Roles of voltage-gated Ca2+ channel subunits in pancreatic cells

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Roles of voltage-gated Ca\(^{2+}\) channel subunits in pancreatic β cells

ABDULLA S. KAZIM

FACULTY OF MEDICINE | LUND UNIVERSITY
Roles of voltage-gated Ca$^{2+}$ channel subunits in pancreatic β cells

Abdulla S. Kazim
Hallmarks of type 2 diabetes (T2D) include elevated blood glucose and free fatty acids (FFAs) as a result of impaired β cell insulin secretion and decreased β cell mass. The glucose-stimulated insulin secretion (GSIS) in β cells is triggered by depolarization-evoked Ca$^{2+}$ entry through voltage-gated Ca$^{2+}$ (Ca$v$) channels. The majority of Ca$v$ channels are believed to reside in cholesterol-rich membrane microdomains called membrane rafts. Ca$^{2+}$ channels consist of the main pore-forming α subunit and three auxiliary subunits, β, αδ, and γ. The roles of the Ca$v$ auxiliary subunits and the membrane rafts in pancreatic β cells are not fully understood, but we have recently shown that the TCF7L2 gene, associated with the strongest genetic risk factor of T2D, regulates Cacna2d1 (αδ1).

This thesis aims to elucidate the roles of β1, β2a, and αδ1 subunits, as well as membrane rafts, in regulating the α1 subunit and, in turn, insulin secretion and β cell survival. Human islets from donors with T2D contained decreased membrane rafts. A similar phenotype was also observed in the diabetic rat model Goto Kakizaki (GK) rat islets. Cholesterol depletion in healthy human islets by cholesterol oxidase (CO) reduced membrane rafts, resembling islets from donors with T2D. Cholesterol depletion resulted in elevated basal insulin release in both human and rat islets. The reason for this appeared to be the declustering of Cav1.2, elevation in basal Ca$^{2+}$ oscillations, and an increase in single-Ca$v$ channel activity as observed in patch-clamp experiments.

When suppressing the Tcf7l2 gene, αδ1 (mRNA and protein) was downregulated and intracellular Ca$^{2+}$ was reduced as measured by confocal microfluorimetry. The decrease in Cacna2d1 expression resulted in Ca$v$ channel internalization in the recycling endosomes. This lowered the whole-cell Ca$^{2+}$ current and decreased insulin secretion.

Human gene expression analysis showed that both Cacnb1 (β1) and Cacnb2a (β2a) genes are abundant in pancreatic islets. When examining the GK rat islets, the expression of both genes was downregulated. Immunoblot experiments showed that high glucose treatment also reduced protein levels of β1 and β2a in INS-1 832/13 cells. Silencing the β1 subunit reduced insulin secretion, which may be due to the observed decrease in whole-cell Ca$^{2+}$ currents. By contrast, β2a suppression did not affect insulin release. When comparing the palmitoylation state of β2a, cells overexpressing the non-palmitoylated β2a had a decreased membrane expression of both β2a and αC. However, overexpression of palmitoylated β2a increased intracellular Ca$^{2+}$, although without affecting secretion. FFA (palmitate) treatment reduced intracellular Ca$^{2+}$ under stimulatory conditions thus decreasing GSIS. Cells that either lack β1 or express excess palmitoylated β2a have increased risk of apoptosis. These data reveal novel roles of membrane rafts and β1, β2a, and αδ1 subunits in regulating Ca$v$ channel trafficking and activity, thus influencing β cell function and survival.

Key words: Type 2 diabetes, β cell, insulin, voltage-gated Ca$^{2+}$ channel, calcium, membrane raft, auxiliary subunits, TCF7L2, cholesterol.
Roles of voltage-gated Ca\textsuperscript{2+} channel subunits in pancreatic β cells

Abdulla S. Kazim
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This thesis is warmly dedicated to:

My mother Durreya and my brothers Huthaifa and Orwa
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## Abbreviations

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<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>1,4-dihydropyridine</td>
<td>DHP</td>
</tr>
<tr>
<td>α-interaction domain</td>
<td>AID</td>
</tr>
<tr>
<td>Adenosine triphosphate</td>
<td>ATP</td>
</tr>
<tr>
<td>AID-binding pocket</td>
<td>ABP</td>
</tr>
<tr>
<td>ATP-sensitive potassium channel</td>
<td>$K_{\text{ATP}}$</td>
</tr>
<tr>
<td>β-interaction domain</td>
<td>BID</td>
</tr>
<tr>
<td>Bovine serum albumin</td>
<td>BSA</td>
</tr>
<tr>
<td>Ca$^{2+}$ modulated protein</td>
<td>Calmodulin</td>
</tr>
<tr>
<td>Calmodulin</td>
<td>CaM</td>
</tr>
<tr>
<td>Calmodulin kinase</td>
<td>CaMK</td>
</tr>
<tr>
<td>Carboxypeptidase E</td>
<td>CPE</td>
</tr>
<tr>
<td>Cholesterol oxidase</td>
<td>CO</td>
</tr>
<tr>
<td>Detergent-resistant membrane</td>
<td>DRM</td>
</tr>
<tr>
<td>Diabetes Mellitus</td>
<td>DM</td>
</tr>
<tr>
<td>Endoplasmic reticulum</td>
<td>ER</td>
</tr>
<tr>
<td>Enhanced chemiluminescence</td>
<td>ECL</td>
</tr>
<tr>
<td>Free fatty acid</td>
<td>FFA</td>
</tr>
<tr>
<td>Fasting plasma glucose</td>
<td>FPG</td>
</tr>
<tr>
<td>Genome-wide association study</td>
<td>GWAS</td>
</tr>
<tr>
<td>Glucose-stimulated insulin secretion</td>
<td>GSIS</td>
</tr>
<tr>
<td>Glucose transporter</td>
<td>GLUT</td>
</tr>
<tr>
<td>Glycosylated hemoglobin</td>
<td>HbA$_{1C}$</td>
</tr>
<tr>
<td>Glycosylphosphatidylinositol</td>
<td>GPI</td>
</tr>
</tbody>
</table>
Goto Kakizaki rat
Green fluorescent protein
Guanylate kinase domain
High voltage-gated Ca\(^{2+}\) channel
Inositol 1,4,5-trisphosphate
Intracellular Ca\(^{2+}\) concentration
IP\(_3\) receptor
Low voltage-gated Ca\(^{2+}\) channel
Low density lipoprotein
Methyl β-cyclodextrin
Mitochondrial Ca\(^{2+}\) uniporter
Mitochondrial Na\(^+\)/Ca\(^{2+}\) exchanger
Neurogenin 3
Non-esterified fatty acid
Oral glucose tolerance test
Pancreatic and duodenal homeobox factor 1
Pancreatic polypeptide
Paraformaldehyde
Phosphatase 2A
Plasma membrane Ca\(^{2+}\)-ATPase
Polyacrylamide gel electrophoresis
Polyvinylidene difluoride
Pore loop
Prohormone convertase
Protein kinase A
Protein kinase C
Red blood cell
RNA-induced silencing complex
Ryanodine receptor

GK
GFP
GK
HVGCC
IP\(_3\)
[Ca\(^{2+}\)]\(_i\)
IP\(_3\)R
LVGCC
LDL
MβCD
mCU
mNCX
Ngn3
NEFA
OGTT
PDX1
PP
PFA
PP2A
PMCA
PAGE
PVDF
P-loop
PC
PKA
PKC
RBC
RISC
RyR
<table>
<thead>
<tr>
<th>Term</th>
<th>Abbreviation</th>
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<tbody>
<tr>
<td>Sarcoendoplasmic reticulum Ca\textsuperscript{2+}-ATPase pump</td>
<td>SERCA</td>
</tr>
<tr>
<td>Signal recognition particle</td>
<td>SRP</td>
</tr>
<tr>
<td>Single-nucleotide polymorphism</td>
<td>SNP</td>
</tr>
<tr>
<td>Sodium dodecyl sulfate</td>
<td>SDS</td>
</tr>
<tr>
<td>Soluble N-ethylmaleimide-sensitive factor attachment protein receptor</td>
<td>SNARE</td>
</tr>
<tr>
<td>Src homology 3 domain</td>
<td>SH3</td>
</tr>
<tr>
<td>Synaptosomal-associated protein 25</td>
<td>SNAP-25</td>
</tr>
<tr>
<td>Synaptotagmin 1</td>
<td>SYT1</td>
</tr>
<tr>
<td>Trans-Golgi network</td>
<td>TGN</td>
</tr>
<tr>
<td>Transcription factor 7-like 2</td>
<td>TCF7L2</td>
</tr>
<tr>
<td>Tricarboxylic acid</td>
<td>TCA</td>
</tr>
<tr>
<td>Triglyceride</td>
<td>TG</td>
</tr>
<tr>
<td>Type 1 diabetes</td>
<td>T1D</td>
</tr>
<tr>
<td>Type 2 diabetes</td>
<td>T2D</td>
</tr>
<tr>
<td>United States dollar</td>
<td>USD</td>
</tr>
<tr>
<td>V-maf musculoaponeurotic fibrosarcoma oncogene homology A</td>
<td>MafA</td>
</tr>
<tr>
<td>V-maf musculoaponeurotic fibrosarcoma oncogene homology B</td>
<td>MafB</td>
</tr>
<tr>
<td>Vesicle-associated membrane protein 2</td>
<td>VAMP-2</td>
</tr>
<tr>
<td>Voltage-gated Ca\textsuperscript{2+} channel</td>
<td>( \text{Ca}_V )</td>
</tr>
<tr>
<td>Voltage-gated K\textsuperscript{+} channel</td>
<td>( \text{K}_V )</td>
</tr>
</tbody>
</table>
Introduction

Diabetes Mellitus

Diabetes mellitus (DM) is a chronic metabolic disorder characterized by elevated blood glucose levels, a state referred to as hyperglycemia. During the 5th century BC, DM was first described as ‘honey-like urine’ [1]. The disease was later termed diabetes and then mellitus in the 2nd and 17th century respectively, followed by the discovery of insulin in the 20th century. Since then, DM has been better understood and thus classified into 3 main types: type 1, type 2, and gestational diabetes.

Type 1 diabetes (T1D), making up 5-10% of diabetic patients, is an autoimmune disease where white blood cells destroy pancreatic β cells causing insulin deprivation [2]. Patients with T1D (also known as insulin-dependent diabetes) are mostly diagnosed as children or adolescents and require regular blood glucose monitoring, a strict diet, and daily insulin injections to maintain normal blood glucose.

Type 2 diabetes (T2D), previously referred to as non-insulin dependent diabetes, is the most common form of the disease accounting for approximately 87-91% of all diabetes cases [3]. In T2D, pancreatic β cells are not destroyed by immune cells but rather lose their function while tissues such as liver, muscle, and fat become resistant to insulin.

A similar phenomenon is also observed in gestational diabetes, a form of diabetes affecting women during pregnancy. Gestational diabetes, occurring in 16% of all pregnancies, increases the risk of pregnancy complications [4]. In addition, patients with gestational diabetes, along with their born child, are more likely to develop T2D later in life. Other types of diabetes also exist, although less common.

