Cell-Specific Ca$^{2+}$ Response
In Pancreatic β-Cells

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Cover illustration:
Calcium imaging in ß-cells with Fura-2
using Openlab 3 software

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Pancreatic β-cells are heterogeneous in many respects, including their secretory responsiveness, glucose sensitivity and metabolic rate. We do not know precisely which mechanisms control functional characteristics, such as time of insulin release (lag-time) or magnitude of the response in each individual cell or islet. The heterogeneity might represent a random phenomenon or could be a manifestation of constant characteristics inherent to individual β-cells. This is an important question because a diminished and delayed first-phase insulin release is an early sign of failing β-cells.

We compared lag-times, initial lowering nadirs and first [Ca^{2+}]_i peak heights in Fura-2 loaded ob/ob mouse β-cells during two consecutive stimulations with glucose. There was a strong correlation of corresponding parameters between the first and second stimulation. Thus, timing and magnitude of the early Ca^{2+} response were individual and reproducible characteristics in β-cells. We then studied Ca^{2+} responses in β-cells from lean mice, diabetic db/db mice and rats stimulated twice with 20 mM glucose and found cell-specific response characteristics also in those cells. This indicates that a cell-specific Ca^{2+} response to glucose is common in rodent β-cells, both normal and diabetic. Another question was whether aggregated β-cells show cell-specific responses. Using the same protocol as for dispersed β-cells, we analysed Ca^{2+} responses in single ob/ob mouse β-cells within a small cluster (3-7 cells), in clusters of medium (about 10 cells) and large size (about 25 cells) and also in intact islets from ob/ob and lean mice. Significant correlations were found between the first and second stimulation for timing and magnitude of [Ca^{2+}]_i rise, and for initial lowering.

Next, we tested if the β-cell response is cell-specific, when induced at different steps of stimulus-secretion coupling. The glycolytic intermediate glyceraldehyde, the mitochondrial substrate KIC, the K_ATP-channel blocker tolbutamide and arginine were used as tools. [Ca^{2+}]_i changes were studied in dispersed β-cells from lean, ob/ob and db/db mice. NAD(P)H responses to glucose and KIC were analyzed as a measure of metabolic flux. With regard to the cell specificity, the correlation between Ca^{2+} and insulin response from individual β-cells was studied using the calcium dye Fluo-3 and Fluozin-3, which is a probe for Zn^{2+}, co-released with insulin. Both timing and magnitude of calcium responses were cell-specific in lean mouse β-cells with all tested secretagogues. β-Cells from ob/ob and db/db mice showed cell-specific temporal Ca^{2+} responses to glyceraldehyde but not to KIC. The lag-time for the Ca^{2+} response to KIC was shorter during the second stimulation in β-cells from hyperglycemic mice. Tolbutamide and arginine induced no cell-specific temporal Ca^{2+} response in dispersed ob/ob and db/db mouse β-cells. However, the timing of tolbutamide-induced response was cell-specific in ob/ob mouse β-cells within intact islets. NAD(P)H changes to glucose were cell-specific in all three mouse models, but the timing of NAD(P)H response to KIC was cell-specific only in lean mice. Thus, a cell-specific response can be induced in normal β-cells at several steps of stimulus-secretion coupling for nutrient-stimulated insulin release. Mitochondrial metabolism generates a cell-specific temporal response in β-cells from lean but not from db/db and ob/ob mice. Cell-specific properties of β-cell ion channels also seem to be affected in these mice.

The relation between mitochondrial mass and parameters of Ca^{2+} responses were investigated in Mitotracker Red and Fluo-3 labelled β-cells using confocal microscopy. Data show that β-cell mitochondrial status may play an important role in determining the timing of [Ca^{2+}]_i changes.

In summary, the early Ca^{2+} response pattern, including the lag-time, the nadir of initial lowering and the height of the first peak response is cell-specific in β-cells. Isolated and functionally coupled β-cells show cell-specific timing of Ca^{2+} responses when stimulated with metabolic and non-metabolic agents. This may be a robust mechanism of importance for the adequate function of β-cells and a basis for the pacemaker function of some cells. A disturbed cell specificity of the mitochondrial metabolism appears to be a marker of β-cell dysfunction in hyperglycemia and diabetes and may explain the delayed insulin release in β-cells from diabetic subjects. Early steps in glucose metabolism seem to be less vulnerable in this regard.

Keywords for indexing: individual β-cells, heterogeneity, islets of Langerhans, cytoplasmic calcium, repetitive glucose stimulation, lag-time, cell-specific, glyceraldehyde, ketoisocaproic acid, tolbutamide, mitochondrial metabolism
ORIGINAL PAPERS

This thesis is based on the following papers, which are referred to in the text by their Roman numerals:


III. Natalia Gustavsson, Gerd Larsson-Nyrén and Per Lindström. Pancreatic β-cells from \textit{db/db} mice show cell-specific [Ca\textsuperscript{2+}], and NADH responses to glucose but not to alpha-ketoisocaproic acid. Pancreas, 31(3), 242-50.

IV. Natalia Gustavsson, Gerd Larsson-Nyrén and Per Lindström. (Manuscript) Cell specificity of Ca\textsuperscript{2+} response to tolbutamide is impaired in β-cells from hyperglycemic mice.

V. Natalia Gustavsson, Golbarg Abedi, Gerd Larsson-Nyrén and Per Lindström. (Manuscript) Timing of Ca\textsuperscript{2+} response in pancreatic β-cells is related to mitochondrial mass.
ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>BSA</td>
<td>bovine serum albumin</td>
</tr>
<tr>
<td>$[Ca^{2+}]_i$</td>
<td>cytoplasmic calcium concentration</td>
</tr>
<tr>
<td>CoA and acetyl-CoA</td>
<td>coenzyme A and its acyl derivative</td>
</tr>
<tr>
<td>cAMP</td>
<td>cyclic adenosine 3' : 5'-monophosphate</td>
</tr>
<tr>
<td>DNAase</td>
<td>deoxyribonuclease</td>
</tr>
<tr>
<td>EDTA</td>
<td>ethylene diamine tetraacetic acid</td>
</tr>
<tr>
<td>IP$_3$</td>
<td>inositol 1,4,5-trisphosphate</td>
</tr>
<tr>
<td>DIC</td>
<td>differential interference contrast</td>
</tr>
<tr>
<td>CCD</td>
<td>charge-coupled device</td>
</tr>
<tr>
<td>$K_{ATP}$-channels</td>
<td>ATP-sensitive potassium channels</td>
</tr>
<tr>
<td>KIC</td>
<td>$\alpha$-ketoisocaproic acid</td>
</tr>
<tr>
<td>SUR1</td>
<td>sulfonylurea receptor 1</td>
</tr>
<tr>
<td>TPEN</td>
<td>tetrakis(2-pyridylmethyl)ethylenediamine</td>
</tr>
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INTRODUCTION

Blood sugar control in mammals is dependent on secretion of hormones from the islets of Langerhans, principally insulin. Pancreatic β-cells measure extracellular glucose concentrations with a high degree of sensitivity and rapidly release insulin in response to a rise in the glucose level. Although the release of insulin is mostly regulated by the glucose level in the plasma, many other nutrients, hormones and neural stimuli can modulate the release. Impairment of the action of insulin causes metabolic disorders, such as glucose intolerance and diabetes. A combination of reduced peripheral insulin sensitivity and β-cell dysfunction characterises type 2 diabetes (non-insulin-dependent diabetes mellitus) (for review Nadal 1999), responsible for 90-95% of all diabetic cases. This disease is associated with disturbances in the insulin release pattern, i.e. a selective loss of first phase secretion, which precedes the other manifestations (Cerasi and Luft 1967), and impairment of the pulsatile rhythm of secretion (Bergsten 2000). The insulin secretory response of type 2-diabetic patients is delayed and decreased (Cerasi et al 1972, Calles-Escandon and Robbins 1987, Polonsky et al 1998). For understanding of the pathogenesis of type 2 diabetes and for development of effective treatment, it is crucial to have knowledge of the mechanisms regulating normal insulin secretion, including the control of the individual β-cell responsiveness.

The structure of a pancreatic islet

Pancreatic islets are mainly composed of insulin-producing β-cells (65-90% in different species), which in most mammals are located in the central part of the islets and surrounded by other hormone-producing cells (Hellerström et al 1960, Pipeleers 1987). Other cell types are glucagon-releasing α-cells, somatostatin-producing δ-cells and pancreatic polypeptide-producing PP-cells, as well as some other rare cell types. Gap junctions connect different β-cells (Orci et al 1973, Int Veld et al 1986) as well as β-cells and other endocrine islet cells, providing a pathway for the cell-to-cell diffusion of hydrophilic molecules, such as ions, metabolites and second messengers (for review Munari-Silem and Rousset 1996). There is evidence that the β-cells in intact islets are functionally coupled and show synchronised activity (Perez-Armendariz et al 1995).
Metabolic signals play an important role in the sequence of cellular events resulting in insulin release during stimulation with glucose and other nutrient secretagogues (Newgard and McGarry 1995, Matschinsky 1996, Tamarit-Rodriguez et al 1998) (Figure 1). Glucose enters through the GLUT transporter (Figure 1). The glucokinase and phosphofructokinase steps are important for the regulation of glycolysis (for review Deeney et al 2000). One of the intermediate glycolytic metabolites is glyceraldehyde phosphate. For experimental purposes glyceraldehyde is often used to by-pass early glycolytic steps. It enters the glycolytic pathway via its phosphorylation by ATP to form glyceraldehyde phosphate. The subsequent metabolism of glyceraldehyde is identical to that of glucose (Figure 1), i.e. metabolised further through glycolysis generating pyruvate and reduced pyridine nucleotides.

