The Intrinsic Rhythmicity of Spike-Burst Generation in Pancreatic β-Cells and Intercellular Interaction within an Islet

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Pancreatic β-cells display cyclic spike-burst activity in response to extracellular glucose of intermediate concentrations (8–15 mM). The spike-burst duration varies with the change in glucose concentration. Insulin release from β-cells is closely related to an increase in intracellular Ca²⁺ [1], which is mainly due to Ca²⁺ influx during spiking activity [2–4]. Thus, spike generation plays a central role in the regulation of insulin release.

The discovery of ATP-sensitive K⁺ channels (Kₐ₅[ATP]) [5] contributed a great deal to understanding the mechanism underlying the electrical activity in β-cells. β-cells are furnished with a low-affinity glucose transporter, GLUT-2 [6, 7]. The elevation of extracellular glucose concentration causes an increase in oxygen consumption [8, 9]. Although it is well established that glucose induces closure of the Kₐ₅[ATP] channels, which leads to the depolarization underlying spike generation, termination of the spiking activity is not fully elucidated. The activation of Ca²⁺-activated K⁺ channels (Kₐ₅[Ca₅V]) was once postulated to be a decisive factor to terminate sustained spike firing [10, 11]. However, patch-clamp and pharmacological observations ruled out such a possibility on the basis that no Kₐ₅[Ca₅V] channel was activated in the physiological range of Ca²⁺ concentration [12], and charybdoxin, a specific blocker of Kₐ₅[Ca₅V] channels, did not cause any change in electrical activity [13]. Intracellular Ca²⁺ measurements [3, 14, 15] failed to show any direct contribution of intracellular Ca²⁺ in terminating the spiking period.

An alternative model, that slowly inactivating the Ca²⁺ current contributes to the termination of spiking activity, has been proposed from the similarity in burst patterns that is seen in certain molluscan and mammalian neurons [16]. In addition to this view, recently, two views are proposed from different laboratories. One is that the bursting pattern may be mediated by a distinct Ca²⁺-dependent K⁺ channel or a slow Ca²⁺-dependent inactivation process of the Ca²⁺ current [17], and the other is that the depletion of endoplasmic reticulum Ca²⁺ deposits results in the activation of an inward current [18] while filling Ca²⁺ deposits with Ca²⁺ causes termination of the sustained depolarization underlying spiking activity.

In this short article, new experimental data are reviewed and an attempt is made to integrate them into a hypothesis for the regulation of complex electrical activity and insulin release.

General Features of Electrical Activity of the Pancreatic β-Cell and Ca²⁺ Oscillation

The electrical characteristics of pancreatic β-cells have been reviewed [19–21]. In brief, the most important findings for building a comprehensive view will be described first. The resting potential for low glucose (lower than 3 mM; which is a subthreshold level for insulin release in most species of animals) is between −60 and −70 mV (Fig. 1). Upon raising the glucose level from the subthreshold level to 8–15 mM, the membrane potential slowly moves to a threshold potential of about −50 mV. When this threshold is reached, rapid depolarization proceeds to a plateau potential (about −40 mV) from which spikes are elicited (the initial spike-train). After sometime, the spiking...
activity stops, and the membrane potential returns to the partially repolarized level. Then the membrane potential starts to oscillate regularly between the partially hyperpolarized level and the plateau potential on which the spikes are superimposed (Fig. 1). The spike frequency is about 10–12 Hz at the onset of spike-burst, but gradually decreases and sometimes incomplete spikes appear in the final stage of the spike-burst period. The crest of spikes varies between -20 and -15 mV. This value is much lower than the equilibrium potential of Ca\(^{2+}\) (\(E_{Ca} \approx 110\) mV). As the glucose level is elevated, the duration of the spike-burst period (the active phase) becomes longer, while the inter-burst interval (the silent phase) becomes shorter. At higher levels of glucose (20–30 mM), the silent phase vanishes completely and spiking is continuous. Although the duration of the active phase is a function of extracellular glucose concentration [22], spike frequencies in the active phase are not dependent on the glucose level.

