Role of the active zone protein, ELKS, in insulin secretion from pancreatic β-cells

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ABSTRACT

Background: Insulin is stored within large dense-core granules in pancreatic β-cells and is released by Ca 2+ -triggered exocytosis with increasing blood glucose levels. Polarized and targeted secretion of insulin from β-cells in pancreatic islets into the vasculature has been proposed; however, the mechanisms related to cellular and molecular localization remain largely unknown. Within nerve terminals, the Ca 2+ -dependent release of a polarized transmitter is limited to the active zone, a highly specialized area of the presynaptic membrane. Several active zone-specific proteins have been characterized; among them, the CAST/ELKS protein family members have the ability to form large protein complexes with other active zone proteins to control the structure and function of the active zone for tight regulation of neurotransmitter release. Notably, ELKS but not CAST is also expressed in β-cells, implying that ELKS may be involved in polarized insulin secretion from β-cells.

Scope of review: This review provides an overview of the current findings regarding the role(s) of ELKS and other active zone proteins in β-cells and focuses on the molecular mechanism underlying ELKS regulation within polarized insulin secretion from islets.

Major conclusions: ELKS localizes at the vascular-facing plasma membrane of β-cells in mouse pancreatic islets. ELKS forms a potent insulin secretion complex with L-type voltage-dependent Ca 2+ channels on the vascular-facing plasma membrane of β-cells, enabling polarized Ca 2+ influx and first-phase insulin secretion from islets. This model provides novel insights into the functional polarity observed during insulin secretion from β-cells within islets at the molecular level. This active zone-like region formed by ELKS at the vascular side of the plasma membrane is essential for coordinating physiological insulin secretion and may be disrupted in diabetes.

Keywords  Pancreatic β-cells; ELKS; Active zone protein; Voltage-dependent Ca 2+ channel; Insulin exocytosis; Ca 2+ influx

1. INTRODUCTION

The hormone insulin is stored in large dense-core granules within pancreatic β-cells and is released by exocytosis in response to elevated extracellular glucose concentrations. Triggered insulin exocytosis appears to involve increases in intracellular ATP concentrations, closure of ATP-sensitive K+ channels, membrane depolarization, and Ca 2+ influx through voltage-dependent Ca 2+ channels (VDCCs) [1]. However, exocytotic “hot spots” and polarized insulin granule exocytosis in β-cells remain elusive. Regarding rodent islets, β-cells are generally structured in rosettes of cells surrounding the central pancreatic venous vasculature [2–4]. There is increasing evidence suggesting that the β-cell plasma membrane encompasses functionally separated compartments, indicating β-cell polarity [5]. In support of this, E-cadherin and GLUT2 have been shown to be located on the lateral regions of the plasma membrane [6,7]. In the vascular apogee (i.e., the side of the plasma membrane opposite the vasculature), PAR-3 and ZO-1 tend to be positioned where adjacent β-cells project their primary cilia [7].

Interestingly, electron microscopy indicates that insulin granules are enriched along the vascular-facing plasma membrane following cellular stimulation [2], supporting the concept of polarized secretion, i.e., insulin is released from the vascular-facing plasma membrane toward the bloodstream in a Ca 2+ -dependent manner. The location of insulin exocytosis is under debate since a two-photon imaging study showed that most events are occurring at the “avascular” compartment of β-cells within intact islets (i.e., cell–cell interfaces away from blood vessels) [8]. However, when two-photon imaging with 3D-measurements were recently employed, it was clear that insulin exocytosis mostly takes place at the vascular-facing plasma membrane of β-cells [9]. Despite these findings, there is still a lack of information on the molecular mechanism of the polarity of the Ca 2+ influx and insulin secretion from pancreatic islets.

In neurons, the release of a transmitter, which has a specific polarity, takes place at specialized active zones on the presynaptic membrane. Active zones are defined as slightly electron-dense regions beneath the presynaptic plasma membrane [10]. The cytomatrix at the active zone contains the molecular machinery necessary for docking and fusion of...
SYNAPTIC VESICLES [11]. THIS UNIQUE AND VITAL STRUCTURE COVERS active-
zone-specific protein families, including RIM1, Munc13-1, Piccolo (i.e.,
Aczonin), Bassoon, CAST (i.e., ERC2), and ELKS (i.e., ERC1) [12,13].
Additionally, VDCCs, cytoskeletal proteins, and SNAREs are associated
with the cytomatrix at the active zone [11]. In particular, the CAST/
ELKS protein family has been reported to directly bind to Bassoon,
Piccolo, and RIM1 and indirectly to Munc13-1 through RIM1, sug-
gest that CAST/ELKS family proteins may act as a platform for a
dynamic multicomplex at the active zone [14,15]. Moreover, growing
evidence suggests the structure and function of the active zones are
controlled by CAST and ELKS to tightly regulate polarized neuro-
transmitter release [15,16]. Interestingly, ELKS but not CAST is also
expressed in pancreatic β-cells and localizes at the vascular-facing
plasma membrane of β-cells within islets [17]. We further found that
ELKS is one of the key players regulating polarized Ca2⁺ influx and
insulin secretion from islet β-cells [18].