Pyruvate undergoes metabolism in the mitochondria and the nucleotides are reoxidised through mitochondrial NADH shuttle systems. An essential step in the stimulus-secretion coupling cascade is ATP production and a change in ATP/ADP ratio. ATP is produced both during glycolysis in the cytoplasm and through oxidative phosphorylation that takes place in mitochondria during the breakdown of pyruvate. α-Ketoisocaproic acid (KIC) can be used to increase mitochondrial metabolism (Fig. 1). KIC is a transamination partner for glutamic acid or glutamine to yield α-ketoglutarate and leucine (Lembert and Idahl 1998). Glucose also increases the production of cytosolic NADH and the reducing equivalents are transported into the mitochondria by the α-glycerolphosphate and malate/aspartate shuttles for rapid ATP synthesis. The rise in ATP-ADP ratio (and, perhaps, other signals) results in closure of ATP-sensitive K channels (K_{ATP}-channels) in the plasma membrane and a depolarisation of the β-cell. Voltage-dependent Ca^{2+}
channels open and the cytoplasmic free Ca\(^{2+}\) level rises, triggering insulin exocytosis (for review, Dunne and Petersen 1991, Misler et al 1992).

Influencing K\(_{ATP}\)-channels with synthetic drugs can also stimulate insulin secretion. Sulfonylureas, which are used for treatment of type-2 diabetic patients, bind to sulfonylurea receptors (SUR1) (Ashcroft and Ashcroft 1992) and in this way block potassium channels (Trube et al, 1986). Sulfonylureas induce a peak of insulin release often followed by a stable period of secretion at a lower rate (Grodsky et al 1969). However, they do not stimulate insulin biosynthesis and even inhibit glucose-stimulated biosynthetic activity in β-cells (Shatz et al 1972, Levy and Malaisse 1975). Arginine directly causes β-cell depolarisation, similarly to cations (Charles et al 1982, Herchuelz et al 1984). In diabetic subjects, the response to sulfonylureas and arginine is less impaired than the response to glucose (Del Guerra et al, 2005).

Glucose-induced insulin secretion is biphasic. There is a transient first phase lasting 5–10 min, which is then followed by a sustained second phase (Curry et al, 1968).

The effect of β-cell secretagogues is not limited to the acute stimulation of insulin release. As an example, glucose may either desensitise or sensitisre the islet to subsequent stimulations depending on concentration and duration of the stimulation (Curry et al 1968, Nesher et al 1989). The terms “time-dependent inhibition” (TDI) and “time-dependent potentiation” (TDP) have been suggested to describe these effects since they require a certain duration of exposure to the stimulatory agent (Nesher et al 1989). These effects seem to operate through a memory induced in the stimulus-secretion pathway and persist after the removal of the stimulator from the medium. Glucose metabolism (Grill et al, 1978) and intracellular pH regulation (Gunawardana and Sharp 2002) may be important cellular mechanisms causing time-dependent effects.

**Ca\(^{2+}\) response in β-cells**

Glucose induces a triphasic Ca\(^{2+}\) response in β-cells (for review Gilon et al 1994). As Fig. 2 shows, a small initial decrease is followed by a large peak increase in [Ca\(^{2+}\)]\(_i\) (Nilsson et al 1988, Gylfe 1989). The initial lowering is caused by sequestration of intracellular Ca\(^{2+}\) to the intracellular pool and is regulated by glucose metabolism (Lund et al 1989). After the peak, [Ca\(^{2+}\)]\(_i\) usually remains increased for as long as the stimulation lasts, and often in the form of oscillations. The timing and magnitude of calcium response are important regulatory factors
because of the close temporal and quantitative relationships between \([\text{Ca}^{2+}]_i\) and insulin secretion patterns (Bergsten et al 1994, Jonas et al 1998). As for insulin release, the \(\text{Ca}^{2+}\) response to glucose is decreased and delayed in diabetic animal β-cells (Lindström et al 1996).

**Heterogeneity concept**

The heterogeneity among pancreatic β-cells is well documented (Salomon and Meda 1986, Herchuelz et al 1991, Van Schravendijk et al 1992). Individual β-cells show different amounts of secretory granules and rough endoplasmic reticulum (Stefan et al 1987) and number of gap junctions (Meda et al, 1980). β-Cells differ in their glucose sensitivity (Kiekens et al 1992) and biosynthetic (Schuit et al 1988) and secretory (Hiriart and Ramirez-Medeles 1991, Pipeleers et al 1994) capabilities. A marked heterogeneity has been noted in the electrical activity among β-cells (Dunne 1990, Kinard et al 1999) and \(\text{Ca}^{2+}\) response pattern (Pralong et al 1990, Gylfe et al 1991, Herchuelz et al 1991, Larsson-Nyrén and Sehlin 1996). Heterogeneity of secretory and electrophysiological responses was found also in intact islets (Aizawa et al 2001). Despite the large number of studies performed on β-cells, the mechanisms underlying functional heterogeneity among pancreatic β-cells are not known. Some evidence suggests that intercellular differences in glucose metabolism and \(K_{\text{ATP}}\)-channel currents are important (Faehling and Ashcroft 1997, Schiut et al 1997). In most of the studies showing heterogeneity in the β-cell population, β-cells were stimulated only once with glucose and the observations were made using unpurified islet cell preparations. These studies have demonstrated group differences but they have not addressed the question of whether different functional characteristics of β-cells within the population reflect inherent qualities of β-cells or random phenomena. Knowledge of the mechanisms determining timing of response in individual β-cells is
important in view of the fact that the first phase of insulin secretion is impaired already in early stages of type 2 diabetes (Cerasi et al 1972).

**Hyperglycemic ob/ob and db/db mice**

The ob/ob mouse mutation was first recognised in 1949 at the Jackson Memorial Laboratory, Bar Harbor, USA (Ingalls et al 1950) and was bred into several mouse sublines (e.g. Umeå ob/ob mouse). The ob/ob syndrome is caused by hyperphagia due to a genetic defect of leptin, a satiety factor that regulates the balance of food intake and consumption (Halaas et al 1995). Characteristic features of this mouse mutation are obesity, hyperinsulinemia and mild hyperglycemia (Stauffacher et al 1967, Stauffacher and Renold 1969). However, these animals do not develop overt diabetes with ketosis (Hunt et al 1976). Islets from ob/ob mice respond normally to stimulators of insulin release (Hahn et al 1974, Hellman et al 1974). One advantage of using Umeå ob/ob mice is the very high proportion of β cells in their pancreatic islets (>90%) (Hellman 1965). Another hyperglycemic animal model is the db/db mouse (Hunt et al 1976). These mice lack functional leptin receptors (Chen et al 1996). Within 6 weeks of age, they develop a severe diabetic syndrome, characterised by obesity, fasting hyperglycemia, glycosuria, and hyperinsulinemia (Berglund et al 1978). Their β-cells show a defective Ca^{2+} response to glucose stimulation (Roe et al 1994).
AIMS OF THE PRESENT THESIS WORK

✓ To determine whether the functional heterogeneity of calcium responses among β-cells represents a cell-specific, inherent quality of pancreatic β-cells or a random phenomenon.

✓ To find out if a cell-specific response can be observed in β-cells from different animal models, in β-cell clusters and intact pancreatic islets.

✓ To determine if Ca\textsuperscript{2+} responses are cell-specific when induced at different steps of the stimulus-secretion coupling.

✓ To test whether the pattern of cell specificity of β-cell response is different in obesity and diabetes.

✓ To investigate if there is a correlation between the mitochondrial mass and the timing and magnitude of calcium response.
MATERIALS AND METHODS

Animals

Pancreatic islets were obtained from non-inbred, 7- to 8-mo-old female $ob/ob$ mice and their lean littermates (Umeå-$ob/ob$), and adult female rats (Sprague–Dawley, 200–250 g). $db/db$ Mice (BKS.Cg-m+/+Lepr$^{db}$) were from Taconic Europe, Ry, Denmark. They were brought to our animal care facility at the age of 6 weeks and were housed until 3-4 months. For experiment series where $db/db$ and lean mice were compared, lean mice were 4-5-mo-old. Principles of laboratory animal care (NIH publication no. 83-25, revised 1985) were followed and the animal care and experimental procedures were carried out in accordance with the standards established by the European Communities Council Directive (86/609/EEC9). The experiments were also approved by the Northern Swedish Committee for Ethics in Animal Experiments.

In overnight fasted $db/db$ mice the blood sugar level was $22 \pm 2$ mM (mean ± S.D., N=33). The urinary glucose level was $17 \pm 2$ mM (N=33). The body weight was $50 \pm 1$ g (N=33). The body weight of $ob/ob$ mice was $73 \pm 2$ g (N=16). The blood sugar in overnight fasted $ob/ob$ mice was $7 \pm 0.4$ mM (N=16), with negative urinary glucose. The body weight of lean mice was $24 \pm 1$ g (N=15) and the blood sugar level was $5 \pm 0.4$ mM (N=15). (Data from Paper IV).

Islet and β-cell preparation

Isolated mouse islets from $db/db$ mice, $ob/ob$ mice and their lean littermates were obtained by collagenase digestion. Single $ob/ob$ and $db/db$ mouse β-cells were prepared by shaking the isolated islets in a Ca$^{2+}$-deficient medium supplemented with EGTA and DNAase (Lernmark 1974). Single lean mouse β-cells were obtained by digestion with 0.025 mg/ml trypsin and 50 µl/ml DNAase. The cells were distributed on polyllysine-coated cover glasses placed in Petri dishes and maintained in tissue culture for 24-48 hours in RPMI 1640 medium supplemented with 10% heat-inactivated fetal calf serum, 60 mg/ml garamycin, 60 mg/ml benzylpenicillin and 2 mM L-glutamine. Subsequent experimental handling was performed in a Krebs-Ringer medium (KRH) having the following composition in mM: 130 NaCl, 4.7 KCl, 1.2 KH$_2$PO$_4$, 1.2 MgSO$_4$, and 2.56 CaCl$_2$ and supplemented with 1mg/ml BSA and 3 mM D-glucose. The medium was buffered with 20 mM HEPES and NaOH to reach pH 7.4 and equilibrated with ambient air. Isolation of rat islets was performed by infiltrating the pancreata with 2.0 mg/ml collagenase via the main pancreatic duct system. The isolated islets were obtained by shaking the pancreas at 37°C. Preparation and culture of rat β-cells was performed as described for the $ob/ob$ mouse β-cells.