Intracellular Ca\(^{2+}\) concentration, [Ca\(^{2+}\)\(_i\)], can be measured by microfluorometry using a fura-2 Ca\(^{2+}\)-sensitive fluorescent probe [23]. When extracellular glucose is increased from the subthreshold level to about 10 mM, intracellular Ca\(^{2+}\) abruptly increases after a small transient decrease, and its elevated level oscillates [3]. Simultaneous recordings of membrane potential and fura-2 fluorescence clearly demonstrated that the phase of Ca\(^{2+}\) oscillation was in accordance with the electrical bursting activity [2]. The removal of extracellular Ca\(^{2+}\) resulted in a fall in the intracellular Ca\(^{2+}\) level [24]. These observations are considered to be strong evidence supporting the view that an increase in Ca\(^{2+}\) influx directly causes elevation at the intracellular Ca\(^{2+}\) level, which is obligatory for the exocytosis of insulin release.

The electrical responses of β-cells to a sudden increase in extracellular glucose consist of an initial spike-train and subsequent cyclic spike-bursts. From the viewpoint of rhythmicity, firstly, the cyclic spike-burst activity will be reviewed. Secondly, we will discuss the initial spike-train in connection with the glucose metabolism on the basis of recent experimental results which may be relevant to the biphasic pattern of insulin secretion response.

**Termination of the Active Phase**

1. **Background inward and outward ionic currents**

In advance of reviewing recent experimental results regarding the termination of depolarization underlying spiking activity, we have to recall the results of experiments carried out about 20 years ago. In those experiments, the electrical membrane resistance was lowest at the beginning of the silent phase, and then gradually increased to a level about twofold of the lowest value [11, 25]. In parallel with the membrane potential jump from the threshold potential to the plateau potential, the membrane resistance decreased drastically, indicating openings of voltage-dependent channels for inward currents. In contrast to a reliable estimation of membrane resistance during the silent phase, there is difficulty in assessing the membrane resistance from the magnitudes of membrane potential displacement induced by passing inward current pulses of constant magnitude during the active phase because voltage-gated channels are rapidly deactivated by the induced hyperpolarization. To overcome this difficulty, Smith et al. [26] conducted perforated-patch whole-cell voltage clamp experiments in cultured single cells. They reported that, at 0 mV glucose, the resting input conductance was 5.1 ± 0.9 nS and that
8 mM glucose decreased the conductance by 80%. Furthermore, they concluded there was no difference in the conductance measured during the plateau when compared to that during the silent phase, showing that an increase in K+ conductance is not likely during termination of the active phase. Unfortunately, because of technical difficulty, the time course of the conductance change in both silent and active phases was not given in their experiments.

The model in which an increase in K+ conductance terminates the plateau potential implicitly assumes the presence of a steady background inward current, while the alternative model, in which inactivation of the background inward current causes a termination of the plateau, implicitly presupposes a steady background outward K+ current.

2. Regenerative nature of repolarization

The spike-burst rhythm can be reset by prematurely applying depolarizing current above the threshold, while supra-threshold hyperpolarizing current applied during the active phase triggers regenerative repolarization [27]. β-cells possess voltage-gated Ca2+ channels. The Ca2+ currents have been characterized in neonatal rat β-cells [28], insulin-secreting cell lines [29], and adult mouse β-cells [30–32]. Rapidly activating Ca2+ currents are inactivated in a Ca2+-dependent manner [29, 33], while slow inactivation (τ ≈ 2.75 s at 0 mV) is Ca2+-independent [31]. Slow inactivation of the Ca2+ currents is proposed to be responsible for sustaining plateau potential [16]. This model seems to be able to describe fully the ionic basis of the plateau potential. However, there are some disagreements between the theoretical spike-burst computed on the basis of slowly inactivated Ca2+ current [34] and actual records. The most prominent discrepancy between them is that the theoretical curve shows progressive elevation of the spike threshold on the plateau potential while actual records show a rather slow fall in the spike threshold after transient elevation during the active phase. Furthermore, if inactivation of the inward current is a time-dependent event, the channel must become inactivated sometime after a long period of depolarization. If the inactivation of Ca2+ channels is the sole factor inducing repolarization, the plateau potential should always be interrupted by repolarization whenever the extracellular glucose concentration is high. However, in reality, at high concentrations of glucose (20–30 mM), the interburst silent phases are completely eclipsed and spiking is continuous. The discrepancy between the actual records and theoretically computed curves implies that factors other than the slow inactivation of Ca2+ channels may play a key role in triggering repolarization of the plateau.

