+} \) release

Another player in the production of depolarization- and agonist-induced \( \mathrm{Ca}^{2+} \) transients in pituitary cells is the mitochondrial \( \mathrm{Ca}^{2+} \) store. Calcium is transported into mitochondria through \( \mathrm{Ca}^{2+} \) uniporters, which are powered by the membrane potential across the inner membrane. Calcium is transported out of mitochondria primarily by \( \mathrm{Na}^+ \)/\( \mathrm{Ca}^{2+} \) exchangers (725). It has been demonstrated that these actions impact the cytosolic \( \mathrm{Ca}^{2+} \) time course in neurons (726–728). In corticotrophs, the rate of \( \mathrm{Ca}^{2+} \) clearance after depolarization-induced \( \mathrm{Ca}^{2+} \) influx is dramatically slowed by mitochondrial uncouplers or inhibitors of the mitochondrial uniporter. This in turn enhances the exocytotic response (729). In gonadotrophs, early work by Hille’s group (719) revealed that \( \mathrm{Ca}^{2+} \) released from the ER is partly taken up by the ER and partly pumped into other intracellular compartments or out of the cells. Subsequent studies by the same group showed that collapsing the mitochondrial inner membrane potential with the protonophore carbonyl cyanide m-chlorophenylhydrazone, a manipulation that inhibits \( \mathrm{Ca}^{2+} \) uptake into mitochondria, slowed or inhibited GnRH-induced cytosolic \( \mathrm{Ca}^{2+} \) oscillations (730, 731).

Although these data demonstrate that mitochondrial \( \mathrm{Ca}^{2+} \) filtering plays a role in the generation of agonist-induced \( \mathrm{Ca}^{2+} \) oscillations, they do not allow one to determine whether this role is active or passive. That is, it is not evident whether the oscillations in mitochondrial \( \mathrm{Ca}^{2+} \) content are required for the production of cytosolic oscillations (mitochondrial \( \mathrm{Ca}^{2+} \) is an active player), or whether cytosolic oscillations would persist if the mitochondrial \( \mathrm{Ca}^{2+} \) level could be clamped at its mean level in the stimulated cell (mitochondrial \( \mathrm{Ca}^{2+} \) is a passive player). Although oscillations in mitochondrial \( \mathrm{Ca}^{2+} \) concentration have been measured in gonadotrophs (730), it is not known whether these oscillations are required for the generation of cytosolic \( \mathrm{Ca}^{2+} \) oscillations. In a mathematical modeling study, a \( \mathrm{Ca}^{2+} \) oscillation model was modified so that oscillations were produced only if a mitochondrial \( \mathrm{Ca}^{2+} \) store was present (732). However, the mitochondrial store played a passive role because \([\mathrm{Ca}^{2+}]_c \) oscillations were produced even when the mitochondrial \( \mathrm{Ca}^{2+} \) concentration was clamped at its mean stimulated value (R. Bertram, unpublished observation).

5. Coupled membrane and \( \mathrm{IP}_3 \)-mediated oscillations

In two studies of GnRH-stimulated rat gonadotrophs, the membrane was voltage clamped, and the frequency of the oscillations in \( \mathrm{Ca}^{2+} \) concentration was measured (269, 283). This procedure allows one to separate the \( \mathrm{IP}_3 \)R-mediated oscillation from the intrinsic membrane oscillation discussed in Section III. It also provides a means to control the rate of \( \mathrm{Ca}^{2+} \) influx by adjusting the holding potential of the cell. When the holding potential is low (hyperpolarized), few voltage-dependent \( \mathrm{Ca}^{2+} \) channels are open, and the \( \mathrm{Ca}^{2+} \) current is small. For larger (depolarized) holding potentials, many \( \mathrm{Ca}^{2+} \) channels are open, yielding a larger \( \mathrm{Ca}^{2+} \) current.

