Small Mouse Islets Are Deficient in Glucagon-Producing Alpha Cells but Rich in Somatostatin-Secreting Delta Cells

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Small and big mouse islets were compared with special reference to their content of glucagon-producing \( \alpha \)-cells and somatostatin-producing \( \delta \)-cells. Areas stained for glucagon and somatostatin were measured in the largest cross section of small (diameter < 60 \( \mu \)m) and big (diameter > 100 \( \mu \)m) islets. Comparison of the areas indicated proportionally more \( \delta \)-than \( \alpha \)-cells in the small islets. After isolation with collagenase these islets were practically devoid of \( \alpha \)-cells. We evaluated the functional importance of the islet size by measuring the \( \text{Ca}^{2+} \) signal for insulin release. A majority of the small islets responded to the hyperpolarization action of somatostatin with periodic decrease of cytoplasmic \( \text{Ca}^{2+} \) when glucose was elevated after tolbutamide blockade of the \( K_{\text{ATP}} \) channels.

1. Introduction

Studies of the endocrine cells in pancreas face special problems dependent on their distribution into islets containing several types of hormone-producing cells. Postnatal growth of the endocrine pancreas is strictly regulated [1, 2]. Independent of age, the total volume of the endocrine pancreas is symmetrically related to the diameter of the islets in various species including mouse [3] and man [4, 5]. Accordingly, a majority of the endocrine cells are situated in the medium-sized and not in the numerous small islets [6, 7].

A key question is how the size of the islets is related to their content of different types of hormone-producing cells. Increase of glucose is a more effective stimulator of insulin release from small rather than from big islets [8]. This observation can be attributed to several factors, including a higher proportion of \( \beta \)-cells. Originally the glucagon-producing \( \alpha \)-cells were considered to be located essentially in small islets [9]. Today we know that a majority of the \( \alpha \)-cells in most species appear in the periphery of big and medium-sized islets [8, 10, 11]. Current knowledge of the cytoarchitecture of small islets is based on studies of islets with a diameter less than 100–150 \( \mu \)m [12, 13]. This size limit is less appropriate by also including medium-sized islets that account for most of the volume of the endocrine pancreas in rats [14], mice [3], and man [4, 5].

The aim of the present study was to compare the cytoarchitecture of small and big mouse islets in both sections of pancreas and after isolation with collagenase. The results indicate differences between small islets and big islets with regard to both proportions of \( \alpha \)-and \( \delta \)-cells and the \( \text{Ca}^{2+} \) oscillations induced by glucose in the absence of functional \( K_{\text{ATP}} \) channels.

2. Material and Methods

The experimental procedure was approved by the Animal Ethics committee of Uppsala University. Pancreas was removed from 3-4-month-old C57BL/6 mice allowed free access to food and water. The islets were studied either when situated in pancreas or after collagenase isolation followed by 1-2 days’ culture. The procedure of isolation was similar to that of Lacy and Kostianovsky [15]. Pancreases were fixed in 10% (vol/vol) formalin and embedded in paraffin. Isolated islets were fixed in 4% (vol/vol) paraformaldehyde followed by embedding in agarose and finally paraffin [16].
2.1 Immunohistochemistry. The paraffin-embedded tissue was consecutively sectioned (5 μm) and mounted on glass slides. Pancreas sections were double stained for glucagon (mouse monoclonal; Abcam, ab10988) and insulin (guinea pig polyclonal; Fitzgerald, Concord, MA, USA) followed by detection with MACH 3 polymer system (Biocare Medical, Concord, CA, USA) and developed with Vulcan Fast Red and 3,3′-diaminobenzidine (DAB), respectively [17]. Highly purified secondary antibodies (donkey anti-rabbit alkaline phosphatase (AP); donkey anti-guinea pig horseradish peroxidase (HRP); Jackson ImmunoResearch Laboratories, West Grove, PA, USA) were used to prevent cross-reactivity during double staining of pancreas with somatostatin and insulin, and development with Vulcan Fast Red and DAB, respectively.

Sections of isolated islets were either immunostained for glucagon or somatostatin followed by detection with MACH 3 polymer systems and developed with Vulcan Fast Red or DAB, respectively. The immunostained sections were counterstained with hematoxylin.