In this review, we provide a brief summary of the recent findings
related to the molecular structures and biochemical properties of the
CAST/ELKS protein family and discuss the potential role(s) of ELKS in
polarized insulin secretion from β-cells.

2. MOLECULAR STRUCTURES AND BIOCHEMICAL PROPERTIES
OF THE CAST/ELKS PROTEIN FAMILY IN THE PRESYNAPTIC
ACTIVE ZONE

CAST was first purified from the rat brain [19] and was independently
identified by the yeast two-hybrid system as ERC2 [20]. CAST has
coiled-coil regions throughout its entire protein structure and contains
a unique three amino acid sequence (IWA) at its C-terminus
(Figure 1A). CAST has been shown to directly interact with the PDZ
domain of RIM1 via this IWA motif [19,20]. Other active zone proteins,
such as Bassoon, Piccolo, and Munc13-1, bind directly to the middle
region of CAST [14,21] (Figure 1A). Additionally, CAST indirectly binds
to Munc13-1 through RIM1 [14]. ELKS is a member of the CAST protein
family, and it has been reported that its encoding gene is translocated
to a different chromosome in thyroid carcinoma [22]. ELKS is named
after its high content of the amino acids E, L, K, and S [22]. In 2002,
three groups independently identified ELKS within different systems
and renamed it CAST2, Rab6IP2, and ERC1 [19,20,23,24]. However,
as this protein was first identified as ELKS [22], we used this termi-
nology within this review to avoid confusion. CAST and ELKS show
relatively high homology (~70% amino acid identity), their molecular
weight is ~120 kDa, and they appear to form oligomers with each other
[25]. As with CAST, ELKS has been reported to directly bind to
Bassoon, Piccolo, and RIM1 and indirectly to Munc13-1 through RIM1
[25]. In addition, the brain-specific variant of Munc13-2, bMunc13-2,
interacts with the C-terminal region of ELKS but not with CAST [26]
(Figure 1A). The physiological significance of the CAST/ELKS-mediated active zone
protein interactions remains unknown, but it is well recognized that
these CAST/ELKS-dependent protein–protein interactions among
active zone proteins may be the molecular basis for the integrity of
presynaptic active zones (Figure 1B). Additionally, disruption of CAST/
RIM1 binding and/or CAST-Bassoon binding have been shown to
significantly impair synaptic transmission in cultured neurons [14].
In addition to these active zone proteins, CAST/ELKS share many synaptic
proteins as binding partners, such as Liprin-α, Rab6, and VDCCs
[23,27,28]. VDCCs such as the N-, P/Q-, R-, and L-types are essential
for neurotransmitter release at presynaptic active zones [29–32].
VDCCs are heteromultimeric protein complexes composed of the pore-
forming α-subunit and the auxiliary γ2, δ-, β-, and γ-subunits
[33,34]. In particular, there is evidence of physical and functional in-
teractions between CAST/ELKS family members and VDCCs. CAST and
ELKS directly interact with the VDCC-β subunit (β4) [28,35], and in a
BHK cell co-expression system CAST slightly shifts the I-V relationship
to the hyperpolarizing direction by about 5 mV without affecting current
density [28]. Consistent with these observations, Kaeber et al. recently
reported that ELKS enhances presynaptic Ca2⁺ influx to boost the
release probability at inhibitory hippocampal nerve terminals with a
Ca2⁺-imaging method [36]. More recently, electrophysiological anal-
yses of the CAST/ELKS and VDCC relationship have been performed
in vivo. For example, the presynaptic Ca2⁺ influx was strongly reduced
within rod photoreceptors of CAST-knockout (KO) and CAST/ELKS-
double-KO mice as analyzed by whole-cell voltage-clamp recordings
[37]. Moreover, in the calyx of Held synapses in CAST/ELKS-double-KO
mice, both VDCC currents and numbers were significantly decreased
as was the readily releasable pool size. Intriguingly, loss of CAST/ELKS
increased the probability of synaptic vesicle release. It has been
suggested that CAST/ELKS complexes regulate the probability of
synaptic vesicle release through post-priming, a late step controlling
synaptic vesicle release. Several other reviews and articles describe
the CAST/ELKS family with additional detail [13,15,16].