Besides demonstrating that \( \mathrm{Ca}^{2+} \) oscillations persist in the absence of membrane potential oscillations, the study demonstrated that oscillations died out if the holding potential was not sufficiently depolarized. This could be explained by a gradual depletion of the ER due to insufficient replenishment of \( \mathrm{Ca}^{2+} \). In other words, redistribution of \( \mathrm{Ca}^{2+} \) between ER and mitochondrial pools is not sufficient to prevent the depletion of the intracellular \( \mathrm{Ca}^{2+} \) in oscillating cells. However, in cells clamped at −60 mV, GnRH-induced \( \mathrm{Ca}^{2+} \) oscillations last for 6–15 min, much longer than \( \mathrm{Ca}^{2+} \) signals in voltage-clamped cells respons-
ing to activation of Ca\(^{2+}\)-mobilizing receptors with a pulse-decay-plateau Ca\(^{2+}\) release. This indicates that depletion of the intracellular Ca\(^{2+}\) takes longer in cells with oscillatory Ca\(^{2+}\) release (283). As stated above, two Ca\(^{2+}\)-handling mechanisms are operative in GnRH-stimulated gonadotrophs: redistribution of Ca\(^{2+}\) within ER and mitochondrial pools, and a Na\(^{+}\)-dependent Ca\(^{2+}\) efflux followed by Ca\(^{2+}\) influx. Thus, it is reasonable to suggest that GnRH-induced baseline Ca\(^{2+}\) oscillation makes gonadotrophs less dependent on Ca\(^{2+}\) influx, in contrast to agonist stimulated lactotrophs and somatotrophs. Consistent with this, our work with neonatal gonadotrophs showed that redistribution of Ca\(^{2+}\) within the cells dominates in GnRH-stimulated cells exhibiting baseline Ca\(^{2+}\) oscillations, whereas removal of Ca\(^{2+}\) from the cells dominates in GnRH-stimulated cells showing a prolonged Ca\(^{2+}\) spike, similar to those observed in agonist-stimulated lactotrophs and somatotrophs (256).

Experiments with cells voltage-clamped at different potentials also showed that the Ca\(^{2+}\) spike frequency increased with increases in the holding potential (for potentials between −60 and −20 mV). The interpretation of this, reproduced with our model, is that the greater Ca\(^{2+}\) influx provided by increased membrane depolarization fills the ER to higher levels, increasing the driving force for the Ca\(^{2+}\) spikes so that \(h\) does not have to rise as high to initiate a new spike. Thus, spikes are produced at shorter time intervals (269, 283).

From the discussion above, it is also evident that Ca\(^{2+}\) oscillations produced by agonist-stimulated gonadotrophs only require Ca\(^{2+}\) flux across the plasma membrane to keep the ER-Ca\(^{2+}\) store replenished; no patterned electrical activity is required. This is unlike other endocrine cells, where Ca\(^{2+}\) oscillations are due to bursting electrical activity (70, 184, 291, 293, 294, 733). However, stimulated gonadotrophs do produce electrical bursting, due to the bidirectional interactions between the plasma membrane and the ER (173, 289, 711, 734). Ion channels in the plasma membrane bring Ca\(^{2+}\) into the cell during each spike, which replenishes the ER and thereby provides coupling from the membrane to the ER.

Coupling from the ER to the plasma membrane is mediated through Ca\(^{2+}\)-activated SK channels (70, 72, 172, 173, 176). In mouse gonadotrophs, BK channels also contribute to the hyperpolarization of the plasma membrane (174). As discussed in Section VIII.A.1, this bidirectional coupling is also present in thyrotrophs and lactotrophs, but in agonist-activated gonadotrophs there are intrinsic activity oscillations in both the plasma membrane and the ER. Furthermore, after the hyperpolarization, gonadotrophs do not return to the tonic spiking state that typically characterizes the unstimulated cell, but instead they exhibit a bursting pattern consisting of tall electrical spikes clustered into episodes. This bursting behavior is illustrated in Fig. 15 using a simple mathematical model from Ref. 716; a more detailed model can be found in Ref. 711. The key feature is the antiphase pattern of electrical activity and Ca\(^{2+}\) spikes. This is due to the inhibitory action of each Ca\(^{2+}\) spike on the plasma membrane; each Ca\(^{2+}\) spike activates a Ca\(^{2+}\)-activated K\(^{+}\) current, which terminates electrical spiking. The electrical spiking resumes once [Ca\(^{2+}\)]\(_{i}\) returns to a low level after the Ca\(^{2+}\) spike. Thus, the Ca\(^{2+}\) oscillator periodically interrupts the plasma membrane oscillator, producing a bursting pattern of electrical activity. Thus, the electrical activity and secretion are out of phase; the former serves to refill the ER, which provides the periodic Ca\(^{2+}\) pulse needed to evoke secretion. However, such a pattern of GnRH-induced electrical activity still does not protect the L-type Ca\(_{\text{v}}\) channels from Ca\(^{2+}\)-dependent inactivation, resulting in smaller amplitude of this current (735).