Measurements of cytoplasmic Ca$^{2+}$ with Fura-2

Cells and islets were loaded with 1µM Fura-2 for 40 min in KRH medium at 37°C. After rinsing in KRH to remove extracellular Fura-2, the cover glasses were mounted as the bottom of an open chamber and placed on the stage of an inverted microscope within a climate box maintained at 37°C. Three microscope systems were used: Nikon
Diaphot-TMD (Bimica), Zeiss IM and Zeiss Axiovert. We have previously found that absolute values for \([\text{Ca}^{2+}]_i\) may differ slightly when measured in the three microscope systems, but the recorded response pattern is similar. Because of this difference, absolute \([\text{Ca}^{2+}]_i\) values were not compared between sets of experiments performed on different microscopes. The cells and islets were continuously superfused at a flow rate of 0.6 ml/min. The Fura-2 signal was successively excited at the wavelengths 340 and 380 nm with band-pass filters in front of a 75-watt xenon lamp. Using Nikon or Zeiss IM, the resulting emitted fluorescence was measured with a photomultiplier at 510 nm. In experiments, performed on Zeiss Axiovert, an image analysis system Openlab 3 (Improvision, Coventry, UK) was used. Fura-2 was successively excited at the wavelengths 340 and 380 nm during 23 ms each using a 75-watt xenon lamp and a Polychrome IV monochromator (TILL Photonics, Germany). Images were acquired by a CCD camera (Orca ER, Hamamatsu, Japan). The interval between successive cycles of 340/380 nm excitation was 1.26 s or 1.75 s (Nikon and Zeiss Axiovert) or 5 s (Zeiss IM). \([\text{Ca}^{2+}]_i\) was calculated from the ratio of 340 nm and 380 nm signals after background subtraction using the equation described by Grynkiewicz et al (1985), with a \(K_d\) of 224 nM.

**Measurements of cytoplasmic \(\text{Ca}^{2+}\) with Fluo-3**

Experiments were performed using the Leica SP2 spectral laser scanning confocal microscopy system (Leica Microsystems, Mannheim, Germany). Intact islets were labelled with the calcium dye Fluo-3 and placed on coverslips on the bottom of an open perfusion chamber. The chamber was put on the stage of the microscope and maintained at 37°C. Islets were continuously superfused at a flow rate of 0.6 ml/min. Cells were first visualized using transmission laser scanning microscopy. Fluo-3 was excited with the 488-nm line of an argon laser. Fluo-3 responses were recorded from peripheral cells in one randomly chosen cross-section in islets using a Leica ×40 oil immersion lens with numerical aperture 1.3. The resulting fluorescence was recorded in a channel set up to detect emitted light in the range 510–600 nm. Islets were stimulated twice with 20 µM tolbutamide as described above. Images were collected at 2-s intervals, and fluorescence signals from individual cells were measured and analyzed as a function of time by using the Leica Confocal software package (Leica Microsystems). Results were plotted as the change in fluorescence intensity in arbitrary units. The images were analyzed using Leica confocal software.

In Paper V, after the detection of MitoTracker staining (see below) calcium responses from isolated β-cells and from cells within intact islets were measured. Cross-talk from the green into the red channel was calculated in cells loaded with Fluo-3 only and was corrected for in subsequent experiments. Cells and islets were pre-perfused during 5 min and then stimulated with 20 mM glucose during 10 minutes. The resulting fluorescence was recorded in a channel set up to detect emitted green light in the range 510–560 nm. The Fluo-3 response in islets was recorded from one randomly chosen cross-section. The lag-time for \(\text{Ca}^{2+}\) rise was defined as the time from addition of stimulus until the onset of fluorescence rise and the magnitude was calculated as the change in fluorescence intensity (ΔF) expressed as a percentage of the basal fluorescence intensity (F0). The magnitude of calcium response was not
analyzed in islets because of difficulties in the precise alignment of cells during the measurement (Paper V).

**Measurement of NAD(P)H fluorescence**

Cells were incubated 40 min at 3 mM glucose without Fura-2 and then placed on the stage of a Nikon Diaphot-TMD (Japan) microscope system. Cells were excited at 340 nm during 125 ms. The resulting NAD(P)H autofluorescence was measured with a photomultiplier at 510 nm and expressed as arbitrary units. The interval between measurements was 1.26 s. β-Cells were stimulated twice with 20 mM glucose or 20 mM KIC during 10 minutes with a resting period of 30 min at 3 mM glucose between the stimulations. Because measurement of NAD(P)H autofluorescence requires more intensive UV light than Fura-2 measurement, the registration period was limited to 4 minutes to reduce the risk of cell damage.

**Experiment design and definition of response parameters**

β-Cells or islets were initially pre-perifused at 3 mM glucose for 15 min. Each β-cell was then stimulated twice over 10-min periods with a resting period of 30 min between the stimulations, except for experiments from Paper V. The duration of stimulation and resting period were chosen to reduce the risk for persisting effects which may influence the second response (Nesher et al 1989) but with sufficient time to record individual profiles also from cells that respond late. All test media contained 3 mM glucose except in studies with 20 µM tolbutamide, which was applied in KRH medium containing 5 mM glucose to increase the amount of responding cells (Jonkers et al 2001). The two stimulation periods from the each cell were compared with regard to the timing and magnitude of changes in [Ca^{2+}]. The lag-time for the initial lowering was defined as the time from addition of stimulus until the first value below the baseline average, calculated during the 3 minutes preceding the stimulation. The depth of initial lowering (nadir) was calculated as the difference between the baseline and the lowest [Ca^{2+}] value. The lag-time for the Ca^{2+} rise, was defined as the time from the addition of stimulus to the first [Ca^{2+}] value above the extrapolated baseline in a continuing rise to a [Ca^{2+}] peak (Fig 2). The magnitude of first [Ca^{2+}] spike, i.e. peak height was calculated as the difference between the baseline and the highest [Ca^{2+}] value during the first peak (Fig. 2). Superimposed spikes on top of the peaks were not included in those measurements. The experiments were not designed to study calcium oscillations.

In Paper V, β-cells were first scanned for MitoTracker staining, and their calcium response was then recorded during 10-min stimulation with 20 mM glucose.

**Measurement of insulin release from intact islets**

Isolated mouse islets were cultured overnight and loaded with Fura-2 during 40 minutes before being placed in a perifusion chamber (25-30 islets) enclosed in an incubator maintained at 37°C (Larsson-Nyrén and Sehlin 1996). After 15-min preperifusion in KRH with 3 mM glucose, two consecutive glucose stimulations were performed with 20 mM glucose during 7 min with a 30 min period at 3 mM glucose
between the stimulations. Effluent fractions were collected over 15 s and 30 s intervals. Insulin was measured by RIA with mouse insulin as the standard (Heding 1966).

**Measurement of insulin release from single β-cells.**

\(\text{Zn}^{2+}\) efflux was measured as a marker of insulin secretion, which is co-released with insulin (Fredrickson and Bush 2001). β-Cells were prepared as described above. \(\text{Zn}^{2+}\) images with FluoZin-3 were acquired with 488 nm excitation and 510 nm emission using the Openlab image analysis system on the Zeiss Axiovert microscope (see Measurements of cytoplasmic Ca\(^{2+}\) in β-cells). Buffer with a reduced background \(\text{Zn}^{2+}\) level was prepared as described by Qian et al (2003). Briefly, KRH was prepared with all ingredients except calcium and magnesium salts. The buffer was then treated with 5g/dl Chelex-100 (Bio-Rad) for 2 h. The pH was adjusted to 7.4 after Chelex treatment and puratronic grade \(\text{CaCl}_2\) and \(\text{MgSO}_4\) were added to the final concentrations. During all experiments and solution preparation, glass containers were avoided to minimize metal contamination. The \(\text{Zn}^{2+}\) chelator 200 nM TPEN was added. Glass slides for cell preparation were soaked in 2 mM EDTA for 2-4 days to minimize \(\text{Zn}^{2+}\) leaching during the experiments (Kay 2004). Cover glasses with Fura-2 loaded cells were washed in 3 ml KRH containing 2µM FluoZin-3, placed as the bottom of the open chamber and transferred to the microscope. 150 µl KRH with Fluozin-3 was added to the chamber. To stimulate cells, 10 µl of tolbutamide stock solution containing 2µM FluoZin-3 was added to give a final concentration of 100 µM tolbutamide. Our preliminary experiments have shown that the final homogeneous concentration of the stimulator in the chamber is reached within about 5s. For controls, KRH buffer without tolbutamide was applied to the cells. Fluozin-3 fluorescence images were collected every 2s and the average fluorescence intensities were measured in a region of interest (ROI) of ∼20 µm around the cell periphery. Cells were identified by DIC images before the measurement and by the low Zn-Fluozin background from the cell membrane. Fluozin-3 has undetectable penetration into the cell, presumably because of the three negative charges on the molecule at physiological pH (Gee et al 2002, Qian et al 2003). Cells were stimulated during 10 min, after which they were perifused with KRH containing 3 mM glucose for 30 min. After that, cells were again stimulated with 100 µM tolbutamide added to the perifusion medium and \([\text{Ca}^{2+}]_i\) was measured as described above. Attempts were not made to measure \(\text{Zn}^{2+}\) efflux and \(\text{Ca}^{2+}\) response simultaneously. \(\text{Zn}^{2+}\) measurements required higher excitation intensity than calcium measurements with Fura-2. The wavelengths used for calcium measurements (380 and 340 nm) can be harmful for cells during intense exposure. Our system does not allow a change of excitation exposure during simultaneous measurements. We have not compared the magnitudes of \(\text{Zn}^{2+}\) and \(\text{Ca}^{2+}\) response because \(\text{Zn}^{2+}\) dissolves quickly in the surrounding medium and the rise in \(\text{Zn}^{2+}\) outflow is difficult to estimate precisely.