Therefore, CAST and ELKS tightly regulate the release of polarized
neurotransmitters by controlling the structure and function of the active
zones; however, other than the brain, the mechanisms by which active
zone proteins contribute to cellular functions—such as hormone
secretion from endocrine cells—remain poorly understood.
3. ELKS LOCALIZES AT THE VASCULAR-FACING PLASMA MEMBRANE OF \(\beta\)-CELLS

\(\beta\)-cells appear to lack active zones, which are ultrastructurally characterized as electron-dense regions of cytoskeletal filaments underneath the plasma membrane. However, active zone proteins, such as ELKS, RIM2, Munc13-1, Piccolo, and Bassoon, and synaptic scaffold proteins, such as Liprin-\(\alpha\), were also found in pancreatic \(\beta\)-cells [9,17,38–41]. Some of these proteins may be involved in the regulation of insulin secretion. Treatment of islets with antisense oligodeoxynucleotides against Piccolo reduced insulin secretion evoked by glucose and a cAMP analog [38]. RIM2 has been demonstrated to regulate the docking and priming of insulin granule exocytosis in a study using RIM2-KO mice [41]. Munc13-1 acts as a priming factor in insulin exocytosis [40]. Thus, identifying the roles of active zone proteins in pancreatic \(\beta\)-cells remains a focus within insulin exocytosis research; however, the functional relationship of these proteins with polarized insulin secretion has not been elucidated.

Because CAST and ELKS have been implicated in the \(\text{Ca}^{2+}\)-dependent exocytosis of neurotransmitters as described above, we hypothesized that CAST and ELKS are potential candidates for localization at insulin exocytotic “hot spots” in \(\beta\)-cells [17]. Islets in rats express ELKS but not CAST proteins, whereas the brain expresses both CAST and ELKS proteins (Figure 2A). In rodents, islets express two major ELKS splice variants: ELKS\(\alpha\) (brain isoform, \(\sim 120\) kDa) and ELKS\(\varepsilon\) (ubiquitous isoform, \(\sim 140\) kDa) [19]. These variants have a distinct C-terminus. ELKS\(\alpha\) has the IWA amino acid motif (Figure 1A).

Double staining for insulin and ELKS showed immunoreactivity of ELKS in insulin-positive \(\beta\)-cells in pancreas sections, indicating that ELKS is most abundant in \(\beta\)-cells (Figure 2B). Higher magnification confocal imaging of islets showed that ELKS was localized at the plasmalemmal region of \(\beta\)-cells, especially those facing blood capillaries labeled with VE-cadherin, a marker of endothelial cells (Figure 2C). However, this pattern differed from the immunostaining pattern of the exocytotic SNARE protein, Syntaxin 1, as it was seen on the entire plasma membrane [42]. In neurons, the t-SNAREs,
Syntaxin 1, and SNAP-25, are similarly present on the entire axonal plasma membrane, and are not specifically localized to terminal nerve active zones [43]. Furthermore, immunogold electron microscopy confirmed that ELKS (labeled with small gold particles) localized to the plasma membrane facing the vasculature and was frequently detected in close proximity to insulin (large gold particles)-containing granules docked on the plasma membrane (Figure 2D). Thus, ELKS localizes to the docking sites of insulin granules at the β-cell plasma membrane, and in particular, accumulates near the vasculature in islets, implying that it plays a role in insulin granule exocytosis.