B. Calcium mobilization and secretion

The release of neurotransmitter from neuronal synapses occurs when Ca\(^{2+}\) that enters through channels in the plasma membrane binds to nearby release sites. This secretion is rapid (less than 1 msec), and the speed of exocytosis is critical for neuronal functions. Thus, it is the localized high-concentration Ca\(^{2+}\) nanodomains that gate release (346, 736), not the bulk cytosolic Ca\(^{2+}\). This allows single APs to evoke transmitter release, although the change in the mean [Ca\(^{2+}\)]\(_{i}\) is small. In contrast, secretion in pituitary cells is slow. Initially, it was postulated that one of three reasons could underlie such slow secretion in endocrine cells: that Ca\(_{\text{v}}\) channels and secretory vesicles are not molecularly colocalized; that the exocytotic machinery is intrinsically slower; or that secretory vesicles are

![FIG. 15. Model simulation of coupled IP\(_3\)R and membrane oscillations.](image-url)
not in close proximity to the plasma membrane to form the fusion pore (342).

As discussed in Section IV.E.3, spontaneous electrical activity in corticotrophs, gonadotrophs, and thyrotrophs does not trigger prominent secretion. In these cells, facilitation of VGCI by Bay K 8644, an L-type Ca\(^{2+}\) channel activator, and high K\(^+\)-induced depolarization of cells generates large amplitude Ca\(^{2+}\) signals and stimulates ACTH, LH, PRL, and GH secretion (345, 737). In melano- 
notrophs, lactotrophs, and somatotrophs, spontaneous electrical activity is sufficient to activate the exocytotic pathway, but in these cells spontaneous APs generate large-amplitude global Ca\(^{2+}\) signals. Single-cell recordings showed that pituitary cells start secreting at submimicromolar [Ca\(^{2+}\)] (738) and therefore have a high Ca\(^{2+}\) affinity (342). This indicates that in pituitary cells secretory vesicles are not colocalized with Ca\(^{2+}\) channels and that global Ca\(^{2+}\) signals are needed to initiate fusion of vesicles with the plasma membrane.