Previous observations on how the number and total volume of mouse islets relate to the islet diameter are shown in Figure 1(a). Referring to these data a comparison was made of the cellular composition in small (diameter < 60 μm) and big (diameter > 100 μm) islets. Areas of glucagon and somatostatin were measured in the largest cross section of the islets with a Leica LMD6000 microsystem (objective 40x). The experimental procedure made it possible to calculate the total areas of α- and δ-cells in islet sections but only to estimate their contributions to the islet volume. Observations in the largest cross section are more representative for small rather than big islets (Figure 1(b)). Even more important, peripherally located cells will contribute more to the islet volume in small rather than in big islets.

The functional importance of the islet size was evaluated by studying the glucose generation of cytoplasmic Ca\(^{2+}\) oscillations mediated by the K\(_{ATP}\) channel-independent pathway. The concentration of cytoplasmic Ca\(^{2+}\) was measured with ratiometric fura-2 technique as previously described [18].
2.2. Statistical Analysis. All values are given as means ± SEM. Probabilities of chance differences between the groups were calculated using Student’s unpaired two-tailed t-test. For all comparisons, a P value < 0.05 was considered statistically significant.

3. Results

The topographic distribution of α- and δ-cells in small and big islets is illustrated in Figures 2 and 3. Both types of cells occurred in the islet periphery, often forming a mantle around a core with β-cells. In the big islets, δ-cells were located close to the capillaries also in the central core. The peripheral location was particularly pronounced for α-cells, whereas the δ-cells were intertwined between β- and α-cells. The difficulties to evaluate the proportions of α- and δ-cells in islets of different sizes are illustrated in Figure 1(b). The approach with measurements in the largest optical cross section will underestimate the number of α-cells in big islets due to their peripheral location. Nevertheless, the glucagon-positive area was twice as large (P < 0.001) in big compared with the small islets in pancreas sections (Table 1). The opposite was found for the area of somatostatin, which was larger in small rather than in big islets (P < 0.05).

Results from measurements in isolated islets are included in Table 1. In the big islets the percentage contribution of the glucagon area was 1.43 ± 0.28 and the small islets were practically devoid of α-cells. In the latter case, only 2 of 52 islets were positive for glucagon. A corresponding analysis revealed that the somatostatin area was larger in small rather than in big islets (P < 0.001).

Attempts were made to decide whether the oscillatory Ca²⁺ signal for glucose-induced insulin release was affected by the islet size. It was found that hyperpolarization with 100 nM somatostatin usually resulted in glucose-dependent Ca²⁺ oscillations in small islets after sulfonylurea blockade of the K_ATP channels (Figure 4). The oscillations appeared not only in the presence of 20 mM glucose but also at 8 or 5 mM. Taken together, the exposure to somatostatin resulted in glucose-dependent oscillations in 89% (42 of 47) of the small and 31% (4 of 13) of the big islets (P < 0.05).

4. Discussion

A reliable but tedious way to study the cellular composition of islets is to measure the area of different types of cells in consecutive thin sections. In the present study the areas stained for glucagon and somatostatin were determined in the largest cross section of the islets. This procedure makes it possible to get a sufficient number of data from small and big islets located in pancreas or isolated with collagenase. Including both the central core and the peripheral mantle of the islets, the approach provides an indirect estimate of the whole islet. It is important to note that the data presented underestimate the contribution of the peripherally located cells, essentially α-cells.
Figure 3: Double immunostaining of a big islet (a) and a small islet (b) in the pancreas for somatostatin (red) and insulin (brown). Immunostaining of an isolated big islet (c) and an isolated small islet (d) for somatostatin (brown). The somatostatin-producing δ-cells are located in both the core and periphery of the islets. Scale bars represent 50 μm.

Table 1: Proportions of α- and δ-cells in small and big islets. Areas stained for glucagon and somatostatin in the largest cross section of small (<60 μm) and big (>100 μm) islets. The data refer to observations made in paraffin sections of pancreas and isolated islets. All values are given as means ± SEM. *P < 0.001 compared with small islets stained for glucagon in the pancreas. †P < 0.05 compared with small islets stained for somatostatin in the pancreas. ‡P < 0.05 compared with small isolated islets stained for glucagon. §P < 0.001 compared with small isolated islets stained for somatostatin.