4. ELKS DEFINES THE FUSION SITE OF INSULIN GRANULES

In single β-cells cultured from dispersed islets, such localization of ELKS was not observed at the plasma membrane. ELKS immunostaining was primarily observed throughout the plasma membrane in single β-cells and MIN6 β-cells, which predominantly express ELKS but not CAST [17]. We further examined ELKS distribution in the plasma membrane of MIN6 cells using total internal reflection fluorescence microscopy (TIRFM) imaging, of which the evanescent field illumination reaches a <100-nm-thick layer immediately adjacent to the cover glass and illuminates only the plasma membrane under our TIRF conditions [44,45]. We found that ELKS immunofluorescence was unevenly but locally distributed at separate clusters within the plasma membrane. Interestingly, double immunostaining for ELKS and insulin suggested that the formar may interact with insulin granule docking sites. However, we previously showed that Syntaxin 1 clusters are essential for the docking and fusion of insulin granules [46,47]. We further explored the interactions between insulin granule docking sites, ELKS, and Syntaxin 1 clusters by triple immunostaining analyzed using TIRFM. Most insulin granules colocalized with either ELKS or Syntaxin 1 clusters, and ~50% of granules docked with ELKS clusters colocalized with Syntaxin 1. In summary, insulin granules preferentially dock with ELKS clusters colocalized with Syntaxin 1 clusters. These results suggest that ELKS may be involved in physiological interactions with insulin granules at docking sites. Does insulin exocytosis occur at the sites of ELKS clusters? Using live MIN6 cells and TIRF images, we analyzed the interaction(s) between docking and fusion of GFP-tagged insulin granules and ELKS clusters labeled by a TAT-conjugated, Cy3-labeled ELKS antibody. Stimulation of MIN6 cells with high KCl showed that the fusion of insulin granules frequently occurred at ELKS clusters (Figure 2E,F), with ~60% of all fusion events occurring at these sites. Thus, fusion of insulin granules occurred selectively at ELKS cluster sites in the plasma membrane of MIN6 cells. The link between ELKS and insulin granule fusion sites was also observed in insulin-secreting INS-1 cells [48]. Of note, attenuation of ELKS expression by RNA interference decreased the number of ELKS clusters within the plasma membrane, thereby suppressing glucose-evoked insulin secretion from MIN6 cells [17]. These results indicate that ELKS may regulate insulin granule exocytosis and is a possible candidate for defining the fusion site of insulin exocytosis.

Taken together with the fact that ELKS localizes at the vascular-facing plasma membrane within islets, these data strongly suggest that ELKS forms an active zone-like region at these locations on the β-cell plasma membrane. To confirm this, we performed experiments using β-cell-specific ELKS-KO (ELKS−/−) mice and in situ Ca2+ imaging.

5. ELKS POTENTIATES OPENING OF L-TYPE Ca2+ CHANNELS AND FIRST-PHASE INSULIN SECRETION

We generated ELKS−/− mice by crossing ELKS floxed (ELKS<sup>lox/lox</sup>) mice [49] with RIP-Cre mice expressing Cre recombinase in pancreatic β-cells [50] in order to examine the potential role(s) of ELKS in pancreatic β-cells. In islets isolated from ELKS−/− mice, the expression levels of ELKS isoforms were markedly decreased, but no changes were detected in terms of the expression of glucokinase, GLUT2, SNARE proteins required for insulin exocytosis, and active zone proteins (Bassoon, RIM2, and Munc13-1). We confirmed reduction of ELKS expression in insulin-positive β-cells in ELKS−/− mice islets via immunostaining of pancreatic sections (Figure 3A).

The oral glucose tolerance test showed impaired glucose tolerance in ELKS−/− mice. Within ELKS−/− islets, insulin secretion evoked by glucose was substantially reduced compared with control islets. Insulin secretion measurements from pancreatic β-cells demonstrated that glucose induces insulin secretion in a characteristic biphasic pattern, with a transient first-phase followed by the second phase sustained component [51–53]. In the perfused pancreata of ELKS−/− mice, there was selective reduction of the first-phase insulin secretion (Figure 3B). Moreover, insulin secretion induced by high K+ depolarization, which evokes first-phase secretion [54], was also decreased in ELKS−/− mice. Our results suggest that ELKS plays a role in glucose-stimulated first-phase insulin secretion. On the other hand, TIRF or electron microscopy did not reveal any change in the number of morphologically docked insulin granules between control and ELKS-KO β-cells. Thus, insulin granule docking was not impaired in ELKS-KO β-cells.

Rapid, marked elevation in Ca2+ influx through the opening of L-type VDCCs is required for first-phase secretion [55,56]. In cultured primary ELKS-KO β-cells, the first glucose-evoked [Ca2+]i rise was decreased compared with that of control β-cells (Figure 3C). [Ca2+]i rise induced by high K+ was also lower in ELKS-KO β-cells (Figure 3D). Nifedipine, an L-type VDCC inhibitor, diminished the glucose- and high K+-evoked increase in [Ca2+]i [57], in control and ELKS-KO β-cells. This suggests that the deletion of ELKS impairs the increase in [Ca2+]i via L-type VDCCs. In addition, the reduced glucose- and high K+-induced [Ca2+]i increase in ELKS-KO β-cells was similarly restored by both ELKS<sup>lox/lox</sup> and ELKS<sup>−/−</sup> gene transfer, suggesting that these isoforms are important for this [Ca2+]i rise in β-cells.