Consistent with this view, activation of Ca\(^{2+}\)-mobilizing receptors in all endocrine pituitary cell types generates global Ca\(^{2+}\) signals and stimulates secretion. It appears that the pattern of Ca\(^{2+}\) mobilization (pulse-decay-plateau response vs. baseline Ca\(^{2+}\) oscillations) is not critical for activation of the exocytotic pathway, as is shown in experiments with rat corticotrophs in which both AVP (the pulse-decay-plateau response) (176) and norepinephrine (baseline oscillations) elicit secretion (682). Also, in L\(\beta\)T2 gonadotrophs, GnRH induces Ca\(^{2+}\) oscillations and hormone secretion, as monitored by measurements of plasma membrane capacitance (42). In rat gonadotrophs, the frequency of the oscillations and level of secretion are dependent on the GnRH level, so there is frequency coding in the Ca\(^{2+}\) response. Furthermore, activation of the exocytotic pathway in GnRH-stimulated gonadotrophs has nothing to do with the bursting electrical pattern exhibited by stimulated cells because secretion occurs during the Ca\(^{2+}\) pulses, when the plasma membrane is hyperpolarized (739, 740). These and other data indicate that two factors contribute to the effective coupling of Ca\(^{2+}\)-mobilizing receptors to secretion in endocrine pituitary cells. First, Ca\(^{2+}\)-mobilizing receptors in pituitary cells generate high amplitude global Ca\(^{2+}\) signals. Second, in corticotrophs and gonadotrophs, the majority of the secretion is from release sites colocalized with IP\(_3\)Rs in the ER (739). In corticotrophs, both intracellular Ca\(^{2+}\) release and VGCI generate a spatial Ca\(^{2+}\) gradient, such that the local [Ca\(^{2+}\)] near the exocytotic sites is about 3-fold higher than the mean [Ca\(^{2+}\)] (741). Thus, the plume of high Ca\(^{2+}\) concentration that forms near a single IP\(_3\)R or a cluster of IP\(_3\)Rs gates exocytosis. This hypothesis could also account for the finding that Ca\(^{2+}\) evokes exocytosis from mela- 
notrophs more effectively when it is released from the ER than when it is introduced through a recording pipette (742). Also, in pancreatic acinar cells, both exocytosis and Ca\(^{2+}\) release from the ER through IP\(_3\)Rs occur at the apical pole of the cell, suggesting some degree of colocalization of release sites and IP\(_3\)Rs (743).

Ca\(^{2+}\)-mobilizing receptors trigger not only Ca\(^{2+}\) signaling but also several other intracellular signaling pathways, which also contribute to the effectiveness of stimulus secretion coupling. One of the major signaling molecules contributing to the control of exocytosis during activation of Ca\(^{2+}\)-mobilizing receptors is DAG. In association with Ca\(^{2+}\), DAG activates PKC, which has been shown to have several effects on gonadotrophs. In cells stimulated by GnRH, activation of PKC by application of phorbol 12-myristate 13-acetate (PMA) slowed down the Ca\(^{2+}\) oscillations and enhanced the SK current (175). These two effects appear to be independent because when unstimulated cells were loaded with IP\(_3\), the PMA had no effect on the frequency of Ca\(^{2+}\) oscillations, yet still enhanced the SK current induced by the Ca\(^{2+}\) oscillations. This suggests that PKC directly enhances SK current and also has an effect on PLC. Another study showed that PMA-activated PKC reduces Ca\(^{2+}\) influx in response to depolarization in gonadotrophs (241). This same study also found that depolarization-mediated secretion was enhanced by PKC, despite the reduction in Ca\(^{2+}\) influx. It is thus apparent that PKC has an action downstream of Ca\(^{2+}\) entry that amplifies secretion.

In chromaffin cells (744) and hippocampal neurons (745), the activation of PKC by phorbol esters or DAG increases the size of the readily releasable pool of vesicles by increasing the rate at which the pool is refilled. To see whether this was the case in gonadotrophs, flash photolysis was used to uncage Ca\(^{2+}\), and both the membrane capacitance and the intracellular Ca\(^{2+}\) level were measured (746). With these measurements, the Ca\(^{2+}\) dependence of exocytosis was established, in both control cells and those in the presence of PMA. There was a substantial left shift of the response curve when PMA was present, but no significant change in the calculated size of the readily releasable pool. Thus, it appears that PKC enhances secretion from gonadotrophs by sensitizing the secretory machinery to Ca\(^{2+}\) (746). This is not totally at odds with the actions of PKC on chromaffin cells and hippocampal neurons because the fact that the readily releasable pool is unchanged by PMA in the face of increased secretion suggests that the rate of refilling of the pool is enhanced by the PMA, as it is in chromaffin cells and hippocampal neurons. The sensitizing actions of PKC in gonadotrophs could account for the finding that secretion is greater when [Ca\(^{2+}\)] is elevated by application of GnRH rather than by
pipette or through photolysis of caged Ca\textsuperscript{2+} (739) because GnRH would activate PKC via DAG. In lactotrophs, PMA increases both granule-to-granule and granule-to-plasma membrane fusion events, which could account for potentiation of secretion by the PKC-dependent pathway (535).