**Measurement of MitoTracker intensity**

Experiments were performed using the Leica SP2 spectral laser scanning confocal microscopy system (Leica Microsystems, Mannheim, Germany). Glasses with cells labeled with Fluo-3 and MitoTracker Red were mounted as the bottom of an open perifusion chamber on the microscope stage and maintained at 37°C. Intact islets were...
placed on coverslips on the bottom of the chamber. In islets, cells were first visualized using transmission laser scanning microscopy. Cells and islets were continuously superfused at a flow rate of 0.6 ml/min and imaged using Leica oil immersion objective lens (×63, zoom 4, for cells and ×40 for islets) with numerical aperture 1.3. The pinhole was adjusted to match the size of one airy unit for each objective and wavelength by the “Airy 1” function of the software. For z-series, the thickness of optical section was 0.6 µm. The pinhole was larger for calcium measurements in isolated β-cells and islets to obtain the thickness of optical section µm. Images of red emission were acquired using excitation wavelength of 543 nm and a 590 nm long pass filter. z-Series scans of about 30 optical sections were made through each cell with 0.5 µm between layers. A maximum intensity projection of the confocal z-series of images of fluorescent images was created. Leica Confocal Software was used to quantify MitoTracker intensity in composite images, which allowed calculation of mean and S.D. of pixel intensities and the construction of three-dimensional and cross sectional profiles. The stained area in single cells was also measured in 3-D projection using a local threshold function and calculated as the proportion of the cell area taken by projected mitochondria. Based on calculation of the area of mitochondria in single sections, we found that a projected area of 60-70% corresponds to 3-4.5% of cell volume. To study the spatial relation between mitochondria and localised increases in [Ca²⁺], the calcium response was compared in areas (regions of interest, ROI) intensively stained with MitoTracker and in not stained and poorly stained areas.

**Chemicals**

Collagenase type I, HEPES and poly-L-lysine were purchased from Boehringer (Mannheim, Germany). BSA, fraction V. EGTA, L-glutamine, KIC, arginine and DNAase were all from Sigma Chemical Co. (St Louis, MO, U.S.A). Culture medium RPMI 1640 was obtained from ICN Biochemicals, and fetal calf serum was from Labkemi AB (Stockholm, Sweden). Fura 2-AM, Fluo-3, FluoZin-3, tetrakis (2-pyridylmethyl)ethilenediamine (TPEN) and Mitotracker Red CMXRos were from Molecular probes Inc. (Eugene, Oregon, USA). Puratronic CaCl₂ and MgSO₄ were from Alfa Aesar (Karlsruhe, Germany). Benzylpenicillin and garamycin were from Schering Co (Kenilworth, NJ, U.S.A). Glyceraldehyde was purchased from The British Drug Houses Ltd, United Kingdom, tolbutamide was from Hoechst AG, Germany. All other reagents were of analytical grade.

**Statistical analysis**

The data are presented as means ± S.E.M. Statistical significance for comparison of responses from the same cell or islet during first and second stimulation was evaluated using Student’s t-test for paired data or the Mann–Whitney U-test. For comparison between sets of experiments two-tailed Student’s t-test for unpaired data was used. Correlation coefficients (r) were computed by using Statworks software (Computer Associates International Islandia, N.Y., USA) or Microsoft Excel. The significance level was set at p less than 0.05. NS (non-significant) means P>0.05.
RESULTS

Glucose-induced Ca$^{2+}$ response in single dispersed β-cells

ob/ob Mouse (Paper I, II and III)

Parameters of Ca$^{2+}$ response in Paper I were calculated in a different way than in papers II and III and are therefore presented in the text. Data from Papers II and III are shown in Table 1.

Comparison of lag-times. In Paper I the lag-time for [Ca$^{2+}$]$_i$ rise was defined as the time from addition of stimulus until the onset of increase. Lag-time for initial lowering was not calculated. The two stimulation periods from each cell were compared with regard to timing and magnitude of changes in [Ca$^{2+}$]$_i$. There was a correlation between lag-times for [Ca$^{2+}$]$_i$ rise with both 10 and 20 mM glucose. The correlation coefficient r was 0.94 for 10 mM glucose, P<0.001, with the averages 169±17 and 201±21 s (P<0.001), for 1st and 2nd stimulation, respectively. For 20 mM glucose the correlation coefficient r was 0.96 (P<0.001), with the averages 170±18 and 183±20 s (P<0.025). Most of the cells reacted with a [Ca$^{2+}$]$_i$ rise within 3 min from the start of stimulation. However, some β-cells did not respond with a [Ca$^{2+}$]$_i$ rise until 7 - 9 min after stimulation. The lag-time during the second stimulation was longer than during the first one both in fast and late responding cells (19% at 10 mM glucose and 8% at 20 mM glucose). The response pattern, including nadir of initial lowering and peak height, was similar in cells with short and long lag-time, i.e. the duration of the lag-time had little influence on the response pattern. Table 1 shows lag-times for [Ca$^{2+}$]$_i$ rise and initial lowering in ob/ob mouse β-cells combined from Papers II and III. The pattern of cell specificity of β-cell response was the same as in Paper I.

Comparison of initial lowering nadirs, peak heights and response patterns. Nadirs of initial lowering showed a correlation coefficient with 10 and 20 mM glucose. The correlation coefficient for nadirs of initial lowering was r=0.93, P=0.001 for β-cells exposed twice to 10 mM glucose (51±3 and 64±4 nM, 1st and 2nd stimulation, respectively; P<0.05) and r=0.79, P<0.001 for β-cells exposed twice to 20 mM glucose (48±2 and 63±2 nM, 1st and 2nd stimulation, respectively; P<0.005). The peak height in Paper I was calculated as the highest [Ca$^{2+}$]$_i$ value during the first peak. The correlation coefficient for peak heights was r=0.51, P<0.01 (average 719±32 and 716±31 nM, 1st and 2nd stimulation) with 10 mM glucose. With 20 mM glucose the correlation coefficient was r=0.77, P<0.001 (average 743±25 and 747±27 nM, 1st and 2nd stimulation, respectively). Peak heights of ob/ob mouse β-cell responses from Paper II and III and their correlation
coefficients are shown in Table 1. In the majority of individual β-cells the pattern of first response was very similar to the second one (Figure 3 in Paper I).

Lean mouse and rat (Paper II)

The lag-times for $[Ca^{2+}]_i$ rise in lean mouse β-cells and rat β-cells were similar to those obtained in single ob/ob mouse β-cells, and there was a significant correlation between the lag-times (Table 1). $Ca^{2+}$ response parameters in glucose-stimulated lean mouse β-cells from Papers II and III are presented in Table 1. The lag-time for initial lowering was shorter in β-cells from lean mice and rats than in ob/ob mouse β-cells, P<0.01 when comparing 78 s with 47 s (lean mouse) or 48 s (rat). There was a correlation between lag-times for initial lowering in lean mouse β-cells (Table 1). The correlation for the nadir of initial $[Ca^{2+}]_i$ lowering was $r=0.1$, NS (32±3 and 26±2 nM Ca$^{2+}$, first and second stimulation, NS). Corresponding values in rat β-cells were $r=0.57$, P<0.01 (39 ±4 and 40 ±4 nM Ca$^{2+}$, NS). The magnitude of the calcium rise was nearly the same in single dispersed β-cells from lean mice and rats as in single dispersed ob/ob β-cells, with a significant correlation within the pairs of stimulation in all three types of β-cells (Table 1).

db/db Mouse (Paper III)

Lag-times for calcium rise in were longer db/db mouse β-cells than in lean mouse β-cells but showed a correlation between stimulations (Table 1). Seventeen out of 30 (17/30) β-cells showed initial lowering during both stimulations, which was fewer than in lean mouse β-cells (21/23) (p<0.001, using Chi-square test with Yates’ correction). Lag-times for initial lowering were cell-specific (Table 1). Nadirs of initial lowering were not correlated ($r=0.26$, NS) and showed no difference between the stimulations (50±16 and 30±7 nM Ca$^{2+}$, first and second stimulation, respectively, NS). Although the magnitude of calcium response was lower than in lean mouse β-cells, peak heights were cell-specific (Table 1). Basal $[Ca^{2+}]_i$ levels were higher than in lean and ob/ob mouse β-cells (130±15 nM vs. 56±3 nM and 93±12 nM Ca$^{2+}$).

We compared timing and magnitude of responses with (17/30) and without initial lowering (13/30). Lag-times did not differ between the groups (189±36 s and 215±36 s, respectively, NS, first stimulation; 152±30 s and 198±33 s, respectively, NS, second stimulation). Responses with initial lowering had higher magnitude. This was observed both during the first (100±25 nM and 221±35 nM Ca2+, P<0.01) and the second stimulation (134±14 nM and 241±44 nM Ca2+, P<0.01). Timing and magnitude of response were cell-specific in both groups.

Glucose-induced Ca$^{2+}$ response in β-cell clusters and islets (Paper II)

Three cluster models were used. First, calcium response was investigated in one single β-cell within a small cluster from ob/ob mouse, referred as ‘single β-cell
within a small cluster’. The cluster consisted of three or four cells in physical contact. Then we enlarged the measurement area and studied temporal Ca\(^{2+}\) profiles in whole medium (about 10 cells) and large size clusters (about 25 cells).