How does ELKS control the entry of Ca2+ into β-cells via type L VDCCs? ELKS and CAST have been shown to bind to VDCC-β1 subunits in mouse islets [58]. In cultured primary MIN6 cells, we found that ELKS binds to VDCC-β1 subunits in islets isolated from ELKS-KO mice via immunoprecipitation (Figure 3E). We generated ELKS±/− mice by crossing ELKS<sup>lox/lox</sup> mice with RIP-Cre mice expressing Cre recombinase in pancreatic β-cells [50] in order to examine the potential role(s) of ELKS in pancreatic β-cells. In islets isolated from ELKS±/− mice, the expression levels of ELKS isoforms were markedly decreased, but no changes were detected in terms of the expression of glucokinase, GLUT2, SNARE proteins required for insulin exocytosis, and active zone proteins (Bassoon, RIM2, and Munc13-1). We confirmed reduction of ELKS expression in insulin-positive β-cells in ELKS±/− mice islets via immunostaining of pancreatic sections (Figure 3A).

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Figure 3: ELKS controls Ca^{2+} influx via L-type Ca^{2+} channels and first-phase insulin secretion. (A) Immunohistochemical staining for insulin and ELKS in pancreatic islets from control and ELKS βKO mice. Scale bars = 50 μm. (B) Insulin secretion in perfused pancreas in response to glucose (16.7 mM). (C, D) Fura-2 Ca^{2+} imaging in β-cells from control and ELKS βKO mice stimulated with glucose (22 mM) (C) and high K^+ (40 mM) (D). (E) ELKS deficiency reduces the L-type voltage-dependent Ca^{2+} current. The current (pA/pF)–voltage (mV) relationships recorded in control and ELKS-KO β-cells with or without of nifedipine are shown (nif: 1 μM). (F, G) G-CaMP8b Ca^{2+} imaging of the β-cell plasma membrane in control (F) and ELKS-KO (G) islets stimulated with glucose (22 mM). The asterisk indicates the vasculature labeled with tomato lectin. (H, I) Typical time course of the fluorescence intensity of G-CaMP8b in the three regions of the β-cell plasma membrane (the vascular-facing plasma membrane, lateral region, and side opposite the vasculature) during glucose stimulation in islets from db/m+ mice (H) and db/db mice (I). Scale bars = 10 μm. Time stamps (min:sec:msec) are shown for each image. Results are presented as the means ± SEM. *P < 0.05; **P < 0.01. Adapted from Ohara-Imaizumi et al. (2019)[18].
regulated by ELKS through L-type VDCCs. In addition, we confirmed the functional effect of ELKS on L-type VDCCs under whole-cell recording using human embryonic kidney (HEK293) cells transfec ted with L-type VDCCs (CaV1.2/β2a/α2δ) in the presence or absence of ELKS overexpression [18]. The Ba^{2+} current density within cells co-expressing L-type VDCCs and ELKS was significantly higher than that in control cells expressing L-type VDCCs without ELKS. Results from the exogenous expression experiments support those of the experiments with the ELKS-KO β-cells showing that ELKS potentiates opening of L-type VDCCs.

As described above, ELKS clusters were identified at insulin granule docking sites colocalized with Syntaxin 1A clusters in the plasma membrane of MIN6 cells [17]. Insulin granules are tethered to an assembly of SNARE proteins near L-type VDCCs [61]. Thus, together with previous findings, our results indicate that ELKS binding to the GK domain of VDCC-β2/3 subunits opens L-type VDCCs. This opening of the L-type VDCCs facilitates the influx of Ca^{2+} and insulin exocytosis during first-phase secretion of insulin from pancreatic β-cells (Figure 4A).