Activation of GnRH receptors in gonadotrophs also leads to stimulation of phospholipase D (747, 748) in a PKC-dependent manner (748). This enzyme stimulates production of phosphatidic acid, which subsequently can be dephosphorylated to DAG by phosphatidate phosphohydrolase to sustain activation of PKC. In chromaffin cells, phosphatidic acid participates in the control of exocytosis (749). Also, in GT1 cells, the phospholipase D signaling pathway contributes to GnRH release (750). Further studies are needed to clarify the relevance of this signaling pathway in secretion from endocrine pituitary cells. One of the mechanisms could be activation of stimulus-transcription coupling by the phospholipase D signaling pathway (751) and facilitation of de novo formation of secretory vesicles. The sustained GnRH-stimulated and VGCI-dependent LH release is completely blocked by wortmannin at concentrations that inhibit phosphatidylinositol 4-kinase, an enzyme that participates in inositol phosphate metabolism (752), suggesting a potential role of PIP\textsubscript{2} in VGCI-dependent gonadotropin secretion. The steroid background also has a profound effect both on stimulus-induced Ca\textsuperscript{2+} mobilization and on Ca\textsuperscript{2+}-sensitivity of exocytosis (42, 753–756).

**IX. Summary**

The introduction of patch clamp techniques (757) was critical in the electrophysiological characterization of numerous voltage-gated Na\textsuperscript{+}, Ca\textsuperscript{2+}, and K\textsuperscript{+} channels in endocrine pituitary cells and their roles in spontaneous electrical activity, as well as in the characterization of receptor channels expressed in pituitary cells. GPCR-controlled electrical activity in single pituitary cells was also extensively studied using patch clamp techniques. The discovery of fluorescent dyes that are suitable for intracellular single-cell recordings, such as Indo-1 and Fura-2 (758), helped with the study of both the AP-driven rise in [Ca\textsuperscript{2+}], and the IP\textsubscript{3}-driven Ca\textsuperscript{2+} release from ER in pituitary cells. Simultaneous measurements of currents and [Ca\textsuperscript{2+}], or membrane potential and [Ca\textsuperscript{2+}], in pituitary cells were important in characterizing the relationship between spontaneous and receptor-controlled electrical activity and Ca\textsuperscript{2+} signaling, as well as synchronization between Ca\textsuperscript{2+} mobilization and electrical activity. The discovery of fluorescent dyes for measurements of single-cell exocytosis (759) also complemented patch clamp capacitative measurements in studies on Ca\textsuperscript{2+}-dependent hormone secretion (760).

These techniques helped to establish that voltage-gated channels provide a basic signaling system for individual pituitary cells. Like neurons, endocrine pituitary cells express numerous voltage-gated Na\textsuperscript{+}, K\textsuperscript{+}, and Ca\textsuperscript{2+}-conducting channels, as well as cation-conducting cyclic nucleotide-modulated and TRP channels. In contrast to neurons, propagation of APs within endocrine pituitary cells is unlikely to occur, and the main function of APs in pituitary cells is to provide a driving force for Ca\textsuperscript{2+} influx through Ca\textsubscript{v} channels. It is now well established that in three of the six endocrine pituitary cells—lactotrophs, somatotrophs, and melanotrophs—spontaneous electrical activity is sufficient to trigger hormone release in the absence of any stimulus. In gonadotrophs, thyrotrophs, and corticotrophs, spontaneous electrical activity is probably not coupled to secretion, at least in a majority of cells in vitro. It is reasonable to conclude that spontaneous electrical activity maintains these cells in a responsive state with [Ca\textsuperscript{2+}] near the threshold level. It is also interesting to note that this division of cells into two groups is not consistent with the embryonic development of endocrine cells (Fig. 1). The pattern of electrical activity (single spikes vs. pseudo-plateau bursting) and the frequency of firing determine the AP secretion coupling in single cells. The channels participating in spike depolarization and repolarization are relatively well characterized, whereas further work is needed to identify channels responsible for the pacemaking activity.

