Histidyl–Proline Diketopiperazine (His–Pro DKP) Immunoreactivity Is Present in the Glucagon-containing Cells of the Human Fetal Pancreas

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Abstract

Histidyl–proline diketopiperazine (His–Pro DKP) cells in the pancreas of human fetuses aged between 12 and 19 wk were localized by the indirect antibody–enzyme method on semithin sections. To study their fine structure, two techniques were used: a superimposition technique consisting of comparison of the same cells in semithin and electron microscopic preparations, and an immunocytochemical technique on ultrathin sections using the unlabelled antibody peroxidase–antiperoxidase method. Our results show that (a) the same cells are positive for both His–Pro DKP and glucagon/glicentin, (b) His–Pro DKP immunoreactive cells possess extremely electron-opaque secretory granules, implying that these cells correspond to the A cells, and (c) His–Pro DKP immunoreactivity is found over the secretory granules. We hypothesize that the two peptides His–Pro DKP and thyrotropin-releasing hormone (TRH) have independent origins, since TRH is found in the B cells.

Introduction

Histidyl–proline diketopiperazine (His–Pro DKP), a cyclic dipeptide initially described in both the hypothalamus and cerebral tissues (1–4), has been detected in various sites outside the central nervous system (5, 6), including the gastrointestinal tract and, more specifically, the endocrine pancreas (7). There has been some controversy over the relationship of endogenous His–Pro DKP to thyrotropin-releasing hormone (TRH). Although it was initially proposed that His–Pro DKP was a metabolite of TRH (8–11), recent studies suggest that most of His–Pro DKP is independent of the tripeptide amide (12, 13). Pancreatic His–Pro DKP appears to be chromatographically and immunologically identical to the synthetic dipeptide (7), but its role is unknown. The cyclic dipeptide is not localized in the insulin-containing cells, since His–Pro DKP content, unlike TRH (14), remains unchanged after the selective destruction of B cells by streptozotocin (7).

In the present experiment we examined the localization of His–Pro DKP immunoreactivity in the human fetal pancreas by light and electron microscopic immunocytochemistry. This model offers a number of potential advantages in clarifying the origin of His–Pro DKP. While pancreatic TRH is easily detected in the fetal and early neonatal period in the rat, the levels gradually decline during the neonatal period and are basely detectable in the adult (15–18). In contrast, the content of His–Pro DKP in the pancreas peaks in the adult rat (12). In light of preliminary studies demonstrating the simultaneous occurrence of both insulin and TRH immunoreactivity in the same islet cells of the human fetal pancreas (19), we have used this system to further clarify the cellular location of His–Pro DKP and its relationship to TRH.

Methods

Tissue collection. Eight pancreatic glands were collected from human fetuses (six females and two males) delivered after legal abortions. The crown-rump length of the fetuses ranged from 6 to 14 cm (i.e., 12–19 wk of gestation) (20). Our study was carried out according to guidelines established by the National Ethical Committee of France.

Tissue preparation. To preserve His–Pro DKP immunoreactivity, a combination of p-formaldehyde (4% wt/vol) and low glutaraldehyde (0.5% vol/vol) was used in the primary fixation step, followed by postfixation with 1% osmium tetroxide as reported for TRH (19). After embedding in araldite, semithin 1-μm and ultrathin 90-nm sections were cut. Semithin sections were dehydrated for 10–30 min with sodium ethoxide (21). Semithin or ultrathin sections of postosmicated tissue were then pretreated with a strong oxidizing agent (as described below) before processing for immunocytochemistry.

Pretreatment of the postosmicated tissue. Before labeling, semithin or ultrathin sections were treated in three different ways: (a) incubation for 10 min with 0.3–10% hydrogen peroxide solution (22); (b) incubation for 10 min with 1–5% periodic acid solution (23); and (c) for 120 min with saturated sodium metaperiodate (24).

Primary antisera. Antiserum to synthetic His–Pro DKP (Peninsula Laboratories, Inc., Belmont, CA) (No. 1068) was raised in a New Zealand White rabbit immunized with the peptide coupled to bovine thyroglobulin by the bis-diazotized benzidine procedure as reported for TRH (25). This antiserum exhibits no cross-reactivity with its constituent amino acids, pGlu-His, TRH free acid, or gonadoliberin in radioimmunoassay, and ~0.1% cross-reactivity with TRH (13). For immunocytochemistry, the antiserum was used at a titer of 1:500.

Antiserum to pure porcine pancreatic glucagon (Novo Research Unit, Copenhagen, Denmark) was prepared at the Unité 264, Montpellier, France. This antibody (No. GAN 8) was obtained from a New Zealand White rabbit inoculated with the peptide conjugated to bovine serum albumin with carbodiimide (26). Scatchard analysis indicates an equilibrium constant of 6.1011 liters/mol and a binding capacity of 638 nmol/liter. The specificity of the antiserum was tested by adding 0.01–100 nmol TRH, His–Pro DKP, human glucagon-like peptides GLP1 and 2, rat pancreatic polypeptide (PP), peptide histidine-isoleucine, peptide histidine–methionine (PHM), peptide tyrosine–tyrosine (Peninsula Lab-
oratories, Inc.), bombesin, neurotensin, somatostatin (Bachem, Bubendorf, Switzerland), secretin, vasoactive intestinal peptide (a gift from Dr. E. Wunsch, Max Planck Institute, Munich, Federal Republic of Germany), synthetic cholecystokinin (a gift from Dr. J. Martinez, Unité 264, Montpellier, France), and rat insulin (Novo Research Institute).