The regenerative nature of repolarization requires the presence of a residual fraction of background inward current channels which will be deactivated in the process of repolarization. The magnitude of the background inward current can be assessed from the slope of the membrane potential change (dV/dt). It should be noted that the maximum slope of regenerative repolarization is usually less in absolute magnitude than the slope of the regenerative depolarization that triggers spike-burst, indicating the ionic membrane current at the phase of regenerative repolarization is less than the current at the phase of regenerative depolarization. When the plateau potential is attained, the background inward current is almost equal to the background outward K+ current. During the process of regenerative repolarization, the voltage-gated Ca2+ channel current may become zero (i.e., the maximum ionic membrane current may correspond to the background Ca2+ current which has been inactivated) unless the K+ current increases. The fact that the ionic membrane current during regenerative repolarization is smaller in amplitude than that during regenerative depolarization raises the difficulty of detecting the background Ca2+ current using the conventional whole-cell clamp technique.

3. What is the trigger of the regenerative repolarization?

To induce regenerative repolarization, the triggering of repolarization is necessary. The triggering of repolarization can be caused by either a decrease in Ca2+ conductance or an increase in K+ conductance. Sulfonylurea (tolbutamide, glybenclamide, or glyburide) specifically inhibits the opening of KATP channels [35–37] and causes continuous spiking activity at intermediate concentrations of glucose. There is substantial evidence that the sulfonylurea receptor could be an integral component of the ATP-sensitive K+ channel in β-cells [38–40]. These observations suggest that opening of the KATP channels is indispensable for triggering regenerative repolarization.

Raising extracellular Ca2+ to 10 mM shortens the duration of the spike-burst in mouse β-cells perfused with a medium containing 15 mM glucose, and restores rhythmic spike-burst activity when continuous spike activity was induced by 30 mM glucose. On the basis of these observations, Henquin [22] has suggested that the metabolic control of the influence of Ca2+ on KATP channels could be a central factor in glucose regulation of electrical activity. A clue to solve the problem of how the triggering of repolariza-
tion occurs was found in the effect of removal of extracellular Ca\(^{2+}\) on the pattern of electrical response. At 11.1 mM of glucose, nominally Ca\(^{2+}\)-free solution induced a change in the response pattern from the typical spike-burst type to the continuous spiking type (Fig. 2). Similar pattern change has been reported by Gilon and Henquin [41] who lowered the extracellular Ca\(^{2+}\) concentration from the normal value (2.5 mM) to 1 mM at 15 mM of glucose. It is known that there are Na\(^{+}\)/Ca\(^{2+}\)-countertransporters in pancreatic β-cells [42, 43]. Thus, it may be reasonable to consider that a decrease in [Ca\(^{2+}\)]\(_i\) results in a decrease in Na\(^{+}\) influx, leading to a fall in cytosolic Na\(^{+}\) concentration, [Na\(^{+}\)]\(_c\), just under the plasma membrane, if Na\(^{+}\) ions are continuously extruded by the Na\(^{+}\)-pump. The decrease in cytosolic Na\(^{+}\) results in the slowing of ATP consumption at Na\(^{+}\)/K\(^{+}\)-pump sites. In addition to this finding, lowering the Na\(^{+}\) concentration in the Ca\(^{2+}\)-free solution to 25 mM immediately suppressed spike generation and induced slowly progressing depolarization after transient hyperpolarization (lowermost trace in Fig. 2). This finding rules out the possibility that the depolarization observed when removing extracellular Ca\(^{2+}\) is due to an increase in Na\(^{+}\) influx. Under the circumstances where the ATP production rate is kept unchanged, the slowing of ATP consumption should prevent the ATP level from falling, leading to persistent depolarization and continuous spiking [24]. The removal of extracellular Na\(^{+}\) from the perfusion solution containing Ca\(^{2+}\) also induced a change in the pattern of electrical activity similar to that caused by the removal of extracellular Ca\(^{2+}\). The substitution of Na\(^{+}\) with either Tris, Li\(^{+}\) or choline caused persistent depolarization on which spikes were superimposed [24, 44]. The re-addition of Na\(^{+}\) induced a marked and long-lasting silent hyperpolarization that was inhibited by ouabain [44]. Simply adding ouabain to the perfusion solution also caused shortening of the silent period or the complete elimination of the silent phase in a manner depending on its concentration [11]. Furthermore, omitting K\(^{+}\) from the perfusion solution had similar effect on the pattern of electrical response as ouabain [45, 46]. These depolarizing effects have been previously explained as the direct consequence of nonoperation of the electrogenic Na\(^{+}\)/K\(^{+}\)-pump. However, the depolarization induced by various means of Na\(^{+}\)/K\(^{+}\)-pump inhibition was always accompanied by the suppression of K\(_{\text{ATP}}\) channel activity [47]. In light of these observations, it may be more rational to deduce that the conditions which cause a fall in ATP consumption always prevent the opening of K\(_{\text{ATP}}\) channels. In other words, the observed continuation of depolarization is a result of the decrease in ATP consumption. Under physiological conditions, a gradual increase in ATP consumption causes a fall in the ATP/ADT ratio, leading to slight repolarization that is brought about by opening a small fraction of the K\(_{\text{ATP}}\) channels. It may be said that the slight repolarization resulting from a fall in the ATP/ADT ratio triggers regenerative repolarization through deactivating the voltage-dependent Ca\(^{2+}\) channels. In support of this view, the electrical membrane resistance is lowest at the beginning of the silent phase [11, 25].