In 2015, DM was estimated to have affected 415 million individuals and was responsible for the deaths of 5 million people worldwide [3]. The estimated cost of treatment and prevention of diabetes and its complications was between USD 673-1,197 billion [3, 4].
Type 2 diabetes

Genetic factors
T2D is a chronic metabolic disorder influenced both by environmental and genetic factors. It is characterized by a combination of insulin resistance and reduced insulin output. Insulin is an integral hormone of the glucose-lowering system driven by pancreatic β cells. So far, 90 risk genes have been identified and associated with T2D [5]. Genome-wide association study (GWAS), a widely used method for gene-association, takes advantage of common single-nucleotide polymorphisms (SNPs) that may vary between diabetic and non-diabetic individuals depending on the SNP examined. The SNPs that associate with T2D are labeled genetic risk factors which can then be used to identify susceptible individuals and take preventive measures prior to the onset of the disease. In 2000, the first gene linked to T2D was identified as CAPN10 (calpain 10) [6]. Following that, many candidate genes were investigated and 3 were found to be associated with the disease, with TCF7L2 (transcription factor 7-like 2) having the strongest association among all ethnic groups [7]. The transcription factor TCF7L2 is part of the Wnt signaling pathway that is involved in regulating glucose-stimulated insulin secretion (GSIS) and β cell growth and survival [8, 9]. Since 2007, many GWAS studies have been conducted to identify novel T2D genetic risk factors [10]. One interesting discovery was the link between T2D and obesity through the FTO (fat mass and obesity-associated protein) gene [11]. This gene is linked to body mass index (BMI) and is responsible for an increase in adiposity and food intake in mice and humans [12, 13]. This is interesting, as patients with T2D have a lipid disorder where cholesterol and non-esterified fatty acids (NEFAs) are elevated in the blood [14-17].

Environmental factors
T2D is an interaction between both genetic and environmental factors. Environmental factors, such as low physical activity and excess calorie intake or unhealthy food intake, could lead to the development of some clinical risk factors like high BMI, elevated fasting plasma glucose (FPG), and high serum concentrations of TG and cholesterol. This would contribute to the development of T2D; however, it is important to consider the genetic variation between individuals as it often explains why some have low BMI and still develop the disease while others who have high BMI do not. Nevertheless, clinical risk factors alone are strong predictors of future diabetes [18]. For example, a combination of high levels of NEFAs in the blood, high BMI, and old age could be indicators of the start of insulin resistance and thus enable early detection and intervention.

The prevalence and risk of diabetes also differs depending on ethnicity [19]. An ethnic group normally shares a common gene pool which allows for certain genetic traits to prevail, including diabetes gene variants. For example, Asians have a lower
obesity threshold than Caucasians, and thus the risk of T2D among Asians is higher [20]. Another example is the difference between two Pima Indian populations, one residing in mountains in northwestern Mexico and the other in Arizona, USA [21]. Although both populations share similar genetic makeup, Arizona Pimas have a much higher prevalence of obesity and T2D than Mexican Pimas. The reason behind this is the different lifestyles associated with the two populations, one a ‘traditional’ while the other an ‘abundance’ lifestyle.

**Insulin resistance**

The cause of T2D is insulin resistance followed by impaired insulin secretion. Obesity is the result of overnutrition and it strongly correlates with insulin resistance [22]. Initially, frequent food intake, and in turn incretin hormones, cause a continuous pancreatic release of insulin into the blood. Insulin-target tissues such as liver, skeletal muscle, and fat, become desensitized to insulin, and hence, reduce glucose uptake. This signals the pancreatic β cell to increase output as the current level is insufficient to lower blood glucose.

**β cell mass and function**

In T2D-susceptible individuals, β cells eventually ‘burn out’ from increased insulin release, thus losing their function and undergoing apoptosis. Many studies have reported a reduction in β cell mass (24-65%) in patients with T2D [23-28]. This is further supported by studies showing a reduced number and size of pancreatic islets from patients with T2D [29]. Early studies in rat have shown that reduced β cell mass is the main contributor to T2D instead of β cell dysfunction, since lower number of β cells equates to less plasma insulin [30]. However, it has been shown that human donors who underwent pancreatectomy (30-50% pancreas removal) did not develop T2D albeit having impaired glucose tolerance, suggesting that β cell dysfunction is the main cause of the disease rather than β cell mass [27, 31-33]. In fact, several studies have reported that the β cell capacity to release insulin has been reduced between 50-97% in patients with T2D [28, 34-36]. Patients with T2D subjected to bariatric surgery or short-term caloric restriction restore their blood glucose within days, supporting the contribution of β cell function rather than mass to this quick restoration [37, 38].

**Toxicity**

Excess energy intake along with a decline in insulin sensitivity results in glucotoxicity and lipotoxicity. The chronically elevated levels of glucose, cholesterol, and NEFAs in the blood exert detrimental effects on insulin-target tissues and pancreatic β cells [39]. For example, liver, muscle, and fat cells release glucose, reduce glucose uptake, and secrete fatty acids into the bloodstream, respectively, under glucolipotoxic conditions. In the case of β cells, elevated glucose
hampers the actions of the transcription factors pancreatic and duodenal homeobox factor 1 (PDX1) and v-maf musculoaponeurotic fibrosarcoma oncogene homology A (MafA), in turn reducing insulin gene expression [40]. In addition, elevated glucose levels increase the demand for insulin output, thus putting load on the endoplasmic reticulum (ER) to synthesize more insulin. This induces ER stress due to the accumulation of misfolded proteins, consequently triggering the unfolded protein response (UPR) [41]. Persistent ER stress and UPR will ultimately result in β cell death via apoptosis. It has been shown that NEFAs hinder insulin release, albeit transiently potentiating secretion [14, 42]. Unlike unsaturated FFAs, saturated FFAs such as palmitate induce ER stress, resulting in cell apoptosis and decreased β cell mass [43]. As in the ER stress, an increase in cellular metabolism increases mitochondrial workload yielding more reactive oxygen species (ROS), resulting in oxidative stress. Islets of T2D patients were shown to have increased markers of oxidative stress which correlated with impairment of GSIS [44]. ROS have deleterious effects in β cells such as disrupting insulin synthesis, mitochondrial membrane, and DNA, as well as increasing ER stress, leading to dysfunction and apoptosis [45-48]. Hyperglycemia also causes a reduction in the number, morphology, and function of mitochondria which in turn diminishes ATP production [49]. Disruption of Ca\(^{2+}\) homeostasis in the form of elevated levels of mitochondrial Ca\(^{2+}\), ER Ca\(^{2+}\) store depletion, reduced Ca\(^{2+}\) influx, and chronic increase in intracellular Ca\(^{2+}\) will negatively impact β cell function and mass [50]. In addition to glucose, oxidized low density lipoprotein (LDL) can also reduce preproinsulin expression [51].

**Diagnosis**

Early detection of T2D risk factors is critical for prevention of T2D. The FPG method is used to diagnose diabetic and pre-diabetic individuals. The fasting takes place for ≥ 8 hours followed by a blood glucose measurement [52]. A similar blood-measuring method, termed oral glucose tolerance test (OGTT), requires the individual to drink a glucose load of 75 g 2 hours prior to measuring blood glucose. A third method has recently been adopted which, instead of plasma glucose, measures hemoglobin A\(_{1C}\) (HbA\(_{1C}\)) [52]. Hemoglobin is a protein found in red blood cells (RBCs) that binds to oxygen. Interestingly, hemoglobin also binds to glucose and transforms into glycosylated hemoglobin, or HbA\(_{1C}\). Hence, HbA\(_{1C}\) is an accurate measure for the average levels of plasma glucose during the last months, since it is stable for 8-12 weeks (the lifespan of an RBC). HbA\(_{1C}\) reflects the long-term plasma glucose levels, as opposed to FPG and OGTT where the short-term response to high glucose is shown. The criteria for diagnosing diabetes is listed in Table 1.
### Table 1 Diabetes and prediabetes diagnosis
Criteria for diagnosing diabetes and prediabetes [53].

<table>
<thead>
<tr>
<th>Method</th>
<th>Prediabetes</th>
<th>Diabetes</th>
</tr>
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<tbody>
<tr>
<td>HbA1c</td>
<td>5.7 – 6.4%</td>
<td>≥ 6.5%</td>
</tr>
<tr>
<td>FPG</td>
<td>5.6 – 6.9 mmol/l (100 – 125 mg/dl)</td>
<td>≥ 7 mmol/l (≥ 126 mg/dl)</td>
</tr>
<tr>
<td>OGTT</td>
<td>7.8 – 11 mmol/l (140 – 199 mg/dl)</td>
<td>≥ 11.1 mmol/l (≥ 200 mg/dl)</td>
</tr>
</tbody>
</table>

### Treatment
Metformin is the most common and preferred treatment for T2D. It is usually the first drug given to patients with T2D, along with diet and exercise recommendations. If the treatment strategy is ineffective, another drug is added to the metformin therapy (e.g. sulfonylurea(SU), DPP4 inhibitors, GLP1 analogs, or SGLT2 inhibitors), and if the treatment target is not reached, a third drug is added to the treatment strategy. However, patients with severe kidney or liver problems avoid metformin as part of their treatment. If all treatment plans fail even after an initial success, insulin would be administered for patients with T2D.

### Complications
T2D is a metabolic disorder that affects critical tissues such as the heart, nerves, blood vessels, kidneys, and eyes, if poorly managed. These complications are categorized into microvascular and macrovascular diseases [54]. Microvascular disease refers to damage to small blood vessels and could lead to blindness from retinopathy, kidney failure from nephropathy, and diabetic foot from neuropathy. Macrovascular refers to damage to the large blood vessels and could lead to cardiovascular diseases such as stroke and heart attack.

### The pancreatic islet

#### Composition
In the pancreas, the hormone-secreting endocrine cells, making up 1-2% of the pancreas, reside in highly vascular punctate regions called islets of Langerhans [55]. There are ~3.2 million islets in the human pancreas, each consisting mainly of α cells (glucagon), β cells (insulin), δ cells (somatostatin), γ cells (pancreatic polypeptide), and ε cells (ghrelin). The β cell distribution in the human islet appears to be more scattered as opposed to rodent islets in which the β cells are more focused in the center [56].
Development

During development, pancreatic progenitor cells differentiate into the 5 different cells mentioned earlier depending on the expression of certain transcription factors. For example, PDX1 and neurogenin 3 (Ngn3) drive the progenitor cells into β progenitor cells. Further development of the β cell takes place with the help of MafB while MafA is vital for mature β cell function [57, 58]. MafA is only expressed in insulin-positive cells while MafB is found in both glucagon and insulin-positive cells prior to birth in mice. However, after birth, MafB is specifically expressed in α cells.

The pancreatic β cell

Insulin

The INS gene expression is regulated by glucose and FFAs via PDX1 and MafA [59]. The gene encodes a 110-amino acid preproinsulin that is targeted to the ER lumen [60]. This takes place when the cytosolic ribonucleoprotein signal recognition particle (SRP) interacts with the hydrophobic signal peptide on the preproinsulin N-terminus, transferring it into the ER lumen via the peptide-conducting channel. The enzyme signal peptidase then removes this hydrophobic end to form proinsulin. With the help of ER chaperone proteins, proinsulin is folded and forms 3 disulfide bonds. Upon reaching the trans-Golgi network (TGN) from the ER, proinsulin is sorted into immature insulin secretory granule (ISG) along with ions such as Ca\(^{2+}\), Zn\(^{2+}\), and H\(^{+}\) and various proteins including carboxypeptidase E (CPE) and prohormone convertase (PC) [60, 61]. The ISG matures when Ca\(^{2+}\) is abundant and the pH of the lumen drops, activating both CPE and PC to trim proinsulin into a 51-amino acid insulin and C-peptide. Prior to cleavage, the C-peptide, namely C chain, was situated between A and B chains. After removal of the C chain, A and B chains become attached by disulfide bonds giving rise to mature insulin.