**Comparison of lag-times.** Lag-times for [Ca\(^{2+}\)]\(_i\) rise and for initial [Ca\(^{2+}\)]\(_i\) lowering were cell-specific in ‘single β-cell within a small cluster’ (Table 1). Responses from the majority of medium size clusters were very similar to those obtained from big clusters. There was a strong correlation between the first and second lag-time for [Ca\(^{2+}\)]\(_i\) rise in both sets of experiments. Lag-times for initial lowering showed a correlation only in medium size clusters. In large clusters, lag-time for [Ca\(^{2+}\)]\(_i\) rise was shorter (<1 min) than in single dispersed β-cells and ‘single β-cells within a small cluster’ (Table 1).

Intact islets showed longer lag-times for [Ca\(^{2+}\)]\(_i\) rise when compared with medium and large size clusters. Lag-times for [Ca\(^{2+}\)]\(_i\) rise were cell-specific in islets from ob/ob but not from lean mice. Lean mouse islets responded with a longer lag-time during the first stimulation, and ob/ob mouse islets responded with a longer lag-time during the second stimulation (Table 1).

**Comparison of nadirs of initial lowering and peak heights.** ‘Single β-cells within a small cluster’ showed cell-specific nadirs of initial lowering in [Ca\(^{2+}\)]\(_i\) (24±2 and 29±2 nM Ca\(^{2+}\), P<0.05 with r=0.59, P<0.001). Initial lowerings were observed during both stimulations in 4/19 medium size clusters (9±3 and 10±2 nM Ca\(^{2+}\), r=0.68) and in 4/22 large clusters (21±9 and 24±9 nM Ca\(^{2+}\), r=0.49). For lean mouse islets the corresponding values were 10±1 and 4±1 nM Ca\(^{2+}\) (P<0.001), with a correlation of r=0.59 (P<0.05). Lag-times for initial lowering were not cell-specific in lean mouse islets. Initial lowerings were seldom observed in islets from ob/ob mice. Peak heights were cell-specific in cluster models and intact islets. The peak heights were reduced as the number of cells in contact increased (Table 1). The first peak height was lower than the second one in clusters and in lean and ob/ob mouse islets (Table 1).

**Insulin release dynamics in ob/ob mouse islets (Paper II)**

We measured insulin release from perifused ob/ob mouse islets to determine if the prolonged lag-time and reduced Ca\(^{2+}\) response observed during the second glucose stimulation corresponded to the time of onset and amount of insulin release, (Fig. 4C and 4D in Paper II). Batches of 25-30 islets were stimulated twice with 20 mM glucose following the same protocol as for the [Ca\(^{2+}\)]\(_i\) measurements. Onset of stimulated secretion was delayed (from 45–60 s to 60–75 s) during the second exposure to 20 mM glucose (P<0.05, using the Mann–Whitney U-test). The total amount of secretion was lower during the second stimulation (0.027±0.005 ng/islet/min and 0.011±0.003 ng/islet/min, first and second stimulation, respectively, P<0.001).
Ca^{2+} response to nutrient and non-nutrient insulin secretagogues in lean, ob/ob and db/db mouse ß-cells (Paper III and IV)

Glyceraldehyde

When single dispersed lean, ob/ob and db/db mouse ß-cells were stimulated with the glycolytic intermediate, 10 mM glyceraldehyde, the Ca^{2+} pattern was similar to glucose-induced Ca^{2+} changes. Lag-times for calcium rise and peak heights were cell-specific in all three animal models (Table 1 and 3).

In lean mouse ß-cells, lag-times for initial lowering were cell-specific and nadirs of initial lowering also showed a correlation (r=0.62, P<0.001; 18±5 and 20±3 nM Ca^{2+}, first and second stimulation, NS). Peak height was smaller during the second stimulation, but peak heights showed a correlation between the first and second stimulation (Table 1).

In ob/ob mouse ß-cells, the response patterns were almost identical during the first and second exposure, but the lag-time for initial lowering in ob/ob mouse ß-cells showed no correlation within the pairs (Table 1). Nadirs of initial lowering were the same during both stimulations (27±4 and 25±3 nM Ca^{2+}, first and second stimulation, respectively, NS) and showed a correlation of r=0.69 (P<0.001).

In db/db mouse ß-cells, initial [Ca^{2+}]i lowerings were observed in only 2 out of 13 cells (2/13). Peak heights were lower compared with glyceraldehyde-induced responses from lean mouse ß-cells, but there was no difference in lag-time for calcium rise (Table 1).

KIC

The mitochondrial substrate, 20 mM KIC, induced cell-specific responses both with regard to timing and magnitude in ß-cells from lean but not from ob/ob or db/db mice (Table 1 and 3). The [Ca^{2+}]i rise occurred earlier than when glucose or glyceraldehyde was applied (Table 1, P<0.001). Initial lowering less pronounced (Lenzen et al 2000) and was observed during both stimulations in 6/16 cells. This is in line with previous observations. There was no correlation between lag-times for initial lowering (Table 1) or between nadirs (r=-0.96, NS).

ob/ob Mouse ß-cells showed a correlation between peak heights (Table 1). Initial lowerings were observed in 25/31 cells during both stimulations. There was no correlation of lag-times for initial lowering (Table 1) or the nadirs (20±3 and 26±3 nM Ca^{2+}, r=-0.14, NS). The rise during the second stimulation occurred with a shorter lag-time than during the first stimulation (Table 1) and had a smaller variation (0-390 s vs.0-155 s).

In db/db mouse ß-cells, peak heights were also cell-specific (Table 1). As in ob/ob mouse ß-cells, the second lag-time was shorter and varied less than the first (75-346 s vs. 68-208 s). Initial lowering was observed during both stimulations in
only 2/14 cells. The Ca\textsuperscript{2+} response to KIC was delayed and reduced when compared with lean mouse β-cells (Table 1, p<0.05).

*Tolbutamide and arginine*

When the K\textsubscript{ATP}-channel blocker, 100 µM tolbutamide was applied, the lag-time and magnitude of [Ca\textsuperscript{2+}], rise were cell-specific in lean mouse β-cells but not in ob/ob or db/db mouse β-cells (Table 1). Unlike glucose, tolbutamide induced responses with a shorter lag-time in β-cells from hyperglycemic mice when compared with lean mouse β-cells. There was no initial lowering of [Ca\textsuperscript{2+}], before the rise, probably because initial lowering is regulated by metabolism (Lund et al 1989). In ob/ob mouse β-cells, the peak height was lower during the second stimulation than during the first one (Table 1). We then tested tolbutamide at 20 µM to find out if the lag-time is concentration-dependent. The lower concentration could provide more heterogeneous responses with regard to timing and reduce the risk for persisting effects after the first stimulation. As with 100 µM tolbutamide, the timing of response was cell-specific only in lean mouse β-cells. Peak heights were cell-specific in all three models (Table 1). The lag-time for [Ca\textsuperscript{2+}], rise was longer than in experiments with 100 µM tolbutamide (Table 1, P<0.01).

Arginine induced cell-specific Ca\textsuperscript{2+} responses in lean mouse β-cells (Table 1). There was no correlation between lag-times for calcium rise during the first and second stimulation in β-cells from db/db or ob/ob mice, but peak heights were cell-specific in db/db mouse β-cells (Table 1). As with tolbutamide, the response in db/db mouse β-cells occurred with shorter lag-times than in lean mouse β-cells and reached a lower magnitude.

Thus, β-cell ion channel function seems to be cell-specific in normal mice but not in hyperglycemic mice (Table 3).
## Table 1

<table>
<thead>
<tr>
<th>Condition</th>
<th>Group Description</th>
<th>Lag-time for initial lowering (s)</th>
<th>Lag-time for calcium rise (s)</th>
<th>Peak height (nM Ca²⁺)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1 stim</td>
<td>2 stim</td>
<td>r</td>
</tr>
<tr>
<td><strong>20 mM glucose</strong></td>
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<td>47±8</td>
<td>62±10</td>
<td>0.85***</td>
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<td></td>
<td>db/db mouse β-cells (30)</td>
<td>42±11</td>
<td>57±10</td>
<td>0.71***</td>
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<td>ob/ob mouse β-cells (38)</td>
<td>68±8</td>
<td>102±14*</td>
<td>0.91***</td>
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<td>rat β-cells (39)</td>
<td>48±8</td>
<td>43±9</td>
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<td>single β-cells within a small cluster (53)</td>
<td>37±5</td>
<td>39±5</td>
<td>0.57***</td>
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<td></td>
<td>medium size clusters (19)</td>
<td>13±5</td>
<td>13±6</td>
<td>0.72NS</td>
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<td>large size clusters (22)</td>
<td>14±4</td>
<td>22±6</td>
<td>0.99***</td>
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<tr>
<td></td>
<td>ob/ob mouse islets (27)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>lean mouse islets (17)</td>
<td>41±4</td>
<td>36±3</td>
<td>0.31NS</td>
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<td>35±4</td>
<td>48±8</td>
<td>0.69**</td>
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<td>94±37</td>
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<td>36±11</td>
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<td>94±36</td>
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<td>-0.04NS</td>
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<td><strong>100 µM tolbutamide</strong></td>
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<td>db/db mouse β-cells (11)</td>
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<td>ob/ob mouse β-cells (24)</td>
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<td><strong>100 µM tolbutamide, insulin and Ca²⁺ response, see explanation in Results, p. 27</strong></td>
<td>lean mouse β-cells (16)</td>
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<td>β-cells in lean mouse islets (30)</td>
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<td>β-cells in ob/ob mouse islets (86)</td>
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<td><strong>10 mM arginine</strong></td>
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<td>ob/ob mouse β-cells (13)</td>
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</table>
Cell specificity of Ca\(^{2+}\) response to tolbutamide in single β-cells within intact islets

We studied responses from individual β-cells on the periphery of Fluo-3-labeled intact islets (Fig. 3) during two consecutive stimulations with 20 µM tolbutamide. We tested 30 cells in 6 lean mouse islets and 87 cells in 10 ob/ob mouse islets. β-Cells within lean mouse islets showed cell-specific responses with a high degree of correlation between lag-times during the first and second stimulation (Fig. 4C in Paper IV). Within each islet, there was only a small variation in lag-time between different cells and between the first and second stimulation (Table 1). Peak heights were cell-specific (Table 1).