We also investigated the molecular mechanism underlying ELKS regulation of VDCC activity. It has been reported that the β subunit associates with the pore-forming VDCC-α1 subunit and contributes to plasma membrane trafficking and voltage-dependent gating of VDCCs [59]. However, surface expression of VDCC-α1 (CaV 1.2) was not affected by ELKS KD in MIN6 β-cells, suggesting that ELKS does not regulate VDCC trafficking to the plasma membrane. In addition, electrophysiological analysis showed that: i) ELKS deletion did not affect steady-state voltage-dependent activation (VDA) or inactivation (VDI) in ELKS-KO β-cells, and ii) the VDI was identical in HEK293 cells transfected with L-type VDCCs present with or without ELKS. These results suggest that ELKS does not affect voltage-dependent gating of L-type VDCCs. Therefore, ELKS may positively regulate L-type VDCCs by a yet unknown mechanism that differs from that of other molecules that have been shown to interact with β subunits, such as RGK (Rem/Gem/Kir) family proteins [62] and RIM1 [63–65]. Further research is needed to clarify the molecular mechanism underlying the control of ELKS-mediated facilitation of L-type VDCCs in β-cells.

Figure 4: Schema for regulation of ELKS during insulin secretion from islet β-cells. (A) ELKS has a role in regulating the opening of L-type VDCCs through direct binding to VDCC-β2/3. (B) In pancreatic islets, ELKS localizes at the vascular-facing plasma membrane of β-cells where it forms a complex with L-type VDCCs for insulin exocytosis. This complex controls the polarity of the initial increase in Ca^{2+} and first-phase insulin secretion into the central venous vasculature. Adapted from Ohara-Imaizumi et al. (2019) [18].
6. ELKS IS IMPORTANT FOR THE INFUX OF Ca^{2+} AND INSULIN SECRETION AT THE VASCULAR-FACING PLASMA MEMBRANE OF β-CELLS

ELKS has been reported to be preferentially enriched at the vascular side of the β-cell plasma membrane, whereas VDCC-α1 (CaV1.2) immunostaining was observed on the entire β-cell plasma membrane, as were exocytic t-SNARE proteins within mouse islets [18]. These data suggest that ELKS may control L-type VDCCs on the vascular-facing plasma membrane in islet β-cells. If so, Ca^{2+} signals via VDCCs should be detectable initially at the vascular-facing plasma membrane during the first phase of insulin secretion. To test this hypothesis, we investigated Ca^{2+} signals at the vascular side of the β-cell plasma membrane in situ in islets using a newly synthesized Ca^{2+} probe employed as an improved high affinity version of GCaMP [66,67], and Dylight 594-labeled tomato lectin, a marker for vasculature [68]. In control islets, Ca^{2+} imaging during glucose stimulation showed that Ca^{2+} signals initially increased at the vascular-facing plasma membrane of the β-cells, and then spread across the entire plasma membrane (Figure 3F). Analysis of this increase in Ca^{2+} in each plasmalemmal region of β-cells showed that the Ca^{2+} signal increased first at the vascular face, with the Ca^{2+} signal gradually increasing initially in the lateral regions and at the side of the plasma membrane opposite the vasculature (Figure 3H). Conversely, ELKS-KO β-cells exhibited a reduced initial increase in Ca^{2+} at the vascular-facing plasma membrane (Figure 3G). Ca^{2+} fluorescence intensity increased similarly in all regions (Figure 3I); these increases in Ca^{2+} were hampered by L-type VDCC blockers within control and ELKS-KO islets. These findings suggest that the plasma membrane of β-cells is functionally heterogeneous in Ca^{2+} rise patterns, and implies that the ELKS-regulated Ca^{2+} rise at the vascular-facing plasma membrane is important for first-phase insulin secretion. We therefore propose that ELKS and L-type VDCCs form an insulin secretion complex at the ELKS-localized vascular side of the β-cell plasma membrane for initial polarized Ca^{2+} influx and first-phase insulin secretion from pancreatic islets (Figure 4B). Use of another Ca^{2+} probe in mouse islets, R-CaMP1h, revealed a Ca^{2+} microdomain in the vascular-facing plasma membrane and in the lateral regions; however, which Ca^{2+} microdomain first appeared remains unclear [69]. The differences from our results may be caused by use of R-CaMP1h in the cytosol. R-CaMP1h has low affinity for Ca^{2+} (Kd: ~1.3 μM) [70], whereas Ca^{2+} affinity (Kd: ~44 nM) [18]. In the future, imaging of the increase in Ca^{2+} and insulin granule exocytosis should be performed simultaneously in islet β-cells. In this case, two-photon imaging of islets transplanted into the anterior chamber of the eye may be a useful tool enabling visualization of in situ β-cell function in a setting where vascular supplies are rewired [71–73].