GSIS

Pancreatic β cells act as glucose sensors to respond to fluctuating levels of glucose. When plasma glucose is at basal levels, the cell is said to be in a resting state with a membrane potential of around -70 mV. In the resting state, the ATP-sensitive K\(^{+}\) channel (K\(_{ATP}\)) remains open allowing diffusion of K\(^{+}\) out of the cell. This keeps the intracellular environment more negative and the voltage-gated Ca\(^{2+}\) (Cav) channel remains inactive, and only low levels of insulin are released. When circulating glucose is high, the 6-carbon sugar is taken up by glucose transporters GLUT1, GLUT3, and possibly GLUT2 to start the triggering pathway in GSIS (Fig. 1). GLUT2 (Km 11.2) is believed to be the main glucose transporter in human
pancreatic β cells. While this may be true in rodents, GLUT2 expression levels in human islets have been found to be low [62]. Therefore, it has been suggested that GLUT2 may not be considered the main glucose transporter in human pancreatic β cells [63]. In addition, GLUT1 (Km 6.9) properties are more in agreement with the dose-dependent curve for GSIS (Km 6.5) suggesting that GLUT2 is likely contributing less, if any, to GSIS in human β cells [62, 64, 65]. Although GLUT1 is currently believed to be the primary glucose transporter in human islets, GLUT3 has shown to be equally highly expressed [63]. Once glucose enters, however, it immediately becomes phosphorylated into glucose-6-phosphate and undergoes glycolysis. Two pyruvate molecules are formed from glucose-6-phosphate which then enter the mitochondria where they are converted into acetyl-CoA, enter the tricarboxylic acid (TCA) cycle, and finally yield chemical energies in the form of adenosine triphosphate (ATP). The rise in ATP:ADP ratio blocks the ATP-sensitive K\(^+\) channel (K\(_{\text{ATP}}\)) causing a more positive membrane potential and thus depolarization [66]. This activates the Ca\(_V\) channel permitting the inflow of Ca\(^{2+}\). The rise in intracellular Ca\(^{2+}\), particularly in regions close to the secretory granules, triggers pulsatile insulin exocytosis [67]. In fact, it has been shown that in mouse β cells, the L-type Ca\(_V\) channel interacts with soluble N-ethylmaleimide-sensitive factor attachment protein (SNARE) proteins residing on the insulin granules [61]. SNARE proteins facilitate granular fusion with the plasma membrane. There are around 10,000 insulin-containing granules in a rat pancreatic β cell with an average of 120 mM insulin concentration [68]. Insulin granules are grouped into two pools; the readily releasable pool (RRP; 1-5%) and the reserve pool (RP; 95-99%). Insulin granules in the RRP localize to the plasma membrane ready for exocytosis, whereas the ones in the RP do not. The first spike of insulin release lasts for approximately 10 minutes and is referred to as the 1\(^{\text{st}}\) phase insulin secretion [68, 69]. By contrast, the 2\(^{\text{nd}}\) phase insulin secretion occurs gradually over a longer period of time. It is believed that the RRP is responsible for the 1\(^{\text{st}}\) phase insulin release while the RP accounts for the 2\(^{\text{nd}}\) phase. This biphasic behavior of insulin secretion allows β cells to immediately respond to a sudden increase in plasma glucose as well as maintaining long-term blood glucose homeostasis.

In addition to the effect of Ca\(^{2+}\) on exocytosis, it also upregulates the insulin gene, INS [70]. This occurs through a separate pathway in GSIS which involves cyclic adenosine monophosphate (cAMP) [71, 72]. The production of cAMP is stimulated by Ca\(^{2+}\), ATP, and/or gut hormones called incretins [73, 74]. Once upregulated, cAMP initiates a downstream signaling pathway activating protein kinase A (PKA) and Epac2A, which stimulate insulin secretion [72, 75]. PKA is believed to activate cAMP responsive element binding protein (CREB) promoting the insulin gene INS for further hormone synthesis [76, 77]. Moreover, in the amplifying pathway, cAMP/PKA signaling may also facilitate the transport of glutamate, a product of glucose metabolism, into insulin granules and stimulate insulin secretion [72].
Membrane rafts

The cell membrane is made of two layers of phospholipids that join to form an inner hydrophobic lipid part surrounded by hydrophilic phosphate heads. The fluidity of the membrane is affected by the number of cholesterol molecules embedded in it; the more cholesterol, the less fluid the membrane is and thus less permeable [78]. The more permeable the membrane becomes, the less control it has over cellular content. The membrane also consists of cholesterol- and sphingolipid-enriched microdomains that are resistant to detergents and hence were termed detergent-resistant membranes (DRMs) or membrane rafts.

Membrane rafts serve as docking platforms for transport proteins and channels, while also aiding in protein interaction and stability [79]. During exocytosis, SNARE proteins form SNARE fusion complexes that depend on membrane rafts to interact and facilitate the fusion of mature secretory granules to release insulin [80]. In addition, (GPI)-anchoring requires membrane rafts to localize and stabilize proteins to the plasma membrane [81]. To identify the functions of membrane rafts,
two cholesterol-targeting agents are widely used, methyl β-cyclodextrin (MβCD) and cholesterol oxidase (CO). When pancreatic β cells were treated with MβCD, the voltage-gated K⁺ channel, \( \text{K}_V \), resulted in reduced K⁺ amplitude and channel activity [82].

Cav channels

The nomenclature of Cav channels has traditionally differed in different fields. Electrophysiologists used a naming system depending on the Cav channel’s biophysical and pharmacological properties (L, P/Q, N, R, T), and biochemists adopted Greek letters to distinguish between the different subunits (α₁, β, α₂δ, γ) [83, 84]. Molecular biologists concurrently used alphabetical letters to name Cav channel genes (CACNA1A-I, CACNA1S) [85]. In 2000, Cav channels were categorized into 3 families, Cav1, Cav2, and Cav3, based on gene sequence analysis [86]. The nomenclature of the 10 Cavα₁ genes and proteins are listed in Table 2. In human pancreatic β cells, the Cav channels largely contributing to GSIS are the L-type (Cav1.2 and Cav1.3) and P/Q type (Cav 2.1), whereas other types such as T-type (Cav3.2) contribute to a lesser extent [87, 88].

<table>
<thead>
<tr>
<th>Type</th>
<th>α₁ (gene)</th>
<th>Cav</th>
<th>Channel Gating</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-type</td>
<td>α₁S (CACNA1S)</td>
<td>Cav1.1</td>
<td>HVA</td>
</tr>
<tr>
<td></td>
<td>α₁C (CACNA1C)</td>
<td>Cav1.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>α₁D (CACNA1D)</td>
<td>Cav1.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>α₁F (CACNA1F)</td>
<td>Cav1.4</td>
<td></td>
</tr>
<tr>
<td>P/Q-type</td>
<td>α₁A (CACNA1A)</td>
<td>Cav2.1</td>
<td>HVA</td>
</tr>
<tr>
<td>N-type</td>
<td>α₁B (CACNA1B)</td>
<td>Cav2.2</td>
<td>HVA</td>
</tr>
<tr>
<td>R-type</td>
<td>α₁E (CACNA1E)</td>
<td>Cav2.3</td>
<td>HVA</td>
</tr>
<tr>
<td>T-type</td>
<td>α₁G (CACNA1G)</td>
<td>Cav3.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>α₁H (CACNA1H)</td>
<td>Cav3.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>α₁I (CACNA1I)</td>
<td>Cav3.3</td>
<td></td>
</tr>
</tbody>
</table>

HVA, high voltage activated; LVA, low voltage activated.

Voltage-gated Ca²⁺ channels (VGCCs) are expressed in the plasma membranes of excitable cells such as nerves, myocytes, retinal cells, and endocrine cells to quickly relay biological and electrical signals such as Ca²⁺ and membrane depolarization. Upon depolarization, the VGCC undergoes a conformational change which either increases or decreases the affinity for extracellular Ca²⁺ [89, 90]. The VGCCs are heteromeric complexes composed of the main α₁ subunit and auxiliary β, α₂δ, and γ subunits that work together to transport Ca²⁺ into the cell (Fig. 2). The VGCCs can be inhibited by channel type-specific Ca²⁺ channel inhibitors, for example, L-type
(1,4-dihydropyridines (DHPs); isradipine) and N-type (ω-conotoxins) CaV channel blockers.

Figure 2 The CaV channel
A schematic illustration of the CaV channel with all its subunits.

Other Ca\(^{2+}\) channels
Other Ca\(^{2+}\) channels also exist in β cells that are voltage-independent and contribute to maintaining intracellular Ca\(^{2+}\) homeostasis. These include ryanodine receptor (RyR), sarcoendoplasmic reticulum Ca\(^{2+}\)-ATPase (SERCA) pump, and inositol 1,4,5-trisphosphate (IP\(_3\)) receptor (IP\(_3\)R) which are found in the ER membrane and regulate Ca\(^{2+}\) transport [91]. Similar to VGCC localization, transient receptor potential (TRP) channel and plasma membrane Ca\(^{2+}\)-ATPase (PMCA) are also situated in the plasma membrane to control Ca\(^{2+}\) transport. The mitochondrial Ca\(^{2+}\) uniporter (mCU) and mitochondrial Na\(^+\)/Ca\(^{2+}\) exchanger (mNCX) reside in the inner mitochondrial membrane to facilitate the transport of Ca\(^{2+}\) across the mitochondrial membrane.

Additional effects of Ca\(^{2+}\)
Aside from promoting insulin maturation and secretion, Ca\(^{2+}\) also activates protein kinase C (PKC) which in turn promotes insulin exocytosis in β cells [92, 93]. Ca\(^{2+}\) can also drive protein phosphorylation through binding to Ca\(^{2+}\) modulated protein (calmodulin (CaM)), forming a complex that regulates CaM kinases (CaMK).
Moreover, Ca$^{2+}$ entry in some cardiac muscle cells can stimulate the release of Ca$^{2+}$ from sarcoendoplasmic reticulum stores [94].

**α1 subunit**

*Structure and function*

Molecular cloning detected 10 genes in humans that encode the pore-forming Ca$_V$α$_1$ (Table 2). The α$_1$ subunit is a ~170-240 kDa membrane protein with 24 transmembrane segments (S1-6) grouped into 4 homologous transmembrane domains (I-IV) (Fig. 3) [95, 96]. In addition, the subunit has 3 intracellular loops, each linking S6 and S1 of 2 transmembrane domains. Interestingly, loop I-II has the α$_1$-interaction domain (AID), the site where the β subunit binds Ca$_V$α$_1$ for trafficking. Furthermore, the SNARE proteins syntaxin 1A, synaptosomal-associated protein 25 (SNAP-25), and synaptotagmin 1 (SYT1) associate with the α$_1$ subunit at the II-III loop, connecting it to the insulin granules [97]. Intriguingly, the α$_1$ subunit pore consists of 4 membrane-embedded pore loops (P-loops), each containing a glutamic acid residue that is responsible for Ca$^{2+}$ entry [98]. During depolarization, voltage sensors such as cationic arginine or lysine residues at S4 cause a conformational change resulting in opening of the α$_1$ pore [90]. Then, upon binding of Ca$^{2+}$ to the extracellular end of the α$_1$ pore, a Ca$^{2+}$ bound to the intracellular end is repelled into the cytosol and replaced with the new extracellular Ca$^{2+}$ [99].