4. Oscillation in the ATP/ADT ratio

During the active phase, intracellular Ca\(^{2+}\) concentration was elevated to about 1×10\(^{-6}\) M. The amplitude of the spikes elicited on the plateau potential was merely about 20 mV, and the capacity of a single β-cell was about 4.5 pf (d=12 μm; 1 μF/cm\(^2\)). The minimum charges carried by the influxed Ca\(^{2+}\) ions were calculated to be 90×10\(^{-12}\) C/spike (=4.5×10\(^{-19}\) mol/spike). This amount of Ca\(^{2+}\) caused a concentration change of 4.97×10\(^{-7}\) M in a cell with a diameter
of 12 μm. On the other hand, taking into account the fact that the peak potential of spikes is about −20 mV, whereas the equilibrium potential of Ca²⁺ (E_{Ca}) is about 110 mV, it may be considered that a significant fraction of electric charge carried by the Ca²⁺ influx is cancelled by outgoing K⁺ current, suggesting that a larger amount of Ca²⁺ ions than calculated above flows into a cell during spike generation. However, microfluorometric experiments show that the elevation in [Ca²⁺]i, induced by glucose stimulation is about 1×10⁻⁶ M at the highest. This calculation implies that a substantial amount of Ca²⁺ ions are extruded from the cell or taken up by intracellular Ca²⁺ buffering organelles at a considerably high rate. In nerve fibers, the most potent machinery functioning to extrude Ca²⁺ is Na⁺/Ca²⁺ exchange [48]. The activation of Na⁺/Ca²⁺ exchange should cause the elevation of intracellular Na⁺. Although the change in Na⁺ concentration in the compartment just under the plasma membrane is believed to be much higher than the change in bulk cytosol, it is difficult to detect the change in the emission strength of the Na⁺-sensitive fluorescent probe, SBFI, if the Na⁺/K⁺-pump is working. Indeed, Gilon and Henquin [49] reported that ouabain caused a detectable rise in [Na⁺], in microfluorometric experiments with SBFI.

Activation of the Na⁺/K⁺-pump is accompanied by the acceleration of ATP consumption. Very recently, oscillation of the ATP/ADP ratio was elegantly demonstrated by Nilsson et al. [50]. They sampled aliquots of islet cells preloaded with fura-2 before and after adding glucose to the cell suspension at 30 s intervals while continuously monitoring the cell [Ca²⁺]i, in the suspension fluorometrically. Both ATP and ADP were assayed by bioluminescent methods [51]. Using this process, it became possible to trace the change in the ATP/ADP ratio in connection with the change in [Ca²⁺]i, which reflected the electrical activity during the period of incubation. The experimental results clearly showed that the initial rise in ATP/ADP ratio preceded the initial rise in [Ca²⁺], and that at a high concentration of Ca²⁺ (an additional 5 mM Ca²⁺), oscillation in the ATP/ADP ratio was in accordance with [Ca²⁺]; oscillation; indicating that a rise in the ATP/ADP ratio causes spiking activity and a fall in the ratio terminates spike generation.