7. ELKS AND DIABETES

Oral glucose tolerance tests have demonstrated impaired glucose tolerance in ELKS-KO mice. Hence, ELKS is required to maintain blood glucose homeostasis, which it accomplishes through its facilitation effect on the Ca^{2+} increase and first-phase secretion of insulin from β-cells. It is intriguing that db/db mice, a murine model of spontaneous type 2 diabetes, demonstrated reduced ELKS expression within islets. These db/db mice displayed higher fasting blood glucose levels and lower insulin secretion from islets compared with nondiabetic db/+ mice. Within this model the glucose-induced [Ca^{2+}], rise showed a slower and decreased peak response versus that of islets from db/+ mice, which was consistent with previous reports [74,75]. These findings led us to suspect the possible involvement of decreased expression of ELKS in diabetes. To assess this idea, we performed in situ G-CaMP8b Ca^{2+} imaging of β-cells within islets from db/db mice and compared these data with that of β-cells from db/+ mice. The Ca^{2+} signal initially increased in the vascular-facing region within db/+ mouse islet β-cells (Figure 3J), similar to that observed in the control mouse islet β-cells. On the other hand, db/db mouse islet β-cells showed an initially impaired Ca^{2+} rise at the vascular-facing plasma membrane, and Ca^{2+} levels were elevated equally throughout the plasma membrane regions, similar to that of the ELKS-KO mouse islet β-cells (Figure 3K). Therefore, we speculated that the downregulated ELKS expression may play a role in impaired insulin secretion within type 2 diabetes via a reduced polarized Ca^{2+} rise. Nevertheless, further studies are needed to understand the relationship between downregulated ELKS expression and diabetes.

8. WHAT IS THE MOLECULAR MECHANISM BEHIND ELKS DELIVERY TO THE VASCULAR-FACING PLASMA MEMBRANE OF β-CELLS?

The findings above raise new questions. What are the molecular cues for ELKS resulting in its localization to the vascular-facing plasma membrane in pancreatic β-cells? How does ELKS organize the secretory hotspots?

For heterogeneous localization of ELKS, there is likely a polarity cue enabling ELKS to localize to the vascular side of the plasma membrane. One possible mechanism may involve communication between β-cells and endothelial cells [76]. Endothelial cells form capillaries and secrete extracellular matrix (ECM) proteins to generate a basement membrane interacting with adjacent β-cells [77,78]. Very recently, Gan et al. [79] reported that contact between the ECM and vascular side of the plasma membrane in β-cells triggers spatially confined activation of β1-integrin-dependent focal adhesion in β-cells, which orients the β-cells and directs targeting of insulin granule fusion to the contact sites. Interestingly, activation of β1-integrin recruits LL5β1, a phosphatidylinositol-3,4,5-triphosphate (PIP3)-binding protein, to the rim of focal adhesion [80], and LL5β1 has been reported to interact with ELKS in Ins-1 β-cells [48]. In addition, the ECM protein laminin interacts extracellularly with VDCCs, which can intracellularly interact further with active zone proteins, including ELKS, Bassoon, and Piccolo, to organize active zones in neuromuscular junctions [81,82]. Therefore, it is possible that focal adhesion activated by local interaction between β1-integrin in β-cells and ECM proteins secreted from endothelial cells localizes ELKS to the vascular side of the plasma membrane via recruitment of LL5β1 and organizes secretory hotspots. However, this should be further examined in β-cells.

9. OTHER ACTIVE ZONE PROTEINS

ELKS has been reported to bind other active zone proteins, implying that these proteins are also present at the ELKS-localized vascular side of the β-cell plasma membrane. Immunostaining research has recently shown that RIM2, Piccolo, and Liprin-α were also present at the vascular-facing plasma membrane of β-cells, although RIM2 and Piccolo are also apparently diffusely located in the β-cell cytosol [9]. We also observed that Bassoon was localized at the vascular-facing plasma membrane of β-cells (unpublished data). Therefore, it is necessary to define the relationship between ELKS and other active zone proteins at the vascular-facing plasma membrane of β-cells.
during polarized insulin secretion within islets. We summarize the current findings below concerning interactions among ELKS and other active zone protein members within the insulin secretory system (Figure 5).