β-Cells within ob/ob mouse islets also showed cell-specific timing of responses (Table 1). The average lag-time was longer than in lean mouse (Table 1) and varied more between individual cells (in 0-350 ob/ob mouse islets vs. 0-128 in lean mouse islets). Thus, individual β-cells within an ob/ob mouse islet showed temporal cell specificity of the Ca\(^{2+}\) response.

NAD(P)H response to glucose and KIC in individual lean, ob/ob and db/db mouse β-cells (Paper III)

Similar to what we found for Ca\(^{2+}\) responses, 20 mM glucose induced cell-specific NAD(P)H responses with regard to timing and magnitude in all three animal models (Table 2). Timing and magnitude of NAD(P)H rise to 20 mM KIC were cell-specific in lean mouse β-cells but not in db/db mouse β-cells (Table 2). β-Cells from ob/ob mice showed no cell specific timing of the response, but the fluorescence increases showed a correlation between the first and second stimulation.
Table 2. Glucose- and KIC-induced NAD(P)H response in β-cells. Data is presented as mean values±S.E.M. for the number of experiments given in parenthesis. *P<0.05, **P<0.005, ***P<0.001.

<table>
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<th>Lag-time 1 (s)</th>
<th>Lag-time 2 (s)</th>
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<th>Fluorescence increase 1 (a.u.)</th>
<th>Fluorescence increase 2 (a.u.)</th>
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<tr>
<td>lean mouse β-cells (11)</td>
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<td>40±5</td>
<td>0.60**</td>
<td>10±2</td>
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<td>11±2</td>
<td>9±1</td>
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<td>lean mouse β-cells (14)</td>
<td>27±3</td>
<td>24±4</td>
<td>0.55*</td>
<td>7±1</td>
<td>4±1</td>
<td>0.72**</td>
</tr>
<tr>
<td>db/db mouse β-cells (11)</td>
<td>49±8</td>
<td>32±4</td>
<td>0.39NS</td>
<td>6±1</td>
<td>6±1</td>
<td>0.40NS</td>
</tr>
<tr>
<td>ob/ob mouse β-cells (15)</td>
<td>32±6</td>
<td>33±7</td>
<td>0.46NS</td>
<td>7±1</td>
<td>5±1</td>
<td>0.56*</td>
</tr>
</tbody>
</table>

Insulin secretion in isolated β-cells from lean and ob/ob mice (Paper IV)

We have also tested to what extent temporal cell specificity of the calcium response is related to insulin release. Zn$^{2+}$ is co-released with insulin and Zn$^{2+}$ efflux was measured as a marker of insulin secretion (Qian et al 2003). The secretory response to 100 µM tolbutamide was recorded during the first stimulation and the Ca$^{2+}$ response was recorded during the second stimulation (Fig. 3 in Paper IV). We found that the time of onset of Zn$^{2+}$ outflow correlated with the lag-time for calcium rise in lean mouse β-cells but not in ob/ob mouse β-cells (Table 1).

Table 3. Temporal cell specificity of β-cell stimulus-secretion coupling in studied models.

<table>
<thead>
<tr>
<th>Model</th>
<th>Glycolysis</th>
<th>Mitochondrial metabolism</th>
<th>Ion channels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal β-cells</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Hyperglycemic mouse β-cells (ob/ob and db/db)</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Mitochondrial state and Ca\(^{2+}\) response in β-cells (Paper V)

The morphology of mitochondria ranged from small grains to large elongated or spindle-shaped structures (Fig. 4A and 4B). Mitochondria were observed mostly around the nucleus, they were unevenly distributed, and many of them seemed to be connected. To compare the mitochondrial mass between lean (N=9) and obese mice (N=22), we analyzed the MitoTracker staining intensity and the area taken by mitochondria projected onto the cell middle section. The projected area percentage was higher in lean mouse β-cells than in \textit{ob/ob} mouse β-cells (72±5 % and 50±4 %, respectively, \(P<0.05\)) but the intensity of MitoTracker staining did not differ (137±14 a.u. and 116±8 a.u., respectively, NS). The intensity showed a correlation with the stained area both in lean and obese mouse β-cells (\(r=0.85, P<0.01\) and \(r=0.42, P<0.05\)).

![Figure 4](image)

**Figure 4.** Active mitochondria in a single lean mouse β-cell stained with MitoTracker Red CMXRos and examined by fluorescence microscopy. Panel A shows a projection of a z-series comprising 24 slices acquired at 0.5 µm intervals. Panel B shows one section from the series.

The intensity of Mitotracker staining in β-cells before the stimulation was analyzed together with the lag-times and magnitude of the calcium response to 20 mM glucose. The mitochondrial mass was inversely correlated with the lag-times for \([\text{Ca}^{2+}]_i\) rise in both lean and \textit{ob/ob} mouse β-cells (\(r=-0.73\) and \(r=-0.43\), respectively, \(P<0.05\)). Thus, cells with short lag-times had a higher amount of active mitochondria before the stimulation. The Fluo-3 increase was lower in β-cells from lean mice than in \textit{ob/ob} mouse β-cells (42±9 % and 85±12 %, \(P<0.001\)).
There was no correlation between the magnitude of responses and the mitochondrial mass in either lean or ob/ob mouse β-cells (r=0.15 and r=0.05, NS).

The [Ca\textsuperscript{2+}]\textsubscript{i} rise occurs as localized [Ca\textsuperscript{2+}]\textsubscript{i} increases within the β-cell (Bokvist et al 1995, Qian and Kennedy 2001). We investigated whether areas of those increases were co-located with mitochondria. Fluo-3 does not enter organelles and therefore we assume that only cytoplasmic free calcium was measured. Responses in different areas of the β-cell cytoplasm were almost simultaneous with the time-scale used but varied in magnitude. Traces with a steep rise and a high magnitude suggested “hot spots” of intracellular calcium dynamics. In some cytoplasmic areas no [Ca\textsuperscript{2+}]\textsubscript{i} increase was observed (Fig. 3A and 3C in Paper V). We tested the correlation between the MitoTracker intensity and the Fluo-3 fluorescence increases in different ROIs. We found an inverse correlation between the magnitude of [Ca\textsuperscript{2+}]\textsubscript{i} rise and the MitoTracker intensity (r=-0.64, P<0.001). It seemed logical, since mitochondria were observed mostly around the nucleus, and the calcium channels are situated on the membrane. We then compared the magnitude of response in ROIs along the membrane near areas densely populated with mitochondria (group 1) and on the opposite side of the cell, where fewer mitochondria were observed (group 2). The magnitude of calcium increases was not correlated with the proximity of active mitochondria (95±29% and 55±13%, group 1 and group 2, respectively, N=9 lean mouse β-cells, NS).

To find out if it is possible to use MitoTracker fluorescence intensity to identify fast-reacting cells within intact islets, we measured the mitochondrial mass in individual cells within islets from lean and ob/ob mice and subsequently registered their calcium response. We tested 26 cells in 3 lean mouse islets and 33 cells in 3 ob/ob mouse islets. There was a narrow range of lag-times for [Ca\textsuperscript{2+}]\textsubscript{i} rise between β-cells within each lean mouse islet. Lag-times in β-cells within ob/ob mouse islets showed a larger variation. In most islets, no correlation was found between the lag-times for [Ca\textsuperscript{2+}]\textsubscript{i} rise and the MitoTracker staining intensity. One of the ob/ob mouse islets showed r=-0.48 (P<0.05, N=18 cells).

DISCUSSION

Paper I reports on the early [Ca\textsuperscript{2+}]\textsubscript{i} dynamics elicited by consecutive glucose stimulations in ob/ob mouse β-cells. The lag-time for [Ca\textsuperscript{2+}]\textsubscript{i} rise, the nadir of initial [Ca\textsuperscript{2+}]\textsubscript{i} lowering and the height of first peak showed a correlation within the pairs when the conditions during the two stimulations were not altered. The results therefore suggest that the response from single β-cells have unique and reproducible characteristics. Heterogeneous Ca\textsuperscript{2+} handling has been observed also in airway epithelial cells (Evans and Sanderson 1999), HIT insulinoma cells (Prentki et al 1988), hepatocytes (Rooney et al 1989) and astrocytes (Yagodin et al 1995).
We analysed variations in lag-time for \([\text{Ca}^{2+}]\) rise between different \(\beta\)-cells and between the two test periods for each \(\beta\)-cell. It was found that at both 10 and 20 mM glucose the second lag period was slightly longer than the first (Paper I), which is interesting because the second stimulation also tended to cause a larger initial lowering in the \(\text{Ca}^{2+}\) level. However, because this enlarged lowering was only seen at 20 mM and not at 10 mM glucose and there was no correlation between the lag-times and initial dips in \(\text{Ca}^{2+}\) level at either 10 or 20 mM glucose, there seems to be no clear-cut relationship between these two glucose effects.

Some \(\beta\)-cells showed a very late response to glucose. Different timing for \([\text{Ca}^{2+}]\) response could be an indication of various degrees of cell damage. It is also possible that damaged cells can show reproducible response patterns. However, cell damage is not likely to be the reason why the response is reproducible and is observed also in clusters of cells and in intact islets (Paper II). It is quite remarkable that the response pattern was similar in early and late-responding cells. This indicates that the lag-time and the pattern of \([\text{Ca}^{2+}]\), changes are controlled by different mechanisms.