The status of the electrical signaling system in pituitary cells in vivo is critically dependent on the release of stimulating and inhibiting neurohormones from the hypothalamus. These agonists act on G\textsubscript{S}- and G\textsubscript{i/o}-coupled receptors expressed in pituitary cells (Fig. 16). Not accidentally, the inhibitory G\textsubscript{i/o}-coupled receptors are expressed predominantly in cells in which spontaneous electrical activity is sufficient to trigger hormone secretion; gonadotrophs express melatonin receptors, but only during embryonic and neonatal life. It has also been established that inhibition of spontaneous electrical activity, not inhibition of AC activity, accounts for down-regulation of basal PRL, GH, and α-MSH release. Solid evidence was obtained to support the concept that GPCRs inhibit electrical activity by inhibiting Ca\textsubscript{v} channels and/or stimulating K\textsubscript{v} channels. In contrast, activation of G\textsubscript{S}-coupled receptors leads to stimulation of electrical activity. Figure 16 shows that the G\textsubscript{S} signaling pathway plays a major role in only two cell types: somatotrophs and corticotrophs. This signaling system changes the pattern of electrical activity by facilitating unidentified Na\textsuperscript{+}-conducting channels and by facilitating Ca\textsubscript{v} conductance, both in a PKA-dependent manner. The relevance of potential phosphorylation of TTX-sensitive Na\textsuperscript{+} channels and cAMP-dependent activation of HCN channels in pituitary cells was critical in the electrophysiological characterization of numerous voltage-gated Na\textsuperscript{+}, Ca\textsuperscript{2+}, and K\textsuperscript{+} channels in endocrine pituitary cells and their roles in spontaneous electrical activity, as well as in the characterization of receptor channels expressed in pituitary cells. GPCR-controlled electrical activity in single pituitary cells was also extensively studied using patch clamp techniques. The discovery of fluorescent dyes that are suitable for intracellular single-cell recordings, such as Indo-1 and Fura-2, helped with the study of both the AP-driven rise in [Ca\textsuperscript{2+}], and the IP\textsubscript{3}-driven Ca\textsuperscript{2+} release from ER in pituitary cells. Simultaneous measurements of currents and [Ca\textsuperscript{2+}], or membrane potential and [Ca\textsuperscript{2+}], in pituitary cells were important in characterizing the relationship between spontaneous and receptor-controlled electrical activity and Ca\textsuperscript{2+} signaling, as well as synchronization between Ca\textsuperscript{2+} mobilization and electrical activity. The discovery of fluorescent dyes for measurements of single-cell exocytosis also complemented patch clamp capacitative measurements in studies on Ca\textsuperscript{2+}-dependent hormone secretion.
channels to the pattern of electrical activity is minimal in cultured cells but may play an important role in intact tissue.

Within the pituitary, there are three additional families of channels contributing to signaling: gap junction channels, receptor channels, and intracellular Ca\textsuperscript{2+} release channels. The role of gap junction coupling is well established in folliculostellate cells, but further work is needed to clarify their relevance in communication among secretory cell types. The potential relevance of these proteins forming hemichannels is also awaiting clarification. At least three types of receptor channels are expressed by pituitary cells: nAChRs, GABA\textsubscript{A}, and P2XRs. In contrast to brain cells, where GABA\textsubscript{A} channels are inhibitory, in pituitary cells they stimulate electrical activity, as do nAChRs and P2XRs. GABA\textsubscript{A} channels are common to all endocrine cells, as are P2XRs, but there is a cell type-specific expression of P2XR subtypes of these channels among pituitary cells. It appears that nAChRs are specific for POMC-producing melanotrophs and corticotrophs. From the electrophysiological point of view, these channels are relatively well characterized. Further work in this field should be focused on physiological conditions under which acetylcholine, GABA, and ATP are released.