In recent years, regions at the N- and C-termini of the α$_1$ subunit have been demonstrated to mediate important functions. Two domains at the C-terminus, proximal and distal C-terminus regulatory domain (PCRD and DCRD), were shown to be involved in channel inactivation [100-102]. In addition, a fragment of the C-terminus was found to regulate Ca$_V$ channel transcription by translocating to the nucleus [103, 104]. The N-terminus is also involved in channel inactivation as it serves as a CaM-binding site in Ca$_V$1.2 and Ca$_V$1.3 channels [102, 105].
Figure 3 The $\alpha_1$ subunit structure
An illustration of the Ca$\text{v}_\alpha_1$ structure in the plasma membrane.

Regulation

Studies on rabbit and rat identified phosphorylation and glycosylation sites in Ca$\text{v}_\alpha_1$, specifically Ca$\text{v}_1.1$, Ca$\text{v}_1.2$, and Ca$\text{v}_2.1$ [96, 106-111]. Phosphorylation was carried out by cAMP-activated PKA and PKC, suggesting an indirect cAMP regulation of Ca$\text{v}_\alpha_1$. Although no glycosylation was observed in Ca$\text{v}_1.1$ and Ca$\text{v}_1.2$, a short form of Ca$\text{v}_2.1$ (95 kDa) was found to be glycosylated. Another Ca$\text{v}$ channel regulatory factor is Ca$^2+$. It achieves channel inactivation by forming a complex with CaM which binds to the C-terminus of the Ca$\text{v}$ channel and regulates channel gating.

Activation and inactivation

Gating is an important property of the Ca$\text{v}$ channel that is regulated by processes of activation and inactivation. Activation occurs when the membrane depolarizes, resulting in opening of the channel. Inactivation, on the other hand, is when the Ca$\text{v}$ channel becomes less permeable. There are two types of inactivation, Ca$^{2+}$-dependent inactivation (CDI) and voltage-dependent inactivation (VDI) [95, 102].

CDI takes place when Ca$^{2+}$ binds channel-tethered CaM, causing it to undergo a conformational change and channel inactivation [89]. The C-terminus-bound Ca$^{2+}$/CaM complex changes structure to bind the N-terminus of the channel, thus blocking Ca$^{2+}$ entry. Therefore, an increase in Ca$^{2+}$ influx increases CDI, reaching ~65% upon full channel activation [112]. The degree of CDI, however, varies depending on the type of Ca$\text{v}$ channel. L-type channels have strong CDI whereas R-type channels have weak CDI.

The other type of inactivation, VDI, depends on the difference in charge across the membrane which is determined by ions like K$^+$, Na$^+$, Cl$^-$, and Ca$^{2+}$.
β subunit

Structure
The human β subunit is encoded by 4 different genes (CACNB1-4) that are translated by free ribosomes [96]. Crystal structures of the subunit identified 5 domains, 2 highly conserved and 3 highly variable (Fig. 4) [113-117]. The highly conserved Src-homology (SH3) and guanylate kinase (GK) domains play an important role in CaVα1 trafficking. In yeast, the GK domain has an active catalytic site [118]. This site is replaced with a hydrophobic AID binding pocket (ABP; also referred to as β-interaction domain or BID) in mammalian GK domain [114-116]. The AID-ABP interaction positions the β subunit near the intracellular end of CaVα1 pore. This positioning of the β subunit allows it to regulate channel inactivation, since the AID N-terminus is very close to the IS6 segment of CaVα1 [113]. The SH3 domain is required for protein-protein interaction. To achieve this, a β sheet blocked by the HOOK domain is exposed via a conformational change in the SH3 domain. Interestingly, studies on Xenopus oocytes show that both SH3 and GK domains interact intramolecularly and disruption of such connection hinders CaVβ from CaVα1 trafficking and gating [119].

![Figure 4 The β subunit domains](image)

An illustration of CaVβ structural domains.

Alternative splicing
Alternative splicing is found in all human CaV subunits, including the β subunit. Each of the 4 β subunit genes (CACNB1-4) exhibits at least 2 alternative splicings, thus increasing subunit variation. Splicing takes place at the exons of the highly variable N-terminus, C-terminus, and HOOK domain. Interestingly, these splice variants differ in distribution. While β2b in rats is found in the brain, heart, and aorta, β2d is explicitly expressed in the heart [120, 121]. Splice variants also vary in expression during development. The expression of β1b in rat brain increases 3-fold during development while that of β2c decreases [120-122].

Alternative splicing gives rise to CaVβ variants that exert unique functions independent of VGCC. The chicken β4c, for instance, was found to lack 90% of the
AID-harboring GK domain and the whole C-terminus [123]. This permits β4c to interact with the nuclear protein heterochromatin protein 1 (HP1) and localize to the nucleus, suggesting its involvement in transcriptional regulation. In fact, in vitro studies in Xenopus oocytes have shown that full-length β subunits are capable of interacting with Pax6(S), a transcription factor required for the development of the eye and nervous system [124]. The interaction translocates the β3 subunit from the cytoplasm to the nucleus and reduces Pax6(S) activity without affecting VGCC properties. Furthermore, β4a forms a complex with B56δ, a nuclear regulatory subunit of phosphatase 2A (PP2A), and translocates to the nucleus to regulate histone dephosphorylation [113, 125].

Localization and function
In the absence of the α1 subunit, most β subunits, except for β2a and β2e, localize to the cytosol [126, 127]. Although the reason behind β2c localization is unknown, localization of β2a is due to it being palmitoylated and anchored to the plasma membrane [128, 129]. The main role of β subunits in VGCC is trafficking, regulating, and increasing surface expression of CaV channels. Regulating the CaV channel involves influencing its activation and inactivation state. Inactivation of the CaV channel is enhanced by the variable HOOK region in the β subunit [114-116]. Because each β subunit has a different HOOK domain, they differ in degree of inactivation of CaV channels.

In contrast to the general role of β subunits, the β3 subunit surprisingly acts as a brake on insulin secretion [130]. A study showed that β cells from β3 knockout mice had enhanced Ca2+ oscillations and improved insulin exocytosis. In addition, these β cells had elevated intracellular Ca2+ due to increased release from intracellular stores via enhanced IP3 formation.

Palmitoylation of β2a
Palmitate is a 16-carbon fully saturated fatty acid that, although deleterious to the β cell when elevated, is important post-translationally. Palmitoylation is a post-translational modification involving the addition of a palmitoyl group onto a protein. Of all the β subunits, β2a is unique in that it contains two cysteine groups in its N-terminal region which can undergo palmitoylation. In general, palmitoylation is a post-translational modification where the fatty acid palmitate attaches to one or more accessible cysteine residues in a protein structure. There are 3 types of palmitoylation: S-, N-, and O-palmitoylation [131, 132]. In S-palmitoylation, palmitate links to cysteine in a reversible manner with the help of palmitoyl acyl transferases (PATs) [133]. N- and O-palmitoylations form amide and oxyster linkages to N-terminus cysteine and serine residues, respectively [132, 134]. Unlike N-palmitoylation, O-palmitoylation involves a monounsaturated palmitate (palmitoleic acid) and is believed to be reversible [134-137]. Depalmitoylation
involves the removal of palmitate from a protein in a reaction catalyzed by thioesterases [138].

It has been reported that at least 10% of human proteins are subjected to palmitoylation [139]. These proteins are involved in signaling, transcription, and in forming ion channels and receptors [140]. One of the important functions of palmitoylation comes from its hydrophobicity, allowing proteins to dock on the inner leaflet of the phospholipid bilayer. As an example, due to the switch between palmitoylation and depalmitoylation, two small GTPases, NRas and HRas, are capable of alternating between the Golgi membrane and the plasma membrane [140-142]. Additional functions of protein palmitoylation include membrane raft-targeting, protein conformational change, and protein-protein interaction [140].

\(\alpha_2\delta\) subunit

Structure

Like the \(\beta\) subunit, the \(\alpha_2\delta\) auxiliary subunit (\(\sim 175\) kDa) is also encoded by 4 different genes (\(CACNA2D1-4\)) in humans, but unlike the \(\beta\) subunit, they are translated by ER ribosomes [84, 96, 143]. At first, the structure of \(\alpha_2\delta\) was determined biochemically and was thought to consist of two different proteins linked by a disulfide bond [144]. However, upon cloning of \(CACNA2D\), it became clear that both proteins emerged from the same gene. The \(\alpha_2\delta\) subunit is synthesized as a continuous polypeptide chain. During processing in the ER and Golgi apparatus, it acquires a disulfide bond between the \(\alpha_2\) and the \(\delta\) parts and undergoes glycosylation at several amino acid residues (Fig. 5) [145]. However, post-translational cleavage by proteases splits the protein into \(\alpha_2\) and \(\delta\), keeping them connected via the disulfide bond [146]. The \(\alpha_2\delta\) subunit consists of 5 domains: N-terminus, C-terminus, von Willebrand factor A (VWA), and 2 chemosensory-like domains (CSDs; or Cache domain) (Fig. 5) [147, 148].
The N-terminus has a signal sequence that guides the newly synthesized $\alpha_2\delta$ into the ER lumen [149]. The C-terminus is hydrophobic and is thought to be a transmembrane domain, although a predicted small sequence of this domain is in the intracellular environment [150, 151]. Proteomic prediction analysis shows that the $\alpha_2\delta$ subunit can be anchored to membrane rafts by glycosylphosphatidylinositol (GPI) which was also confirmed by many biochemical studies [152, 153]. Interestingly, the VWA domain, with the help of its metal ion-dependent adhesion site (MIDAS) motif, is involved in protein-protein interaction with extracellular matrix proteins and cell-adhesion proteins [154]. The MIDAS motif binds a divalent cation such as Ca$^{2+}$ or Mg$^{2+}$ causing a structural change in the subunit and allowing it to interact with other proteins. Although the VWA is an $\alpha_2\delta$ subunit domain, it has been found in other proteins that require protein-protein interaction, for example in integrins. Lastly, the CSDs were also discovered in bacteria and serve as multiple nutrient sensors [147].

**Localization**

The $\alpha_2\delta$ subunit, similar to the $\beta$ subunit, is expressed in excitable tissues like skeletal and cardiac muscles, brain, endocrine tissue, and retina [155, 156]. In human and mouse pancreatic islets, the predominant CACNA2D is the CACNA2D1 [157-159].
Table 3 Ca\textsubscript{V}α\textsubscript{2}δ tissue distribution

Tissue distribution of α\textsubscript{2}δ subunit.