\(\beta\)-Cells from normoglycemic mice and rats also showed reproducible \([\text{Ca}^{2+}]\) changes (Paper II). Although mouse and rat pancreatic islets differ with regard to glucose-induced changes in cytoplasmic calcium (Antunes et al 2000), a cell-specific response at the single cell level was observed in both species. The lag-times for \([\text{Ca}^{2+}]\) rise were very similar in single dispersed \(\beta\)-cells from both lean and \(ob/ob\) mice and from rat. However, the lag-time for initial lowering was nearly twice as long in \(ob/ob\) mouse \(\beta\)-cells, when compared with \(\beta\)-cells from lean mice and rat. This suggests that the metabolic signal(s) differ for onset of initial lowering and later \([\text{Ca}^{2+}]\) rise.

In Paper II we show that the timing of the \(\text{Ca}^{2+}\) response and the magnitude of response can be reproduced during a second stimulation both in \(\beta\)-cells in clusters of different size and in intact islets. We also find that the lag-time for initial lowering and \([\text{Ca}^{2+}]\) rise is reduced when the number of cells in contact is increased. The shorter lag-time may depend on cell-to-cell transfer of signals. With the experimental set up it was not possible to study \(\text{Ca}^{2+}\) profiles from individual cells within either medium or large size clusters. We, therefore, cannot say how many fast reacting \(\beta\)-cells (Paper I) need to be in contact to reduce the lag-times to the extent observed when comparing single \(\beta\)-cells and medium size clusters. Jonkers et al (1999) have shown that the number of cells responding to a particular glucose concentration is enhanced in clusters of as few as 2–50 cells.

A delayed and reduced second \([\text{Ca}^{2+}]\), response in \(ob/ob\) islets corresponds to the findings on insulin secretion. A later response during the second stimulation conforms to previous finding of a stepwise reduction in insulin release dynamics in response to repeated glucose stimulations in perfused pancreata from C56BL/KsJ-mice (Berglund 1980). It may be that inhibitory signals produced during the first exposure are more effective when the cluster architecture is intact. Ionic events and diffusible compounds, such as NO (Grapengiesser et al 1999), may carry such inhibitory signals.
The strong reproducibility of the early \([\text{Ca}^{2+}]_i\) changes suggests that each ß-cell has a cell-specific capacity to respond to a glucose challenge within a certain time and with a unique response pattern. Individual ß-cells may have different roles in intact pancreatic islets. It has been suggested that there are subpopulations of ß-cells, which act as sensors within islets capable of initiating an appropriate response in other, less sensitive cells (Hiriart and Ramirez-Medeles 1991, Van Schravendijk et al 1992). The possibility that a subset of ß-cells have an actual pacemaker function has also been discussed (Cook 1984, Palti et al 1996, Aslanidi et al 2002) but the mechanisms underlying such a function are unclear. ß-Cells maintain their individual responses when they are functionally coupled in clusters and islets. This in line with the study by Jonkers et al (1999) suggesting the possibility that the fastest cells in an islet entrains all the other ß-cells, to which it is electrically coupled. However, intercellular coupling or paracrine influences in vivo may also erase individual differences of ß-cells and create a functionally more homogenous population.

An important question was whether cell specificity of ß-cell response is impaired in hyperglycemia and type 2 diabetes (Paper III). We found that \(db/db\) mouse ß-cells show cell-specific responses to glucose. Thus, cell specificity is not by itself a sign of normal ß-cell function. However, when several insulin secretagogues were tested, ß-cells from normal and hyperglycemic mice showed different patterns. Cells from lean mice showed cell-specific \(\text{Ca}^{2+}\) responses to all tested stimulators, which enter the stimulus-secretion coupling at different steps. This indicates that the response of each individual cell is under tight control and that the activity of several steps is cell-specific. Data on NAD(P)H responses to glucose and KIC suggest that cell specificity of nutrient-induced calcium response is determined by steps not later than mitochondrial metabolism. ß-Cells from \(ob/ob\) and \(db/db\) mice showed no cell-specific timing of NAD(P)H responses when stimulated with KIC. There was also no cell-specific lag-time for calcium rise in ß-cells from hyperglycemic mice with KIC, tolbutamide or arginine. Bypassing glycolysis or the whole metabolism caused a loss of temporal cell specificity in hyperglycemic mouse ß-cells but the magnitude of calcium response was still cell-specific. This allows us to make some further assumptions. Firstly, timing and magnitude of ß-cell response are controlled by different mechanisms. Secondly, mitochondrial function and ion channel function seem to be cell-specific in normal ß-cells but not in \(db/db\) and \(ob/ob\) mouse ß-cells. Glycolysis signals can generate a cell-specific response also when mitochondrial metabolism cannot. Thirdly, tests for cell specificity of ß-cell responses can be an important tool for studies of normal and disturbed ß-cell function.

ß-Cells from \(db/db\) mice showed delayed and reduced glucose-induced \([\text{Ca}^{2+}]_i\) responses and higher basal \([\text{Ca}^{2+}]_i\) levels, when compared with lean mouse ß-cells. This is similar to what has been shown earlier on insulin release and \(\text{Ca}^{2+}\) response for animals with hereditary and chemically induced type 2 diabetes (Berglund et al 1984, Roe et al 1994, Lindström et al 1996, Zaitsev et al 1997, Marie et al 2001). Mitochondrial dysfunction in type 2 diabetes has also been described (Maechler and Wollheim 2001, Malaisse 1993). The altered mitochondrial metabolism in \(db/db\) mouse ß-cells in experiments with KIC can be a sign of such dysfunction.
However, such changes were found also in ob/ob mice, whose \( \beta \)-cells respond adequately to glucose stimulation (Hahn et al 1974, Hellman et al 1974) and where most aspects of mitochondrial metabolism are normal (Idahl and Lembert, 1995). Therefore the altered mitochondrial function could be a compensatory mechanism to prolonged hyperglycaemia.

Exogenously added glyceraldehyde enters the glycolytic pathway via its phosphorylation in the triokinase reaction to yield glyceraldehyde phosphate (MacDonald 1989). The subsequent metabolism is probably identical to that of glyceraldehyde phosphate derived from glucose. Glyceraldehyde can cause acidification of the \( \beta \)-cell, which may affect the cell function (MacDonald 1989). The \( \text{Ca}^{2+} \) response to glyceraldehyde was cell-specific in lean, ob/ob and \( \text{db/db} \) mouse \( \beta \)-cells, and the lag-time for calcium rise was the same with glyceraldehyde as with glucose. It is therefore possible that later parts of glycolysis are more important than the earliest metabolic steps including the glucokinase reaction for determining whether a cell reacts fast or slow. Lag-time for initial lowering was shorter in the presence of glyceraldehyde than in the presence of glucose. As already suggested above, various aspects of the \( \text{Ca}^{2+} \) response are probably controlled by different steps of the stimulus-secretion coupling, i.e. the lag-time for the lowering in \( \text{Ca}^{2+} \) may be controlled by early metabolic steps closer to the glucose phosphorylation, whereas lag-time for \([\text{Ca}^{2+}]\) rise is determined by later metabolic events. This may be why the lag-time for initial lowering was not reproduced during the second stimulation in studies with glyceraldehyde.

KIC is a transamination partner, which feeds into mitochondrial metabolism through \( \alpha \)-ketoglutarate (Lembert and Idahl 1998) and induces a rapid increase in mitochondrial NADH generation (Panten 1975). KIC shortened the lag-time for calcium rise during the second stimulation in ob/ob and \( \text{db/db} \) mouse \( \beta \)-cells but not in lean mouse \( \beta \)-cells. The lag-times for the calcium rise in \( \beta \)-cells from hyperglycemic mice showed a smaller variation during the second stimulation than during the first stimulation. Therefore, the difference between lag-times for calcium rise was more pronounced in cells responding late during the first stimulation. Precise mechanism of the accelerating effect of KIC on the calcium response in ob/ob and \( \text{db/db} \) mouse \( \beta \)-cells remains to be clarified, but this observation suggests an intriguing possibility to specifically improve \( \beta \)-cell function in diabetic animals.

Normal \( \beta \)-cells had unique and reproducible, i.e. cell-specific, calcium responses also to the non-metabolic stimulators tolbutamide and arginine. This indicates cell specificity not only of the stimulus-secretion coupling, as a whole, but also of different parts of it. While cell specificity seems to be a robust phenomenon in normal \( \beta \)-cells, ob/ob and \( \text{db/db} \) mouse \( \beta \)-cells show no cell-specific timing of calcium response when stimulated with tolbutamide or arginine. The mechanism of action of arginine is a direct depolarisation of the \( \beta \)-cell membrane similar to cations (Herchuelz et al 1984, Charles et al 1982). This means that the function of \( \text{Ca}^{2+} \) and maybe \( K^+_{\text{ATP}} \) channels is cell-specific in \( \beta \)-cells from normal mice but not in ob/ob and \( \text{db/db} \) mouse \( \beta \)-cells. Cell-specific function of calcium channels may be linked to the number of channels and/or differences in channel properties, such
as the gating. Previous reports have shown different results regarding calcium channel function in β-cells from animals with conditions similar to type 2 diabetes. A reduction of L-type Ca\(^{2+}\) currents and a decreased expression of alpha 1-subunit of L-type Ca\(^{2+}\) channels was observed in the Zucker diabetic fatty rat (Roe et al 1996). Other studies report an increased activity of L-type Ca\(^{2+}\) channels in the GK rat (Kato et al 1996). A lower expression of Ca\(^{2+}\) channels can explain the reduced calcium response in type 2 diabetes but not the loss of temporal cell specificity, which we found in db/db and ob/ob mouse β-cells. This indicates that other alterations of Ca\(^{2+}\) channels are present in β-cells hyperglycemic mice (Kato et al 1996).

The failure of db/db and ob/ob mouse β-cells to show cell-specific timing of response to a K\(_{\text{ATP}}\)-channel blocker (tolbutamide) may be due to increased activity of K\(_{\text{ATP}}\)-channels and impaired electrical activity (Ashcroft and Rorsman 2004). The cell-specific magnitude of response to KIC and tolbutamide in β-cells from hyperglycemic mouse indicates that cell-specific timing and magnitude of the Ca\(^{2+}\) response are controlled by different mechanisms.