Patch-clamp experiments in the cell-attached configuration confirmed that both removing extracellular Ca²⁺ and lowering extracellular Na⁺ induced suppression of the K(ATP) channel activity at 8.4 mM of glucose [47]. This observation also suggests elevation of the ATP level. Larsson et al. [52] re-examined membrane conductance using the same method used by Smith et al. [26], and unquestionably demonstrated that membrane conductance was much larger at the beginning of the silent phase as compared to that at the midpoint of the active phase. During the silent phase, intracellular Ca²⁺ concentration gradually falls, indicating that almost all of the voltage-dependent Ca²⁺ channels are closed in this period. Electrogenic Na⁺/Ca²⁺ countertransport may contribute to the background inward current to some extent. A leak from the intracellular Ca²⁺ deposits may maintain the resting Ca²⁺ level. It is not likely that the non-selective cation channel contributes substantially to the background current because activation of this channel requires Ca²⁺ concentrations higher than 10⁻⁴ M [53].

The spike-burst activity observations cited above lead to the following view on periodical spike-burst activity: Elevation of the intracellular Ca²⁺ level due to spiking activity causes an increase in the Na⁺ influx through activating Na⁺/Ca²⁺-countertransport. The result elevation of intracellular Na⁺ accelerates ATP consumption through activation of the Na⁺/K⁺-pump, which causes a fall in the ATP/ADP ratio. Slight repolarization resulting from the opening of a small fraction of K(ATP) channels triggers regenerative repolarization, which terminates spiking activity. In the course of the progress of repolarization, a residual fraction of voltage-gated Ca²⁺ channels are almost completely deactivitated. During the silent phase, the intracellular Ca²⁺ concentration gradually declines, leading to a fall in [Na⁺], which causes a gradual rise in the ATP/ADP ratio through decrease in ATP consumption. The gradual rise in the ATP/ADP ratio leads to gradual depolarization back to the threshold of low-threshold Ca²⁺ channels, of which the opening causes regenerative depolarization underlying the next spike-burst. The K(ATP) channel has a twofold role: during the silent phase, its gradual closing causes gradual depolarization to the threshold for the plateau potential underlying spiking activity; and during the active phase, its gradual opening triggers regenerative repolarization. The functional coupling of the Na⁺/Ca²⁺-countertransporter and Na⁺/K⁺-pump induces oscillation of the ATP/ADP ratio under conditions where the ATP production rate remains unchanged and voltage-gated Ca²⁺ channels are functioning. A part of these conditions is fulfilled by keeping the glucose supply constant.

The Initial Spike-Train

1. Response of β-cells to glucagon

Single cells in an intact islet always display the initial spike-train and the subsequent spike-burst activity in response to a sudden increase in extracellular glu-
cose [54], while an isolated single β-cell does not have the initial spike-train [26]. The mechanism of the initial spike-train preceding cyclic spike-burst activity has not attracted much attention from researchers up to now.

It is known that the amount of glycogen, the main deposits of carbohydrates, in islet cells depends on the extracellular glucose concentration, and that dibutyl-cAMP reduces the glycogen content, whereas adrenaline increases the level of islet glycogen in the presence of 16.7 mM glucose [55]. Single pancreatic β-cells purified by autofluorescence-activated cell-sorting have been reported to release insulin at a rate five times less during nutrient stimulation as compared to intact islets [56]. cAMP levels are low in purified β-cells, and glucagon induces a rise in the cAMP level [57]. The nutrient-induced insulin release from purified β-cells is noticeably increased by the addition of either dibutyl-cAMP, glucagon or pancreatic α-cells [56]. These findings suggest that the glucagon released from α-cells affects the secretory activity of β-cells lodged in the same islet through elevating the cAMP content of the β-cells.

Glucagon enhances the glucose-induced electrical response of the β-cells in intact islets. Upon adding glucagon, the plateau duration increases and the silent phase becomes shorter [58, 59]. The change in electrical activity may be caused by either activation of the voltage-gated Ca²⁺ channel or stimulation of the glucose supply. It has been reported that a slowing of the decay and an increase in the peak Ca²⁺ channel current response to step depolarization is induced by cAMP in both cardiac myocyte and β-cells [60, 61], probably through A-kinase-dependent phosphorylation. On the other hand, the effect of glucagon on the glucose metabolism is well defined in liver cells, stimulating the phosphorylase of glycogen to yield glucose-1-phosphate [62]. The phosphorylase kinase which activates glycogenphosphorylase is the first enzyme to be established as a cAMP-dependent protein kinase [63].