| Islets in pancreas | Isolated islets |
|-------------------|-----------------|
|                   | Small islets    | Big islets | Small islets | Big islets |
| Glucagon          |                 |           |              |            |
| Number of animals | 5               | 5         | 6            | 4          |
| Number of islets  | 52              | 65        | 52           | 44         |
| Area stained for glucagon (%) | 3.22 ± 0.51 | 8.61 ± 0.50* | 0.1 | 1.43 ± 0.28† |
| Somatostatin      |                 |           |              |            |
| Number of animals | 5               | 5         | 4            | 5          |
| Number of islets  | 68              | 51        | 35           | 50         |
| Area stained for somatostatin (%) | 4.60 ± 0.48 | 3.37 ± 0.22‡ | 5.71 ± 0.58 | 3.54 ± 0.24§ |

The size distribution of the islet volume is essentially symmetrical in birds [19], rodents [3, 4, 14], and man [4, 5]. A major part of the endocrine mouse pancreas is insulin-producing β-cells, the number of which usually increases when required for glucose homeostasis [20]. In the obese-hyperglycemic syndrome the increase results in 10 times more β-cells with maintenance of the symmetrical volume relation between the big and small islets [3]. In human diabetes the symmetry is maintained [21] despite a reduced islet mass [22]. The present study indicates that the glucagon-positive area is more than twice as large in big compared with small islets situated in the pancreas. Further evidence that big islets contain more α-cells was obtained from studies of islets isolated with collagenase and kept in culture for 1-2 days. In this case the small islets were practically devoid of α-cells but rich in δ-cells. The reason why isolation of islets results in a
Figure 4: Effects of 100 nM somatostatin on the cytoplasmic Ca$^{2+}$ concentration in one big and two small islets after elevation of glucose in the presence of 1 mM tolbutamide. Exposure to somatostatin resulted in glucose-dependent oscillations mediated by periodic interruption of elevated Ca$^{2+}$ in 42 of 47 of the small islets and in 4 of 13 of the big islets.

lower content of α-cells is open for discussion. It is possible that the peripheral location makes the α-cells particularly vulnerable to the isolation process [23].

Somatostatin-producing δ-cells are topographically related to insulin-producing β-cells in several species including rodents [19]. We now confirm the reports that δ-cells are located not only in the periphery but also close to the capillaries penetrating into the central part of the big islets [20, 24]. The area stained for somatostatin is proportionately larger in small rather than in big mouse islets. The difference was particularly pronounced after collagenase isolation.

The finding that small islets are rich in somatostatin-producing δ-cells but practically devoid of α-cells raises the question how this affects the oscillatory Ca$^{2+}$ signal for insulin release. Periodic entry of Ca$^{2+}$ can be generated not only by elevation of glucose followed by closure of K$_{ATP}$ channels and depolarization. In the K$_{ATP}$ channel-independent alternative, the high membrane resistance makes the β-cells sensitive to hyperpolarization resulting in oscillations due to periodic interruption of the entry of Ca$^{2+}$. Somatostatin is one of several G-protein coupled agonists known to hyperpolarize β-cells [25]. We now report that exposure to somatostatin in the presence of sulfonylurea blockade of the K$_{ATP}$ channels promotes glucose-dependent oscillations more often in small rather than in big islets. Similar effects were seen after hyperpolarization obtained by activation of the α$_{2}$-adrenergic receptors (unpublished data). Kilimnik et al. [22] have reported that the proportion of α-cells is higher in large rather than in small islets also in humans. Moreover, these authors observed preferential loss of large islets in patients with type-2 diabetes. Accordingly, the reduction of islet mass in diabetes results in a proportionally larger decrease of glucagon-producing α-cells than of β- and δ-cells, a situation supposed to counteract hyperglycemia [26]. Our observation that small islets are prone to oscillate in response to glucose provides additional support for the clinical importance of the islet size.

5. Conclusions

Big islets situated in the pancreas contain proportionally more α-cells than small islets. In contrast, small islets situated in the pancreas contain proportionally more δ-cells than big islets. This difference was even more pronounced in isolated islets. Moreover, isolated small islets are practically devoid of α-cells but rich in δ-cells. Our Ca$^{2+}$ measurements revealed
that exposure to somatostatin in the presence of sulfonylurea blockade of the $K_{ATP}$ channels promoted glucose-dependent oscillations more often in small than in big isolated islets.

**Abbreviations**

ATP: Adenosine triphosphate  
DAB: 3,3’-Diaminobenzidine  
AP: Alkaline phosphatase  
HRP: Horseradish peroxidase.

**Competing Interests**

The authors of this paper have no conflict of interests to disclose.

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