All pituitary cells have an additional system to control intracellular calcium, composed of the calcium-conducting channels expressed in the ER membrane. Endocrine pituitary cells express at least 15 subtypes of G\textsubscript{q/11}-coupled
GPCRs (Fig. 16) as well as several receptor tyrosine kinases, whose activation leads to mobilization of intracellular \( \text{Ca}^{2+} \) in an IP3-dependent manner. \( \text{Ca}^{2+} \) mobilization provides an additional security system for these cells to control hormone secretion, which is a calcium-dependent process. The pattern of \( \text{Ca}^{2+} \) release signaling is cell type specific, but not receptor type specific. Gonadotrophs have the most sophisticated \( \text{Ca}^{2+} \) mobilization pathway; they release \( \text{Ca}^{2+} \) in an oscillatory manner in response to activation of any of the \( \text{Ca}^{2+} \)-mobilizing receptors expressed in these cells, as well as in response to injection of IP3, with a frequency of spiking determined by IP3 concentrations. Norepinephrine-stimulated corticotrophs also release \( \text{Ca}^{2+} \) in an oscillatory manner, but at lower frequency. In all other pituitary cell types, \( \text{Ca}^{2+} \)-mobilizing receptors trigger \( \text{Ca}^{2+} \) release in a nonoscillatory manner, raising the question of why the sister cells respond differently to activation of the \( G_{q/11} \) pathway.

From the physiological point of view, electrically driven \( \text{Ca}^{2+} \) signals in somatotrophs, lactotrophs, and melanotrophs resemble the signaling pathway of neuronal cells, requiring high \( \text{Ca}^{2+} \) in extracellular medium and APs as a driving force for \( \text{Ca}^{2+} \) influx and secretion. In these cells, \( \text{Ca}^{2+} \) mobilization is a supplementary pathway to VGCI to up-regulate secretion, and oscillations in \( [\text{Ca}^{2+}] \), are achieved by periodic activation of \( \text{Ca}_i \) channels. The \( \text{Ca}^{2+} \) release pathway provides only a transient source for nonoscillatory elevation in \( [\text{Ca}^{2+}] \), due to the continuous opening of the IP3Rs in the presence of agonist, and \( \text{Ca}^{2+} \) influx through \( \text{Ca}_i \) channels is critical for sustained \( \text{Ca}^{2+} \) signaling. It is unlikely that capacitative \( \text{Ca}^{2+} \) entry is the major driving force for sustained \( \text{Ca}^{2+} \) influx. Further studies are needed to identify channels involved in sustained depolarization in these cells, including TRP channels that could be activated by \( \text{Ca}^{2+} \)-mobilizing receptors. GH cell lines behave similarly, suggesting that from the \( \text{Ca}^{2+} \) signaling point of view they are good cell models.

Gonadotrophs, on the other hand, resemble skeletal muscle cells, relying on \( \text{Ca}^{2+} \) mobilization for a prolonged period and with VGCI controlling the “excitability” of the ER membrane during receptor activation. In these cells, oscillations in \( [\text{Ca}^{2+}] \), are generated by periodic activation of IP3Rs during continuous stimulation of \( \text{Ca}^{2+} \)-mobilizing receptors due to bidirectional actions of cytosolic \( \text{Ca}^{2+} \) on the gating of these channels. Conservation of intracellular \( \text{Ca}^{2+} \) is achieved by its redistribution between ER and mitochondria. In contrast to skeletal muscle cells, there is a “leak” of \( \text{Ca}^{2+} \) from the cells, and VGCI is temporally separated from \( \text{Ca}^{2+} \) mobilization by periodic activation of SK channels. The \( \text{Ca}^{2+} \) signaling properties of gonadotrophs are not preserved in \( \alpha T3-1 \) and \( \beta T3 \) cells, indicating their limited use in the characterization of \( \text{Ca}^{2+} \)-dependent cellular processes. Corticotrophs are likely to have both plasma membrane- and ER-dependent oscillators operative, but further work is needed to clarify the mechanisms controlling these oscillators. Finally, the characterization of \( \text{Ca}^{2+} \) signaling pathways in thyrotrophs still is in a preliminary stage.

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