β-Cells from type 2 diabetic subjects show a delayed Ca\(^{2+}\) response to glucose due to metabolic disturbances, but they react with a normal lag-time when challenged with arginine or sulfonylureas (Del Guerra et al 2005, Zaitsev et al 1997). β-Cells from db/db mice in fact responded to tolbutamide and arginine with a shorter lag-time than lean mouse β-cells and a similar tendency was found also in ob/ob mouse β-cells. It may be that chronic hyperglycemia slows down the metabolic rate but increases the sensitivity of K\(_{\text{ATP}}\) or Ca\(^{2+}\) channels (Kato et al 1996).

The timing of Zn\(^{2+}\) outflow recorded during the first stimulation with tolbutamide in lean mouse β-cells correlated with the timing of calcium response recorded during the second stimulation. We showed that β-cells repeat their calcium response pattern during consecutive stimulations with tolbutamide. It has also been found that cytoplasmic calcium changes and Zn\(^{2+}\) release are linked with regard to both timing and magnitude (Quan et al 2004, Wollheim and Pozzan 1984). We therefore conclude that the timing of insulin release during consecutive stimulations is cell-specific in cells from lean mice. No such correlation was found in ob/ob mouse β-cells.

β-Cells within intact islets showed cell-specific timing of their calcium responses. Fast responding β-cells therefore may have a pacemaker function. It was recently claimed that β-cells within an islet show identical temporal responses (Kuznetsov et al 2005). We could identify fast and slowly responding cells both in lean and ob/ob mouse islets. The variation was rather small in lean mouse islets, but in ob/ob mouse islets the range of lag-times was almost as large as in isolated β-cells. This could be due to a disturbed stimulus-secretion coupling or to a damage of cell contacts and may explain the disorganization of calcium oscillations found in ob/ob mouse islets (Ravier et al 2002). We cannot exclude that part of the variation in lag-time could be caused by the handling of the tissue during the islet isolation procedure. β-Cells may vary in their functional characteristics within pancreatic islets, because they originate not from a single stem cell but from a
number of different low-differentiated precursors (Lechner and Habener 2003). We have previously reported that intact islets, as a whole, show a specific response. Taken together, this indicates that both the cell specificity of each β-cell and the functional connection between cells is important for islet-specific behaviour and adequate insulin release.

In Paper V, we could now show that there is a correlation between mitochondrial mass before the stimulation and the timing of the β-cell cytoplasmic calcium response. This was done using Mitotracker Red and Fluo-3. There may be several mechanisms by which the mitochondrial state influences the timing of calcium response. Firstly, mitochondria are the main source of production of ATP, which plays a key role in pancreatic β-cell glucose signalling. Secondly, before the generation of ATP from the glucose molecule, ATP is required for glucose phosphorylation to yield glucose-6-phosphate and fructose-2,6-bisphosphate. It has been suggested that these reactions depend on sufficient β-cell ATP content prior to the glucose challenge (Ortsäter et al. 2002). As a consequence, a large amount of active mitochondria could enhance the glycolysis. Thus, a larger mitochondrial mass can provide a higher rate of both glycolytic and mitochondrial metabolism. It is difficult to say which metabolic step is more important with regard to determining the lag time for calcium rise. The role of glycolytic vs. mitochondrial metabolism in glucose-stimulated insulin secretion is controversial. Mertz et al. (1996) found that glycolytic ATP alone can sustain glucose-induced insulin release, but Ortsäter et al. (2002) found that glycolysis alone could neither initiate nor maintain it. Mitochondrial metabolism may stimulate insulin release also through other mechanisms (Maechler and Wollheim 2001, MacDonald et al. 2005).

The mitochondrial Ca$^{2+}$ uptake, following the rise in cytoplasmic Ca$^{2+}$ (Rutter et al. 1993) may activate mitochondrial metabolism (Ainscow and Rutter 2001, Rutter et al. 1996), providing a mitochondrial potentiation, which may have an important role in the response of islets to glucose. ATP is important for many processes in stimulated β-cells, such as sequestration of Ca$^{2+}$ before the [Ca$^{2+}$], rise (19), insulin granular movement and the exocytotic process (Wollheim et al. 1996). Decreased ATP levels result in opening of K$_{ATP}$ channels (Kohler et al. 1998). ATP is also suggested to be a secreted messenger in communication between β-cells (Hellman et al. 2004).

As our results from Paper III show, the cell specificity of mitochondrial metabolism is blunted in β-cells from hyperglycemic mice in a way that affects timing of response but not the magnitude. There was no correlation between mitochondrial mass and the magnitude of calcium response. As it was suggested above, it is possible, that other factors, such a number and opening state of voltage-dependent calcium channels have a large influence on the magnitude of calcium rise. There may also be a technical explanation for this. Changes in [Ca$^{2+}$], are indicated by changes in the absolute values for Fluo-3 fluorescence. Because of the dye degradation and leakage, such a system is more subject to experimental variation, when compared with ratiometric measurements, such as with fura-2. An existing correlation between mitochondrial mass and [Ca$^{2+}$], peak height may therefore be difficult to detect using Fluo-3.
According to electron microscopy studies, β-cells from 3-5-month-old ob/ob mice have fewer and larger mitochondria than lean mouse β-cells (Diani et al 2004). Other studies have not found such differences (Jörns et al. 1997). It has been suggested that β-cell dysfunction results from prolonged exposure to high glucose and elevated free fatty acid levels, leading to oxidative stress and apoptosis (Evans et al 2003). ob/ob Mice are hyperglycaemic, but we did not find differences in the localization of mitochondria or MitoTracker staining intensity when compared with β-cells from normoglycemic mice. As shown previously (Paper II and III), the timing of the Ca^{2+} response to glucose did not differ between β-cells from lean and ob/ob mice.

Calcium rise was less pronounced in the central cell areas, where most mitochondria were localized. This is not unexpected since calcium rise is caused by Ca^{2+} influx through plasma membrane channels. It supports the hypothesis that mitochondrial calcium uptake also has an impact on the cytoplasmic calcium signal (Duchen 2004). Thereby the activity of calcium-dependent processes may be partly regulated by mitochondria. It could also reflect that mitochondrial activation requires a smaller calcium increase than exocytosis does.

Voltage-dependent calcium channels are localized to specific membrane domains, so called “hot spots” (Qian and Kennedy 2001). We found no evidence that calcium increases occur in areas close to aggregates of mitochondria. Our data therefore suggest that hot spots are not connected to localised increases in ATP production.

Cells within the intact islet of Langerhans may function as a metabolic syncytium, secreting insulin in a coordinated and oscillatory manner in response to external fuel (Pipeleers 1987). Since the pacemakers in the islets are likely to be the cells with the shortest lag-times, there could be a possibility of identifying pacemaker cells within an islet on the basis of a larger mitochondrial mass. However, we saw no correlation between the MitoTracker intensity and the lag-times in most islets. With Fluo-3, we studied only the outer cells in the islet. For all we know, the signal propagation may not be radial (from a single source) and the source of propagation may be located near the surface of the islet. The range of lag-times between individual β-cells is very narrow in lean mouse islets, which makes it difficult to detect a correlation. In one ob/ob mouse islet, β-cells with a higher amount of active mitochondria responded to stimulation with a shorter lag-time. Synchronicity in ob/ob mouse islets is lower than in normal mouse islets. This may be attributed to an impaired cell coupling (Raviet et al 2002), but there is also a disturbed cell specificity of the response to mitochondrial stimulation.

High concentrations of MitoTracker (µM) may reduce mitochondrial respiration, but only a small effect was observed at the low concentration range (Buckman et al 2001). Other studies suggest that prolonged exposure to low concentrations of MitoTracker has little or no effect on mitochondrial function (Waters and Smith 2003).

Thus, we found that mitochondrial mass varied considerably between β-cells and that mitochondrial mass was related to the timing of calcium response in both lean and ob/ob mouse β-cells. The state of mitochondria before the stimulation
seems to determine the timing of secretory response and may be an important part of the explanation for the heterogeneity of β-cell response.

Normal β-cells show cell specific responses to all tested stimuli. β-Cells from hyperglycemic mice have cell-specific timing and magnitude of responses to glucose and glyceraldehyde, but they show no temporal cell specificity when stimulated with a mitochondrial substrate. This suggests an altered mitochondrial function. The loss of cell-specific timing of response also at the step of calcium channel opening suggests that the mechanism(s) for setting the individual response characteristics is impaired in diabetic mouse β-cells. Temporal cell specificity for all steps of the stimulus-secretion coupling may therefore be important for normal function. The timing of response in β-cells may be controlled by their mitochondrial state.
SUMMARY

1. The heterogeneity of β-cell responses is based on unique and reproducible characteristics of β-cells.

2. Cell specificity of the β-cell response is found in different rodent models. Aggregated β-cells also show cell-specific responses.

3. Normal β-cells show cell-specific Ca\(^{2+}\) responses when stimulated at different steps of the stimulus-secretion coupling. This means that not only the stimulus-secretion coupling, as a whole, is cell-specific but also different parts of it, such as mitochondrial metabolism, glycolysis and ion channel function.

4. β-Cells from hyperglycemic mice show cell-specific timing and magnitude of responses to glucose and glyceraldehyde but they lose cell specificity of the timing when stimulated with a mitochondrial substrate or with depolarizing agents. This indicates that the mechanism(s) for setting the individual response characteristics is impaired in diabetic mouse β-cells. This may be due to the altered functions of calcium channels and mitochondria. Cell specificity for all steps of the stimulus-secretion coupling may be necessary for normal β-cell function.

5. β-Cells with a larger number of active mitochondria react faster to stimulation. The lag-time for calcium rise may be determined by the mitochondrial state.
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