9.1. RIM2
RIM2 was first identified as a binding partner for Epac2, a cAMP sensor in MIN6 cells [83]. A study using RIM2-null mice revealed that RIM2 is necessary for insulin granule docking to the plasma membrane and for the subsequent priming step, a process required for insulin granules to acquire competence for exocytosis in response to a Ca\(^{2+}\) rise during insulin secretion from β-cells [41]. RIM2 is a multidomain protein interacting with several proteins through its functional domains. While RIM1 interacts with VDCC-β in neurons [65], RIM2 interacts with VDCC-α\(_1\) (Ca\(_{\alpha}1.2\)) and suppresses the VDI of VDCC currents [41,84]. RIM2 has also been reported to interact with the C-terminal IWA region of ELKS. Because ELKS binds directly to VDCC-β through the N-terminal region, RIM2 is not likely involved in the ELKS control of VDCCs. Similar to ELKS\(\alpha\) (with the IWA region), ELKS\(\alpha\) (lacking IWA region) rescued the impairment of the [Ca\(^{2+}\)]\(_i\) increase in ELKS-KO β-cells [18]. However, the relationship between ELKS and RIM2 in VDCC regulation remains unclear.

9.2. Munc13-1
Munc13-1 is a mammalian homolog of the Unc13 synaptic protein in Caenorhabditis elegans [85]. As revealed by experiments with C. elegans [86] and mammalian neurons [87], Munc13-1 is required for the exocytosis priming step of insulin granules [39,88]. Given that Munc13-1 is dispensable for the docking process in neurons and chromaffin cells [87,89], the docking process of insulin granules is likely to be independent of Munc13-1. Munc13-1 contains a C1-domain (homologous to the phorbol ester- and diacylglycerol-binding region of protein kinase C), three C2-domains (Ca\(^{2+}\)-binding and protein-protein interaction domains), and a calmodulin binding sequence and a Syntaxin-binding domain essential for the priming process [90]. Munc13-1 interacts with the VDCC-α\(_1\) (Ca\(_{\alpha}2.2\)) through the C2-domain C2B to regulate its gating properties in nerve terminals [91]. Whether Munc13-1 can physically interact with the α-subunit of VDCCs in β-cells remains uncertain. However, competitive inhibition experiments using the C2B domain of Munc13-1 revealed that functional interaction between Munc13-1 and Ca\(_{\alpha}1.2\) is likely important for arrangement of L-type VDCCs near the docked insulin granules in order to facilitate granule exocytosis [92]. ELKS has been reported to indirectly bind to Munc13-1 through RIM1 [12,16]; hence, it is important to unravel the functional relationship between these three proteins within Ca\(^{2+}\) influx and insulin secretion. Notably, it has been reported that Munc13-1 levels in pancreatic islets were reduced in type 2 diabetic patients as well as type 2 diabetic model rats [40,93].

9.3. Bassoon/Piccolo
Bassoon and Piccolo are large scaffolding proteins present in synaptic active zones. These proteins are structurally related multidomain proteins with 10 highly conserved regions [94]. Bassoon interacts with ELKS, and its binding is crucial for the docking of insulin granules to the plasma membrane with subsequent insulin secretion from MIN6 cells [17]. Piccolo is involved in insulin secretion evoked by the cAMP analog, 8-Br-cAMP, but not by glucose alone in MIN6 cells [38]. Both Bassoon and Piccolo interact with VDCC-α\(_1\) directly or indirectly, and Bassoon affects VDCC activity [84,95]. Both Bassoon and Piccolo interact with a multitude of proteins, including ELKS, RIM2, and Munc13-1.

9.4. Liprin-α
The scaffold protein, Liprin-α, localizes to the vascular-facing plasma membrane in β-cells [9], but so far there are no reports revealing the role of Liprin-α in insulin secretion. Because Liprin-α is required for neurotransmitter release in synapses [96,97], and it interacts with ELKS, RIM, and CASK [98], it is also potentially involved in insulin secretion from β-cells.

10. CONCLUSIONS
Our analyses indicate that ELKS has a role in opening L-type VDCCs at the vascular-facing plasma membrane of β-cells, which leads to an initial polarized Ca\(^{2+}\) influx and first-phase insulin secretion from islets. Our model further clarifies active zone protein control of the polarity of the Ca\(^{2+}\) influx and insulin secretion from β-cells. ELKS maintains blood glucose homeostasis through its facilitation effect on the Ca\(^{2+}\) rise and first-phase secretion of insulin from β-cells. ELKS expression is reduced in diabetic mouse islets with impaired polarized increase of Ca\(^{2+}\) at the vascular-facing plasma membrane of β-cells and insulin secretion. Considering these findings together, ELKS may be a therapeutic target for improvement of glycemic control in type 2 diabetes, which includes abnormalities in first-phase insulin secretion. Further experiments are required to investigate these possibilities in human islets, which differ from mouse islets in structure and have β-cells interspersed with other endocrine cells [99,100].

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CONFLICT OF INTEREST

None declared.
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