<table>
<thead>
<tr>
<th>Subunit</th>
<th>Human</th>
<th>Mouse</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mRNA</td>
<td>Protein</td>
</tr>
<tr>
<td>α\textsubscript{2}δ\textsubscript{1}</td>
<td>Brain</td>
<td>Heart</td>
</tr>
<tr>
<td></td>
<td>Heart</td>
<td>Kidney</td>
</tr>
<tr>
<td></td>
<td>Pancreas</td>
<td>Spleen</td>
</tr>
<tr>
<td></td>
<td>Testis</td>
<td>Testis</td>
</tr>
<tr>
<td>α\textsubscript{2}δ\textsubscript{2}</td>
<td>Brain</td>
<td>Heart</td>
</tr>
<tr>
<td></td>
<td>Heart</td>
<td>Kidney</td>
</tr>
<tr>
<td></td>
<td>Pancreas</td>
<td>Lung</td>
</tr>
<tr>
<td>α\textsubscript{2}δ\textsubscript{3}</td>
<td>Brain</td>
<td>Heart</td>
</tr>
<tr>
<td></td>
<td>Heart</td>
<td>Testis</td>
</tr>
<tr>
<td></td>
<td>Skeletal muscle</td>
<td>Kidney</td>
</tr>
<tr>
<td>α\textsubscript{2}δ\textsubscript{4}</td>
<td>Brain</td>
<td>Heart</td>
</tr>
<tr>
<td></td>
<td>Heart</td>
<td>Testis</td>
</tr>
<tr>
<td></td>
<td>Skeletal muscle</td>
<td>Small intestine</td>
</tr>
<tr>
<td></td>
<td>Liver</td>
<td>Liver</td>
</tr>
<tr>
<td></td>
<td>Adrenal gland</td>
<td>Adrenal gland</td>
</tr>
<tr>
<td></td>
<td>Pituitary gland</td>
<td>Pituitary gland</td>
</tr>
<tr>
<td></td>
<td>Liver</td>
<td>Retina</td>
</tr>
<tr>
<td></td>
<td>Lung</td>
<td>Retina</td>
</tr>
</tbody>
</table>

Function

The study of the auxiliary subunit α\textsubscript{2}δ in pancreatic islets is minimal compared to in neurons and muscles. In general, the subunit facilitates Ca\textsubscript{V} channel surface expression and turnover, decreases the opening time of the α\textsubscript{1} subunit (i.e. increases inactivation), and increases Ca\textsubscript{V} current with the aid of the MIDAS motif [160-165]. Knockout of α\textsubscript{2}δ\textsubscript{1} subunit in mice, via reduction of L-, P/Q-, N-, & R-type Ca\textsubscript{V} channel currents, hindered the first and second phase insulin secretion, as well as decreased β cell mass [166]. The effect was sex-dependent as male mice developed diabetes while female mice solely had a higher risk of the disease [167]. This was due to the increase in basal insulin release in female mice, which had a preventive effect on diabetes development. A recent study showed that trafficking and activity of neuronal Ca\textsubscript{V}2.2 appeared to be dependent on the post-translational cleavage of the α\textsubscript{2}δ\textsubscript{1} subunit [168]. Surprisingly, this modification is unnecessary for α\textsubscript{2}δ\textsubscript{1} transport to the plasma membrane.

Additionally, because of the α\textsubscript{2}δ subunit 3 arginine (RRR) motif near the VWA domain (Fig. 5), the α\textsubscript{2}δ subunit is capable of interacting with gabapentin, an anti-epileptic drug [169, 170]. In vivo studies on rat brain neurons showed that the binding of gabapentin to the α\textsubscript{2}δ\textsubscript{1} subunit lowered Ca\textsuperscript{2+} currents. The α\textsubscript{2}δ subunit can also block and enhance the actions of ω-conotoxins (painkillers) and DHP
antagonists, respectively [165, 171]. In case of ω-conotoxins, the role of the subunit, specifically the α2 part, is believed to involve blocking the drug binding site on the CaV channel [171, 172]. As for the DHP antagonists, the α2δ subunit increases channel inactivation and since DHP antagonists generally prefer binding to inactivated channels, the affinity of antagonist binding to channels associated with α2δ subunit increases [173].
Aims

CaV channel auxiliary subunits are an important element in trafficking and regulating the main α1 subunit. However, their role in pancreatic β cells remains unclear. This thesis investigates the roles of β1, β2a, and α2δ1 subunits in pancreatic β cells. The thesis also addresses the role of membrane rafts in CaV channel function.

The specific aims are as follows:

1. To explore the role of membrane rafts in regulating voltage-gated Ca²⁺ channel (CaV) activity and insulin release in β cells.

2. To examine the role of Tcf7l2 in regulating the expression of α2δ1 subunit and the subsequent effect on CaV channel trafficking and activity in β cells.

3. To investigate the role of the β1 subunit in regulating CaV channel activity and the resulting effect on insulin secretion and β cell survival.

4. To study the role of palmitoylation of the β2a subunit in regulating CaV channel activity and the ensuing effect on insulin secretion and β cell survival.
Materials and Methods

RNAseq and MicroArray

Microarray
Microarray technique is used to detect gene expressions among a library of transcripts. This method of gene expression analysis has the advantage of being quick, robust, and reliable. It is currently cheaper than RNA sequencing (RNAseq). However, it requires prior knowledge of the desired transcript and, thus, it is not ideal for finding novel genes, structural variations, or isoforms. The data produced by a microarray method only indicates relative expression and as such should not be used for quantification purposes.

RNAseq
RNAseq technique is also used to detect gene expressions, however, it extracts data from the transcriptome pool and thus does not prior knowledge of a sequence. This is of particular importance as it allows discovering novel genes, isoforms, structural variations, or transcripts. This relatively new method has the advantage of being highly sensitive compared to microarray, however, it has a higher cost. It also has the advantage of providing absolute quantifications instead of relative expressions. This, however, demands more time for data analysis and larger storage space. Because of it being relatively new, there is no standard RNAseq protocol, and therefore, data are harder to compare.

Protein Quantification

Immunoblotting and immunohistochemistry methods use antibodies specific to a desired protein to semi-quantify and visualize the protein, respectively.
Immunoblot

Immunoblot (or western blot) is composed of 3 stages: running, transfer, and detection.

Running
Briefly, this stage involves loading an amount of protein onto a sodium dodecyl sulfate-polyacrylamide gel to perform electrophoresis (SDS-PAGE) after being denatured with DTT and heat. The denaturing step is important for migration of the protein as it breaks the sulfide bonds from secondary and tertiary structures, making the protein linear. The proteins will migrate, due to their negative charge, from the cathode (negative) to the anode (positive) ends (top of gel to bottom) with the help of ions in the running buffer. Depending on the molecular weight of the protein, it will migrate at a certain speed on the gel. This is because SDS-PAGE gels contain a percentage of polyacrylamide which gives the gel structure a certain sized ‘holes’ through which proteins migrate. The more acrylamide, the more rigid the gel is and the smaller these ‘holes’ are. This means that only smaller sized proteins will reach faster towards the bottom of the gel while the larger ones will get held back. Using a gradient gel, such as 4-15%, is often useful to capture different sized proteins.

Transfer
The gel is then placed, along with a polyvinylidene difluoride (PVDF) membrane, in a sandwich cassette such that the gel is closer to the cathode while the PVDF membrane is closer to the anode. This is to ensure the transfer of proteins from the gel onto the PVDF membrane i.e. from negative end to the positive end. Current is then applied to the sandwich cassette submerged in Tris-based transfer buffer, allowing the proteins to migrate onto the membrane. This is called wet transfer as opposed to dry or semi-dry transfers where the setup is slightly changed. One of the advantages of a PVDF membrane over a nitrocellulose membrane is the ability to prevent proteins from passing through the membrane (overtransfer). This is especially useful if studying two or more proteins with vastly different sizes as smaller proteins transfer at a faster rate than larger ones. The transfer stage is vulnerable because a mere tiny bubble can render the membrane useless.

Detection
After transfer, the membrane is incubated with a blocking agent containing around 5% protein such as skimmed milk or bovine serum albumin (BSA). The 5% protein in milk binds non-specifically to spaces on the PVDF membrane that are left unbound. This reduces background noise during detection as these spaces, if unblocked, may bind to antibodies during incubation. Next, the membrane is incubated first with primary and then with secondary antibodies that will
specifically bind to the desired protein. After that, the protein bands are visualized under ultra violet light using enhanced chemiluminescence (ECL) and analyzed using appropriate softwares.

**Immunohistochemistry**

Immunohistochemistry uses a similar concept to immunoblot. The Detection stage is similar while Running and Transfer are replaced with Fixation. Although this method is not considered quantifiable, it is useful for detecting protein localization.

*Fixation*

Cells are fixed with 4% paraformaldehyde (PFA) and permeabilized with a detergent such as saponin. Permeabilization of the plasma membrane is required for antibody entry and binding of intracellular proteins.

The following stage will be similar to immunoblot Detection stage where a blocking agent, primary and secondary antibodies are used. The fixed cells are then visualized under a confocal microscope.

**Ca\(^{2+}\) Quantification**

*Ratiometric vs non-ratiometric*

Ca\(^{2+}\) is an essential contributor to GSIS and thus quantifying it is of great interest. Ca\(^{2+}\) quantification is achieved with two main methods: ratiometric and non-ratiometric.

*Ratiometric*

This technique measures free intracellular Ca\(^{2+}\) ions using a ratiometric fluorescent dye called aminopolycarboxylic acid or Fura-2. The fluorescence from the dye can be used to quantify Ca\(^{2+}\), since upon excitation at 340 nm and 380 nm, the ratio of emission at these wavelengths is directly proportional to the amount of free Ca\(^{2+}\) bound to Fura-2. The advantage of this ratiometric technique is the elimination of confounding factors such as dye concentration, bleaching, change in focus, variations in laser intensity, and cell thickness. However, this technique is more difficult in measuring and processing data as it requires specific settings that are only available with some microscopes.
Non-ratiometric

This technique is used to detect free intracellular Ca$^{2+}$ in a non-quantifiable way using a fluorophore such as Fluo-5F. The excitation and emission are at 494 nm and 516 nm wavelengths, respectively. The fluorescence intensity may reflect the amount of free intracellular Ca$^{2+}$. However, the fluorescence can be influenced by other factors such as change in focus, variations in laser intensity, and dye concentration.

Knockdown and Overexpression

One of the most common methods to determine protein function is to attempt to either eliminate the protein or over-produce it. This is attained by knockdown and overexpression techniques.

Knockdown

The protein expression can be reduced (knocked down) by up to 90-95% but not completely eliminated with standard cell manipulation techniques (a complete elimination will be termed knockout and is achieved by DNA-editing techniques). Protein knockdown takes place at the mRNA level where a small interference RNA (siRNA) targets the desired mRNA and activates the RNA-induced silencing complex (RISC) machinery to break down the mRNA, preventing it from being translated. The results are compared to the control siRNA which does not correspond to any known RNA sequence and theoretically should pose no change to cellular physiology.

Overexpression

With this technique, the protein is overexpressed using a plasmid. This plasmid can either carry a tag-attached-protein or the protein sequence alone. A popular tag used for detection is green fluorescence protein (GFP). The plasmid-encoded protein is expressed using the cell’s transcription and translation machineries.
Results and Discussion

Paper I

Results

Type 2 diabetic islets show decreased plasma membrane cholesterol content and membrane rafts

In type 2 diabetes (T2D), the aberrant blood lipid profile has been suggested to contribute to pancreatic β cell dysfunction. To investigate the importance of cholesterol-enriched membrane rafts in pancreatic islets, the sphingolipid dye ATTO-SM and the cholesterol dye filipin were used to stain healthy Wistar and diabetic GK rat islets (Fig. 1a-e in Paper I). GK rat islets showed a marked decrease in filipin and ATTO-SM stainings (~40% and ~60% respectively) compared to healthy islets. This suggests a depletion in cholesterol-enriched membrane rafts in rat islets under diabetic conditions (p < 0.1, n = 3 rats/condition; Fig. 1b & 1d in Paper I).