The fluorescence measurement of endogenous reduced pyridine nucleotides (NAD(P)H) in isolated islet cells is a good measure of glucose metabolism. Raising the concentration of glucose from 3 to 15 mM causes an approximate 50% increase in the autofluorescence, and the mannose 1-phosphate that inhibits glucose phosphorylation [8] causes a marked decrease in the fluorescence [41]. A similar change in NAD(P)H induced by glucose has been reported in single isolated β-cells [64]. Ca²⁺-sensitive mitochondrial dehydrogenase is known to stimulate NAD(P)H formation [65]. However, it has been confirmed that NAD(P)H fluorescence increases before Ca²⁺ entry in response to glucose [66], indicating that an increase in glucose itself is more closely relevant to the increase in NAD(P)H than Ca²⁺.

At 2.8 mM of glucose, 8.6 × 10⁻⁸ M glucagon scarcely affected either NAD(P)H or electrical activity (unpublished observation). In contrast, adding glucagon to the perfusion solution containing 11.1 mM glucose caused a 30% increase in NAD(P)H in the presence of 2.5 mM Ca²⁺ (Fig. 3), whereas in the absence of extracellular Ca²⁺, the addition of glucagon had little effect on NAD(P)H fluorescence [59]. Furthermore, when being stimulated by glucagon containing 11.1 mM glucose, the removal of extracellular Ca²⁺ caused NAD(P)H fluorescence decrease gradually until it reached the level attained before glucagon had been added. The resupplementation of Ca²⁺ restored NAD(P)H fluorescence. All the calmodulin antagonists tested (chlorpromazine, W-7, and trifluoperazine) similarly suppressed the glucagon-stimulated increase in NAD(P)H fluorescence [59]. These observations suggest that cAMP, together with Ca²⁺/calmodulin, stimulates NAD(P)H formation even in β-cells, probably through accelerating the glucose supply from glycogen.

2. Effect of cAMP on exocytosis

Studies on permeated β-cells indicate that cAMP increases insulin release even when [Ca²⁺]ₗ is held constant [67]. In connection with this, Åmmalä et al. [61] reported that, in whole-cell perforated patch-clamp experiments, cAMP caused an increase in Ca²⁺ current (about 30%) to step depolarization with a very large capacitance change (an increase of 175%), which means a 175% increase in the membrane fusion of secretory granules to the plasma membrane. Taking into account the fact that the intracellular trafficking of secretory granules is an ATP-dependent event [68, 69], the very large increase in capacitance change to step depolarization (representing insulin release) induced by cAMP is believed to reflect elevation of the ATP level. In contrast to the effect of cAMP, adrenaline is known to suppress insulin release and electrical activity [70] through stimulation of α-adrenoceptors, and also induce the elevation of islet glycogen content in the presence of 16.7 mM glucose [55], suggesting that the stimulation of α-adrenoceptors suppresses glucose utilization by activating glycogen synthesis. In fact, clonidine, an α₂-adrenoceptor agonist, induces a fall in the ATP/ADP ratio [50]. These findings also support the view that regulation of both the synthesis and breakdown of glycogen is closely related to the control of electrical activity by affecting
the ATP production rate. It may be too premature to derive any concrete view on the mechanism of the initial spike-train from the data reviewed so far, however, observations in glucagon experiments provide important information for drawing a working model of the initial spike-train.

3. Biphasic response of insulin release

It is well known that the phase of the initial spike-train is in accordance with the first phase of insulin-release response, and the subsequent period of rhythmical spike-bursts is in accordance with the second phase of secretion response. At present, there is no generally accepted view on the mechanism of this biphasic phenomenon. The integration of data reviewed above allows us to speculate the mechanism as follows.