Are the membrane rafts expressed differently between α and β cells? To elucidate this, we stained dispersed Wistar islets with ATTO-SM and found that, in comparison with α cells, β cells showed approximately 300% higher membrane raft intensity (p < 0.5, n = 3 rats/group; Fig. 1e in Paper I).

To explore whether these results in rat translated to the human situation, human islets from donors with T2D were stained and found to display a similar membrane raft phenotype as that of GK rat islets. A significant decrease (~50%) in ATTO-SM staining was observed in islets from donors with T2D as oppose to healthy islets (p < 0.1, n = 3 donors/condition; Fig. 1f-g in Paper I). A reduction in membrane rafts was also observed in healthy human islets treated with the cholesterol-depleting enzyme cholesterol oxidase (CO), which resulted in ~40% reduced ATTO-SM intensity (p < 0.1, n = 3 donors/condition; Fig. 1f-g in Paper I). CO dosage and treatment time were optimized for activity and cell toxicity on INS-1 832/13 cells prior its use on islets (Suppl. Fig. 1 in Paper I).
**Disruption of membrane rafts causes increased basal insulin secretion**

The main functions of β cells are to sense changes in blood glucose and secrete insulin in order to maintain euglycemia, i.e. normal blood glucose. To evaluate the effect of CO on β cell function, we measured glucose stimulated insulin secretion (GSIS) in human and rodent islets as well as in INS-1 832/13 cells following CO treatment. For comparison, the effects of the widely used cholesterol-depleting agent MβCD were studied in parallel. First, optimal time was determined using a time-dependent experiment where INS-1 832/13 cells were treated with CO for 0.5, 1, and 2 hours. A 1 hour CO treatment time was selected as the enzyme influenced both basal (2.8 mM) and stimulated (16.7 mM) insulin secretions at that time point (Fig. 2a). Next, INS-1 832/13 cells were pretreated with either MβCD or CO and insulin release was measured. Under both conditions stimulated secretion was slightly increased. However, the main finding was that basal secretion was greatly elevated compared to the control condition (Fig. 2b in Paper I). As a result, the stimulatory effect of glucose, expressed as fold-change between 16.7 mM and 2.8 mM glucose, was decreased by half compared to control. In agreement with the cell line data, human and rat islets subjected to CO treatment showed elevation in basal insulin secretion (8-fold in human and 3-fold in rat) compared to control (Fig. 2c-d in Paper I). MβCD treatment, on the other hand, showed little effect on basal secretion in human and rat islets.

Since membrane-clustering of the SNARE protein syntaxin 1A facilitates insulin secretory granule exocytosis, the importance of membrane rafts for this process was assessed. Indeed, INS-1 832/13 cells treated with CO showed a scattered syntaxin 1A localization as opposed to its native membrane association. The ratio of syntaxin 1A membrane expression to the intracellular level was greatly reduced from 3.2 ± 0.8 to 1.0 ± 0.2 in CO-treated cells compared to control (Suppl. Fig. 2 in Paper I).

**Activation of [Ca\(^{2+}\)], oscillations in CO-treated cells under depolarizing and resting conditions**

After membrane depolarization, voltage-gated Ca\(^{2+}\) (Ca\(_V\)) channels trigger insulin release by allowing Ca\(^{2+}\) entry. This causes a rise in intracellular Ca\(^{2+}\) ([Ca\(^{2+}\)]) stimulating exocytosis. Due to the importance of this step, Ca\(^{2+}\) entry into INS-1 832/13 cells and depolarization-evoked increases in [Ca\(^{2+}\)], were tested with or without membrane raft dispersion. To examine the Ca\(^{2+}\) signaling, cells were incubated with the Ca\(^{2+}\) fluorophore Fluo-5F, and depolarized with 70 mM K\(^+\) to trigger Ca\(^{2+}\) influx and rise in [Ca\(^{2+}\)]. Compared to control cells, the cells treated with CO had higher [Ca\(^{2+}\)], after K\(^+\) stimulation (p < 0.001, n = 3; Fig. 3a-c in Paper I). When investigating Ca\(^{2+}\) oscillations under resting conditions, the Ca\(^{2+}\) spikes were 85% more frequent in CO-treated cells compared to control cells under resting condition (p < 0.5, n = 3; Fig. 3d-e in Paper I). To determine whether Ca\(^{2+}\) stores contribute to this rise, thapsigargin (TG) was used. TG is an inhibitor of the
sarcoendoplasmic reticulum Ca\(^{2+}\)-ATPase (SERCA) pump, thus preventing Ca\(^{2+}\) entry into the ER. The effect of CO on Ca\(^{2+}\) oscillations was also observed in resting TG-treated cells (Fig. 3d & 3f in Paper I).

To check whether the increased [Ca\(^{2+}\)]\(_i\) is due to altered localization, clustering, or expression of Cav1.2, INS-1 832/13 cells were transfected with EGFP-Cav1.2 and subjected, or not, to CO treatment. Interestingly, CO treatment caused disruption of Cav1.2 clustering on the plasma membrane, but neither localization nor expression were affected (p < 0.5, n = 3; Fig. 3g-i in Paper I).

**CO treatment increases Ca\(^{2+}\) influx via Cav channels**

Cav channel activity can either be measured as whole-cell or single-channel currents. The whole-cell current measures the activity of all Cav channels in the cell, whereas the single-channel current measures the activity of a single Cav channel. To assess the effect of CO on whole-cell Ca\(^{2+}\) currents, whole-cell patch-clamp recordings were performed in INS-1 832/13 cells. An increase in Ca\(^{2+}\) current was observed with membrane cholesterol oxidation (Fig. 4a-b in Paper I). Next, single-channel recordings were performed to determine whether the increase in whole-cell Ca\(^{2+}\) current is due to an increase in number or activity of single Cav channels. We confirmed that the latter suggested mechanism was indeed responsible for the CO-induced Ca\(^{2+}\) influx, as single-channel activity was upregulated when INS-1 832/13 cells were treated with CO (Fig. 4c in Paper I).

**A prolonged elevated glucose disperses membrane rafts**

Glucotoxicity is considered a major contributor to T2D pathogenesis. In order to observe the effect of glucose on membrane rafts, INS-1 832/13 cells were subjected to glucose treatment and stained with ATTO-SM. Like T2D islets, the rat β cell line displayed, in a time-dependent manner, a severe reduction in membrane rafts under glucotoxic conditions (Fig. 6 in Paper I).

**Discussion**

Pancreatic β cells secrete insulin following rapid Ca\(^{2+}\) influx by voltage-gated Ca\(^{2+}\) (Cav) channels. The composition of cholesterol-rich membrane rafts is essential for the function and localization of many proteins, including Cav channels. Sphingolipids are constituents of membrane rafts that are synthesized from palmitate and serine. Palmitate treatment has effects on insulin secretion, including increasing basal insulin release [174, 175]. However, palmitate has many effects in the β cell and the increase in basal secretion may be by other means than by disrupted membrane rafts.
Healthy human islets show normal membrane raft expression (Fig. 1). However, once the healthy islets are depleted of cholesterol, they exhibit dispersed membrane rafts. Could high blood glucose influence membrane raft integrity? Indeed, islets from donors with T2D display much lower membrane raft expression when compared to healthy islets (Fig. 1 in Paper I). A study showed that INS-1 cells exposed to high glucose displayed a reduction in membrane cholesterol content and dismantled membrane rafts, further supporting our findings [176].

To understand the role of these rafts in pancreatic β cells, we have disrupted the microdomains using the enzyme cholesterol oxidase (CO). This enzyme selectively oxidizes membrane cholesterol to 4-cholesten-3-one thus dispersing membrane rafts [177, 178]. Despite the numerous studies reporting that insulin secretion is affected by changes in membrane cholesterol content, they are inconsistent. One study has shown that MβCD treatment in rodents promoted insulin release [80]. On the contrary, another study has shown that MβCD treatment lowered insulin exocytosis in mouse β cells [82]. Furthermore, a report demonstrated reduced insulin secretion under conditions of excess cholesterol [179]. Interestingly, MβCD-mediated cholesterol depletion restored insulin secretion under that condition. A possible overarching explanation that would reconcile these divergent observations would be that the relation between cholesterol content and insulin secretion is bell-shaped. When cholesterol is at normal levels, β cells secrete insulin normally. If, however, cholesterol levels were increased or depleted, secretion will be compromised. To further clarify an area with such conflicting results, an alternative tool also used for manipulating membrane cholesterol composition, CO, was utilized. Unlike data from MβCD treatment, CO data are more consistent between human/rat islets and INS-1 832/13 cells.

Basal insulin secretion rises in human/rat islets and β cells when treated with CO (Fig. 2 in Paper I). A similar effect was observed in mouse islets after silencing caveolin-1 (Cav-1), a protein component of specialized rafts called caveolae. Dispersion of membrane rafts was discovered to also increase basal glucagon secretion in α cells. These data further emphasize the importance of membrane rafts in controlling basal insulin exocytosis in β cells. Moreover, exocytotic SNARE proteins require intact membrane rafts to function. We have shown that dispersing the rafts with CO indeed delocalizes the SNARE protein syntaxin 1A from the plasma membrane to the endosomes, at least in INS-1 832/13 cells (Suppl. Fig. 2 in Paper I).

The mechanism behind the effect of membrane raft dispersion on basal insulin secretion is only partially elucidated. We show that Ca²⁺ oscillations under both resting and stimulatory conditions are elevated in cells depleted of membrane cholesterol (Fig. 3d-f in Paper I). Consequently, depolarization-evoked intracellular Ca²⁺ levels are also increased (Fig. 3a-c in Paper I). The reason for this could be an
increase in either the number of active CaV channels or an increased activity in every single CaV channel. In fact, we found that the increase in Ca\(^{2+}\) influx is driven by the longer opening time of each single channel (Fig. 4c in Paper I). This is in line with a study showing that the gating of Kir channels and the current density of N-type CaV channels are affected by membrane cholesterol composition [180-182]. Membrane raft dispersion also promotes CaV1.2 declustering (Fig. 3g-h in Paper I). A way to better understand this is if one regards the SNARE protein complex as a gate-keeper for insulin exocytosis and as clustering agents for CaV channels. Hence, a rise in blood glucose could induce membrane raft dismantling leading to a reduced membrane expression of SNARE proteins. In turn, this would result in CaV channel declustering and elevated basal Ca\(^{2+}\) spikes leading to dysregulated basal insulin exocytosis. In support of this theory, a study in mouse MIN6B1 cells showed that a rise in basal insulin secretion was associated with an elevated cytosolic Ca\(^{2+}\) [183]. To conclude, the work presented in this paper has added important insight into the mechanism behind elevated basal insulin secretion upon membrane raft dispersion in pancreatic \(\beta\) cells and islets.

**Paper II**

**Results**

*TCF7L2 controls expression of Cacna2d1/\(\alpha_2\delta_1\)*

An intronic genomic region close to the *TCF7L2* gene has been identified as the strongest genetic risk factor for T2D. We have previously shown that this transcription factor regulates multiple genes including the CaV auxiliary subunit gene *Cacna2d1* [184]. To expand our understanding of this link, we investigated the effect of Tcf7l2 gene expression on Cacna2d1. Rat INS-1 832/13 cells and islets were transfected with siRNA targeting Tcf7l2. Indeed, both \(\alpha_2\delta_1\) gene expression and protein levels were downregulated in INS-1 832/13 cells and rat islets following Tcf7l2 silencing (Fig. 1 in Paper II).

*Silencing Cacna2d1 prevents trafficking of CaV1.2 to the plasma membrane*

The \(\alpha_2\delta_1\) subunit is suggested to be involved in CaV channel trafficking and regulation. To study the effect of silencing Cacna2d1 on CaV1.2 in insulin-secreting cells, we suppressed Cacna2d1 and quantified the gene expression and protein levels of CaV1.2. Interestingly, the reduced levels of \(\alpha_2\delta_1\) in rat \(\beta\) cells resulted in reduction of CaV1.2 surface expression by \(\sim50\%\) while the total amount of CaV1.2 remained unchanged (Fig. 2 in Paper II). Since one of the roles of \(\alpha_2\delta_1\) is CaV channel trafficking, CaV1.2 can be envisioned to be entrapped in the ER after protein
translation, and we therefore investigated the localization of Ca\textsubscript{V}1.2 in cells lacking \(\alpha_2\delta_1\) subunits. However, we found that Ca\textsubscript{V}1.2 was retained in recycling endosomes, suggesting that the \(\alpha_2\delta_1\) subunit mainly acts to translocate the channel from the recycling endosome back to cell surface.

Silencing Cacna2d1 affects Ca\textsuperscript{2+} influx and exocytosis
Regulating Ca\textsubscript{V} channel activity is crucial for normal \(\beta\) cell function. To test the effect of Cacna2d1 suppression on depolarization-evoked Ca\textsuperscript{2+} current, INS-1 832/13 cells with knocked-down Cacna2d1 gene expression were incubated with the Ca\textsuperscript{2+} fluorophore Fluo-5F and depolarized with 70 mM K\textsuperscript{+} to trigger Ca\textsuperscript{2+} influx and consequent rise in intracellular Ca\textsuperscript{2+} ([Ca\textsuperscript{2+}]\textsubscript{i}). Silencing Cacna2d1 or Tcf7l2 in clonal \(\beta\) cells lowered Ca\textsuperscript{2+} entry leading to a fall in [Ca\textsuperscript{2+}]. The Ca\textsuperscript{2+}-lowering effect was also observed in cells treated with the anti-epileptic drug gabapentin (Fig. 4A-D in Paper II); this drug is believed to inhibit \(\alpha_2\delta_1\) subunit function. To follow the Ca\textsuperscript{2+}-lowering effect further, we voltage-clamped mouse \(\beta\) cells from a -70 mV holding potential to membrane potentials between -50 and 10 mV. Cells that were either Cacna2d1-silenced or gabapentin-treated showed a significant reduction (~33%) in whole-cell Ca\textsuperscript{2+} current (Fig. 4E in Paper II). To explore the implication of this on \(\beta\) cell function, we tested insulin secretory function. Upon silencing Cacna2d1, INS-1 832/13 cells secreted less insulin under both resting and stimulated conditions (Fig. 4F in Paper II).

Discussion
TCF7L2 is a transcription factor involved in the Wnt/\(\beta\)-catenin signaling pathway and regulates several genes involved in pancreas development, including ISL1, MAFA, and PDX1. It has been reported that TCF7L2 affects insulin secretion in rodent and human pancreatic islets [184]. Moreover, silencing Tcf7l2 reduces \(\beta\) cell survival and impairs GSIS [185]. We have recently shown that the Cacna2d1 (\(\alpha_2\delta_1\)) gene is also regulated by TCF7L2 [184]. The auxiliary subunit \(\alpha_2\delta\) is involved in voltage-gated calcium (Ca\textsubscript{V}) channel trafficking and gating. For this reason, we hypothesize that \(\alpha_2\delta_1\) subunit may contribute to the proper execution of GSIS in pancreatic \(\beta\) cells.

In this paper, we show that \(\alpha_2\delta_1\) gene expression and protein levels are downregulated upon Tcf7l2 silencing (Fig. 1 in Paper II). Since the \(\alpha_2\delta\) subunit aids in Ca\textsubscript{V} channel trafficking, we explored the effect of \(\alpha_2\delta_1\) subunit removal on Ca\textsubscript{V}1.2 localization. Knocking down Cacna2d1 in INS-1 832/13 and mouse \(\beta\) cells resulted in decreased Ca\textsubscript{V}1.2 plasma membrane expression caused by retention of the channels in recycling endosomes (Fig. 2C-E & 3 in Paper II). Our data is in line with work presented in HEK293 cells using a co-expression system [186].
The $\alpha_2\delta$ subunit carries a three arginine (RRR) motif at the von Willebrand factor A (VWA) domain allowing it to bind to the anti-epileptic drug gabapentin. The drug structurally resembles the neurotransmitter GABA but does not bind to GABA receptors. Instead, chronic exposure to gabapentin resulted in reduced surface expression of $\alpha_2\delta_1$ and $\alpha_1$ subunits leading to lower Ca$^{2+}$ current in mouse neurons [169, 187]. In consonance with this, our data confirmed the inhibitory effect of gabapentin on Ca$^{2+}$ activity and consequently reduced intracellular Ca$^{2+}$ in rodent pancreatic $\beta$ cells (Fig. 4 in Paper II). The degree of inhibition on Ca$^{2+}$ influx due to gabapentin was similar to that observed in the $\alpha_2\delta_1$ knockdown group (Fig. 4E in Paper II).

The consequence of $\alpha_2\delta_1$ gene expression silencing is a significant reduction in insulin secretion (Fig.4F in Paper II). However, gabapentin treatment had no influence on secretion (data not shown). The reason behind the failure of gabapentin to affect secretion remains unclear. A possible explanation could be the involvement of gabapentin in pathways such as GABA production, that may compensate for the decrease in insulin secretion driven by the loss of $\alpha_2\delta_1$ subunit. GABA may have an autocrine effect on pancreatic $\beta$ cells by binding to GABA$\_A$ receptor which in turn promotes insulin release [188-191]. Seemingly contrasting this notion, we observed a decrease in Ca$^{2+}$ currents when cells were treated with gabapentin. This, however, is due to the nature of the patch-clamp technique where the membrane potential of the cell is controlled by the experimenter, preventing any glucose- and K$^+$-stimulated depolarizations from taking place. In the case of the insulin secretion experiments, however, the membrane potential of the cell was not voltage-clamped. This means that any effect involving increased production of GABA and an autocrine action via GABA$\_A$ receptors should become manifest. In favor of this view, gabapentin was reported to induce severe hypoglycemia in six patients (diabetic and non-diabetic), suggesting it having insulinotropic effects [193]. In conclusion, TCF7L2 is important for maintaining proper $\beta$ cell function as it regulates $\alpha_2\delta_1$ subunit, and in turn influences Ca$_V$ channel trafficking and activity.
Paper III

Results

Gene expression of Cavβ1 in rat islets
There are four auxiliary β subunits in the human genome. The expression of these β subunits may vary depending on the tissue being explored. Gene expression analysis confirmed the expression of all β subunits in human islets (Fig. 1B in Paper III). In order to understand the role of the β1 subunit in pancreatic β cells, we examined the expression level of Cacnb1 in Wistar and GK rat islets. Islets from the diabetic model showed more than 40% reduction in Cacnb1 expression (Fig. 1C in Paper III). The results were reproduced by Western blot when rat-derived INS-1 832/13 cells and rat islets were treated with 20 mM glucose or 1 mM palmitate, respectively (Fig. 1D-E in Paper III).

Insulin secretion is reduced in beta cells with reduced Cavβ1 expression
To better understand whether the expression of the β1 subunit in rat INS-1 832/13 cells influences insulin secretion, we tested the effect of β1 suppression on β cell secretory function. GSIS was markedly reduced in cells lacking β1 subunits (Fig. 2 in Paper III).

Cavβ1 is required for maintaining healthy intracellular Ca^{2+} levels
The β subunits have been suggested to regulate Cav channel activity, as well as their translocation to the plasma membrane. To find the reason for the disrupted insulin secretion upon β1 silencing, we measured intracellular Ca^{2+} levels by Ca^{2+} imaging by using the Ca^{2+} fluorophore Fluo-5F. Cells were incubated with Fluo-5F and stimulated with 70 mM K^+ to allow observation of depolarization-evoked intracellular Ca^{2+} elevations by confocal microfluorimetry. We also measured whole-cell Ca^{2+} currents using the patch-clamp technique. Interestingly, Ca^{2+} influx was significantly lowered in the absence of β1 subunit (Fig. 3C in Paper III). This also resulted in a drastic reduction in K^+-stimulated intracellular Ca^{2+} levels by ~66% in rat β cells (Fig. 3A-B in Paper III). Because β cell function was compromised, β cell survival in Ca^{2+}-depleted environments was explored. When silencing Cacnb1, β cells suffered apoptosis which suggests an important role of the β1 subunit in β cell survival (Fig. 4 in Paper III).
Discussion

The β auxiliary subunit is believed to regulate CaV channel trafficking and gating in excitable tissues. However, the mechanism behind this remains unclear. Our gene expression data have shown that CACNB1 encoding the β₁ subunit is abundant in human pancreatic islets (Fig. 1A-B in Paper III). The expression of Cacnb1 in GK rat compared to Wistar rat, however, was greatly reduced (Fig. 1C in Paper III). This decrease in gene expression was also observed at the protein level in INS-1 832/13 cells and rat islets following chronic exposure to high glucose and palmitate, respectively (Fig. 1D-E in Paper III). Taken together, these results indicate an intriguing role of the β₁ subunit as a target of detrimental environmental factors that may lead to T2D.

Insulin secretion and β cell survival were both explored after silencing the β₁ subunit. Indeed, the data show a reduction in insulin secretion by ~75% in Cacnb1-silenced cells while an increase in apoptosis was observed (Fig. 2 and 4 in Paper III). False elevation of insulin secretion could take place under apoptotic conditions due to cell rupture and leakage. Normally, this is evident as strongly increased basal insulin secretion and loss of regulated insulin release. However, to further minimize the risk of this confounding factor, an additional washing step was added and then insulin secretion was normalized to total protein. Further investigation was made to identify the reason for β cell dysfunction and toxicity. Intriguingly, the patch-clamp and Ca²⁺-imaging experiments demonstrate that β cells lacking the β₁ subunit had significantly lower Ca²⁺ influx and depolarization-evoked cytosolic Ca²⁺ levels (Fig. 3 in Paper III). This argues against Ca²⁺-induced cell toxicity, but underscores that Ca²⁺ homeostasis is essential for β cell function and survival. It can be speculated that under diabetic conditions such as hyperglycemia, the β₁ subunit expression decreases in β cells, leading to intracellular Ca²⁺ starvation. In an attempt to restore Ca²⁺ homeostasis, Ca²⁺ from the ER and mitochondria are released into the cytosol. However, in doing so, these compartments deplete their Ca²⁺ stores and trigger stress elements causing apoptosis [194]. In support of this view, depletion of the ER Ca²⁺ stores was shown to trigger ER stress and programmed cell death [195].