At low concentrations of glucose, α-cells are secreting glucagon, which induces a rise in the cAMP level in β-cells lodged in the same islet but causes little change in the membrane potential. This is in accordance with the observation that, in a state of low blood glucose, an increased glucagon release from α-cells does not induce elevation of the blood insulin level, and hence does not lead to hypoglycemia but rather maintains the glucose level within the physiological range. When extracellular glucose is raised, the membrane depolarizes to a threshold for the plateau potential on which Ca^{2+}-spikes are superimposed, resulting in the rapid elevation of [Ca^{2+}]. The increased [Ca^{2+}]; together with cAMP, which has already been increased, acts to activate enzymes relevant to glycogenolysis in the β-cells. The stimulation of glycogenolysis causes a rise in ATP production. As long as the glucose supply from stored glycogen remains elevated, depolarization continues, notwithstanding that spikes are continuously elicited, and the level of [Ca^{2+}]; stays so high that acceleration of ATP consumption at the Na^{+}/K^{+}-pump sites results. On switching to the high-glucose medium, however, the glucagon release from α-cells is soon suppressed and, consequently, the cAMP level in the β-cells falls. The fall in cAMP level is followed by deactivation of the enzymes responsible for the breakdown of glycogen. Under the conditions where ATP consumption is accelerated, a decrease in the glucose supply from stored
glycogen leads to a fall in the ATP/ADP ratio, which triggers regenerative repolarization. After cessation of the glucose supply from intracellular deposits, glucose influx from the extracellular solution becomes the sole fuel source for ATP synthesis. This period may correspond to the second phase of insulin-secretion response.

4. Effect of acetylcholine

Unlike the response to glucagon, β-cells respond to acetylcholine even at low glucose concentrations [71]. The pattern of response is very similar to that observed when glucose is suddenly increased, consisting of the initial spike-train and subsequent spike-burst [49, 71]. Muscarinic stimulation increases Na$^+$ influx [49]. Thus, acetylcholine-induced depolarization is believed to be due to an increase in the background inward Na$^+$ current. An increase in the background inward current means that the electrical response can be elicited at even low concentrations of glucose. When acetylcholine is present in extracellular medium, the ATP/ADP ratio may gradually fall and oscillate in a range relatively lower than that in glucose-stimulated cells. This point remains to be verified.

Summary

The pancreatic β-cell has four types of Ca$^{2+}$ channel (L-type, T-type, low-threshold slowly inactivating, and low-threshold non-inactivating Ca$^{2+}$), although the low-threshold non-inactivating Ca$^{2+}$ channel has not yet been confirmed experimentally. Beside these, there are at least three types of K$^+$ channels ($K_{ATP}$, $K_{(Ca,V)}$, and $K_{(V)}$), and transporters (GLUT-2, Na$^+$/Ca$^{2+}$-countertransporter, and Na$^+$/K$^+$-pump) as schematically shown in Fig. 4.

Opinions on the mechanism of spike-burst are converging to the following view: At intermediate glucose concentrations, the intracellular ATP/ADP ratio oscillates in the following way. A gradual rise in the ATP/ADP ratio causes gradual progression of depolarization to the threshold for the low-threshold Ca$^{2+}$ channels, of which the opening causes regenerative depolarization to the plateau potential on which spikes (the L-type Ca$^{2+}$ channel contributes to spike firing) are superimposed. During the active phase, a fall in

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**Fig. 4.** Schematic drawing of receptors, channels, transporters, and the enzyme systems related to the glucose metabolism. Details are in the text. The T-type Ca$^{2+}$ channel is not shown.
the ATP/ADP ratio follows a gradual rise in ATP consumption. Slight repolarization due to the opening of a small fraction of K_{ATP} channels triggers regenerative repolarization. With the progress of repolarization, a residual fraction of voltage-gated Ca^{2+} channels (low-threshold non-inactivating) are deactivated. During the silent phase, a gradual rise in the ATP/ADP ratio leads to gradual depolarization back to the threshold for the next spike-burst.

There are still a diversity of views regarding the mechanism of the initial spike-train. On the basis of observations made in various laboratories including ours, we propose the following working model: At low concentrations of glucose, α-cells secrete glucagon which induces a rise in cAMP in β-cells lodged in the same islet. A rise in cAMP itself does not activate the enzymes relevant to glycogenolysis, but merely prepares to activate the enzymes. When extracellular glucose increases, Ca^{2+} spikes are elicited. Influened Ca^{2+} ions, together with cAMP, work to activate the enzymes, resulting in an additional supply of fuel for ATP synthesis. After sometime, the cAMP level falls back to a low level and the additional glucose supply from stored glycogen stops. This reaction sequence may be the mechanism behind the initial spike-train. To substantiate this working model, it may be important to elucidate the dependence of the phosphorylase kinase and glycogenphosphorylase activities on the Ca^{2+} in β-cells.

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