Paper IV

Results

Gene expression of Caβ_{2a} in human and rat islets

Gene expression data showed that among the β₂ isoforms expressed, the levels of the β₂ₐ isoform were the highest in human pancreatic islets (Fig. 1A & C in Paper
IV). A comparison in expression levels of β2a between healthy Wistar and diabetic GK rat islets was performed. Since diabetic islets showed reduced β1 expression, we speculated that the β2a subunit expression would reveal a similar outcome. In support of our hypothesis, GK rat islets proved to express much less Cacnb2a compared to Wistar rat islets (Fig. 1B in Paper IV). To examine the effect of high glucose on β2a expression, we performed Western blot on INS-1 832/13 cells following chronic glucose exposure, and could conclude that chronic glucose exposure leads to a decrease in β2 subunit levels (Fig. 2C in Paper IV).

**Palmitoylation is required for CaVβ2a tethering to the plasma membrane**

The β2a subunit is post-translationally modified by palmitoylation. We tested whether the state of palmitoylation is essential for β2a function. INS-1 832/13 cells were made to overexpress either wildtype palmitoylatable (β2a; WT) or non-palmitoylatable (β2aCAS; mutant), GFP-tagged β2a subunit. Whereas the WT β2a was inserted in the plasma membrane, non-palmitoylatable β2aCAS was largely localized to the cytosol (Fig. 2A in Paper IV). We then followed this aberrant localization by investigating the cellular localization of the CaV channel α1C subunit (CaV1.2). To this end, we used COS-1 cells, which lack endogenous CaV channels. The CaV-free COS-1 cells were co-transfected with both α1C and β2a or β2aCAS GFP-tagged plasmids. The CaV1.2 channel was incapable of tethering to the plasma membrane in COS-1 cells expressing the mutant β2a (Fig. 2B in Paper IV). This demonstrates the importance of β2a palmitoylation in regulating CaV channel trafficking to the plasma membrane. We further investigated the effect of palmitoylation state of β2a on insulin secretion and β cell death. INS-1 832/13 cells overexpressing WT β2a secreted normal amounts of insulin with respect to glucose. However, overexpressing WT β2a also induced apoptosis, an effect not observed using the β2aCAS isoform (Fig. 3E-F and 4F-G in Paper IV).

**Overexpression of CaVβ2a upregulates basal intracellular Ca^{2+} concentration**

Since β2a palmitoylation state affected CaV1.2 localization, we further investigated the effect of β2a palmitoylation state on intracellular Ca^{2+} levels. Cells were incubated with Fluo-5F and stimulated with 70 mM K+ to induce membrane depolarization and Ca^{2+} influx. Using confocal microscopy, we observed a more than 100% increase in basal intracellular Ca^{2+} concentration in cells overexpressing WT palmitoylatable β2a, but not the mutant (Fig. 3A-D in Paper IV). The high K+-stimulated [Ca^{2+}], reveals an important role for β2a subunit in Ca^{2+} homeostasis and β cell survival.
Discussion

T2D is a metabolic disease characterized by high levels of glucose, cholesterol, TG, and palmitate in the blood. Palmitate is a fully saturated free fatty acid (FFA) that has many biological functions such as being a precursor for sphingolipid synthesis and necessary for palmitoylation of proteins. Furthermore, this study demonstrates the capacity of WT palmitoylated β2a, but not the mutant, for proper trafficking of both β2a and pore-forming α1C CaV channel subunits to the plasma membrane (Fig. 2A-B in Paper IV). Interestingly, excess of palmitoylated β2a also elevates cytoplasmic Ca2+ and induces cellular apoptosis (Fig. 3 in Paper IV). This rise in intracellular Ca2+, however, did not stimulate GSIS (4F-G in Paper IV). A possible explanation for the paradoxical absence of an effect of β2a overexpression on insulin secretion could be due to a downregulated expression of other β subunit isoforms that are required for secretion. This decreased expression could, therefore, counteract the potential increase in insulin release. However, excess membrane expression of palmitoylated β2a upregulates intracellular Ca2+ levels, leading to apparent Ca2+ overload and toxicity (Fig. 3 in Paper IV). Therefore, the possibility of a counteracting effect on insulin secretion by other β subunits cannot explain this secretion-ineffective rise in cytosolic Ca2+.

Another possible explanation for the failure of β2a-mediated increases in intracellular Ca2+-signals to stimulate insulin secretion is a β2a interaction with dynamin. Dynamin is suggested to be important for normal endocytosis and exocytosis, and thereby insulin secretion [196]. It has been shown that the conserved SH3 domain in β subunits could bind to dynamin [197]. It could be proposed that palmitoylated β2a interacts with dynamin leading to either a partial inhibitory effect on insulin exocytosis or a ‘kiss-and-run’ where the insulin granule alternates between exo- and endocytosis without fully releasing its insulin content.

Interestingly, it has been reported that ER stressors such as cytokines and thapsigargin could increase cytosolic Ca2+ while having minimal or no effect on insulin secretion [198]. On the other hand, metabolic stressors such as FFAs and high glucose are able to increase both intracellular Ca2+ and insulin secretion. One could speculate that excess palmitoylated β2a could trigger ER stressors via a mechanism involving the palmitoylated state of β2a since β2aΔS had no effect on cytosolic Ca2+. What we can conclude with certainty is that the palmitoylation state of β2a can influence intracellular Ca2+ levels and excess of the palmitoylated subunit induces β cell death.
Conclusion

1. The work presented in Paper I has partly explained the mechanism behind elevated basal insulin secretion upon membrane raft dispersion in pancreatic β cells, a phenomenon that appears relevant for the increased basal insulin secretion observed in T2D. Membrane rafts are required for proper clustering of Cav1.2 and surface expression of the SNARE protein syntaxin 1A. Declustering of Cav1.2 is suggested to lead to dysregulated basal Ca\(^{2+}\) oscillations and increased single-channel activity and hence a rise in basal insulin release.

2. The work presented in Paper II revealed the importance of TCF7L2 in maintaining proper β cell function as it regulates α\(_2\)δ\(_1\) subunit expression, and in turn influences Cav channel trafficking and activity. Cells lacking α\(_2\)δ\(_1\) subunit show reduced surface expression of Cav1.2 and decreased Ca\(^{2+}\) currents, leading to reduced insulin secretion.

3. The work presented in Paper III concluded that β\(_1\) subunit expression is required for Ca\(^{2+}\) homeostasis. Cells with low β\(_1\) subunit expression exhibit not only low Cav channel activity and impaired insulin secretion, but also apoptosis. Thus, the expression of β\(_1\) subunit is essential for β cell function and survival.

4. The work presented in Paper III showed that the palmitoylation state of β\(_{2a}\) subunit is important for proper Cav channel trafficking and activity. Elevated expression of palmitoylated β\(_{2a}\), however, can induce cell death via elevated cytosolic Ca\(^{2+}\). Therefore, the palmitoylation state of β\(_{2a}\) may also affect β cell survival.
Future Perspectives

Introduction
CaV channels play a crucial role in regulating basal and stimulated insulin secretion in β cells. The α₁ pore-forming subunit of CaV channels is believed to depend on auxiliary subunits not only for membrane trafficking but also for channel activity. In this thesis, novel roles for the auxiliary subunits α₂δ₁, β₁, and β₂a in controlling β cell Ca²⁺ homeostasis, insulin secretion, and survival were identified.

Membrane rafts and exocytosis
To better understand the mechanism behind the elevation in basal secretion following membrane raft dispersion, the members of exocytotic machinery should be further investigated. The expression, localization, and clustering of SNARE proteins such as syntaxin 4, SNAP-25, and synaptotagmin-7 may be explored by implementing qPCR, immunohistochemistry, and immunoblotting. Furthermore, using other cholesterol- or sphingolipid-depleting agents such as nystatin and myriocin to disrupt these rafts might help in better understanding the role of membrane raft composition in regulating β cell function.

CaV channel regulation by α₂δ₁
More studies are needed to further elucidate the role of α₂δ₁ subunit in regulating CaV channels. For example, the α₂δ subunits are believed to anchor on the cell surface via GPI-anchoring in membrane rafts. It would be interesting to explore the role of membrane rafts in regulating α₂δ₁ subunit and the implications on β cell function. In addition, the α₂δ₁ subunit affects whole-cell Ca²⁺ currents. However, to further explore this effect single-channel and capacitance measurements would be considered.

The data in Paper II are based on reducing Tcf7l2 or Cacna2d1 expression. It would be of interest to investigate the effect of overexpressing Tcf7l2 on Cacna2d1 and in turn the effect on CaV channel trafficking and activity. Since TCF7L2 is associated with T2D, it would be interesting to investigate the role of elevated glucose and FFAs on the expression and localization of α₂δ₁ subunit.
β₁ subunit and insulin secretion

To further explore the effect of silencing β₁ subunit on Ca²⁺ signaling, it would be interesting to dissect the cause for the decrease in whole-cell Ca²⁺ currents by using single-channel measurements as in Paper I. Moreover, the role of the β₁ subunit in Caᵥ channel trafficking could also help elucidate the Ca²⁺ signaling effects. If one would follow the effect of β₁ subunit on β cell survival, exploring cellular stress pathways such as ER and mitochondria stress may be considered. It is possible that TCF7L2 not only regulates α₂δ₁ expression but also the expression of the β subunits. Lastly, β₁ overexpression could be applied in order to identify a potential rescue effect, developing our understanding of the role of β₁ subunit in β cells.

β₂a and Ca²⁺ overload

To explore the effect of β₂a on Ca²⁺ signaling, patch-clamp techniques could be utilized to determine the whole-cell and single-channel activity. Moreover, the absence of effect of β₂a on insulin secretion could be further investigated via capacitance measurements that would reflect exocytotic activity. These experiments would explain whether the rise in intracellular Ca²⁺ is due to the number or activity of single Caᵥ channels, and whether exocytosis is compromised or not. Lastly, the hypotheses presented earlier, dynamin and ER stressors, are also worth investigating to better understand the role of β₂a and its palmitoylation state in β cell function and survival. It is also important to confirm the results from Paper I-IV using animal models and human islets to better relate to our physiology.

TCF7L2 and membrane rafts

The transcription factor TCF7L2 is harbors the strongest genetic risk factor for T2D, and hence, investigating this transcription factor is key to understanding the disease. TCF7L2 is regulated by the Wnt/β-catenin signaling pathway and a decrease in this signaling could potentially lower insulin secretion and cause β cell death. Interestingly, disruption of the membrane raft results in decreased Wnt/β-catenin signaling in HEK293 cells [199]. Indeed, as shown in Paper I, dispersion of membrane rafts increases basal insulin release, which is an indication of dysregulated granular exocytosis of a type that bears resemblance with the phenotype observed in T2D. Therefore, further studies are encouraged to better understand the link between membrane rafts, Wnt/β-catenin signaling, and TCF7L2.

Final remarks

In conclusion, this thesis has demonstrated that Caᵥ channel auxiliary subunits are targets of environmental stressors relevant to T2D. Their altered expression leads to impaired β cell function and consequently increases the risk of T2D. Gene auxiliary subunits, namely α₂δ₁, are also targets for genetic risk factors of T2D. Taken
together, understanding the role of Cav subunits in T2D can ultimately lead to better preventive strategies or improved T2D treatment.
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References


[107] K.S. De Jongh, B.J. Murphy, A.A. Colvin, J.W. Hell, M. Takahashi, W.A. Catterall, Specific phosphorylation of a site in the full-length form of the alpha 1 subunit of the cardiac
L-type calcium channel by adenosine 3',5'-cyclic monophosphate-dependent protein kinase, Biochemistry, 35 (1996) 10392-10402.


Roles of voltage-gated Ca\textsuperscript{2+} channel subunits in pancreatic β cells

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