None of these peptides demonstrated cross-reactivity with the antiserum (Fig. 1). The relative affinities for COOH-terminal glucagon fragment F 13–26 obtained by endopeptidase digestion and for porcine oxytocinomodulin G-37 (G-37) (supplied by Dr. D. Bataille, Unité 264, Montpellier, France) were 0.1 and 2%, respectively. Dilutions of a crude rat intestinal extract shared the same pattern of cross-reactivity as G-37 (Fig. 1). For immunocytochemistry, the anti-hormone serum was used at a titer of 1:5,000.

The following specific rabbit primary antisera were also used: anti-insulin No. 24-2 (kindly donated by Dr. R. Assan, Hôpital Bichat, Paris, France) diluted 1:5,000; anti-somatostatin No. 19608 (kindly donated by Dr. M. P. Dubois, Institut National de la Recherche Agronomique; Nouzilly, France) diluted 1:1,000; anti-bovine PP No. 146-6 (kindly donated by Dr. R. E. Chance, Lilly Research Laboratories, Indianapolis, IN) diluted 1:5,000; and anti-glicentin No. R-64 (kindly donated by Dr. A. J. Moody, Novo Research Institute) diluted 1:5,000. Full characterization of these primary antisera have been reported elsewhere (27–32).

Immunocytochemistry on semithin sections: light and electron microscopy. The immunocytochemical procedure performed on serial 1-μm sections using the goat anti-rabbit immunoglobulin G conjugated to peroxidase (33) was applied as described previously (30). Semithin sections were examined in a Zeiss light microscope. For electron microscopy, immunoreactive cells were photographed and the corresponding ultrathin sections of the same cells, contrasted with lead citrate, were examined in an Elmskop 101 electron microscope. The mean values (in nanometers) of granule profile diameter were characterized with the aid of the 95% confidence limits (mean±1.96 [SD]/√n).

Immunocytochemistry on ultrathin sections. The unlabeled antibody peroxidase-antiperoxidase method (34) was applied as described previously (22). The grids were contrasted with lead citrate before observation.

Primary antisera specificity controls. To test for specificity, antiserum to His-Pro DKP was absorbed with the corresponding antigen (Peninsula Laboratories, Inc.) at concentrations of 10–1,000 nmol/ml undiluted antiserum for 1 h at 37°C. Absorbed antiserum or nonimmune serum were used instead of freshly diluted primary antiserum on control sections. All staining was abolished by absorption of primary antiserum at 250 nmol. Absorption of the anti-His-Pro DKP with heterologous hormones (pancreatic and brain peptides) or hormone fragments (also see reference 19) gave no change in the reaction. The immunoreactivity due to anti-COOH-terminal–specific glucagon, anti-glicentin, anti-insulin, anti-somatostatin, or anti-PF was quenched by the corresponding antigen (also see reference 19).

Results

Technical aspects of His−Pro DKP immunostaining. Among the different oxidizing agents tested, sodium metaperiodate was found to be the most suitable, giving dense labeling over perfectly preserved structures (Figs. 2–4). The restoration of His−Pro DKP immunoreactivity with sodium metaperiodate was found to be time dependent: after 10 min of treatment on deresinated semithin sections, intense labeling was observed, but unsatisfactory results were obtained after 20 min. On ultrathin sections, maximal labeling intensity was reached between 30 and 60 min. A slight decrease in immunostaining occurred when 1–5% periodic acid was used, whereas the least reactivity was obtained with 0.3–10% hydrogen peroxide.

Location of His-Pro DKP immunoreactive cells. Antiserum to His-Pro DKP stained endocrine-type cells in the pancreatic gland of 12–19-wk-old human fetuses. The immunoreaction was present in all the pancreatic islets of the female and male fetuses investigated. At the light microscope level, immunoreactive cells were interspersed among the islet cell clusters at the 12th week of gestation (Fig. 2 A), and later (16 wk) inside the recognizable islets of Langerhans (Fig. 2 C). His-Pro DKP immunoreactivity was entirely confined to the peripheral parts of the typical islets (Fig. 2 C). It was present throughout the pancreatic gland, but its distribution was heterogeneous. Immunoreactive cells were numerous in the superior part of the head, in the body and in the tail, but were only very scarce in the lower posterior part of the head. The exocrine tissue of the fetal pancreas was unaffected. Also, no nerve cells or fibers within the pancreas contained His-Pro DKP immunoreactivity.

Interrelationship of cells immunoreactive to His-Pro DKP and other hormone antisera. Consecutive semithin sections of 12–19-wk-old fetuses were stained alternately with His-Pro DKP and COOH-terminal–specific glucagon antisera. As shown in Fig. 2, the same cells in the islet cell clusters (12 wk) or in the typical islets (16–19 wk) revealed the simultaneous occurrence of both His-Pro DKP and COOH-terminal glucagon immunoreactivity. Similar results were obtained after substitution of glicentin antisera (that specifically recognizes the glucagon precursor; also see reference 35) in place of the COOH-terminal–specific glucagon antisera. In contrast, the His-Pro DKP immunoreactive cells were distinct from insulin- and somatostatin-containing cells in the superior part of the head, in the body, and in the tail. No His-Pro DKP immunoreactivity could be detected in the numerous cells containing pancreatic polypeptide of the lower posterior part of the head.

Ultrastructural characterization of His-Pro DKP immunoreactive cells. A superimposition technique, using semithin sections for immunocytochemistry and consecutive ultrathin sections for ultrastructural characterization, were then adopted. His-Pro DKP immunoreactive cells observed in the consecutive ultrathin sections all shared a number of characteristics. They possessed a population of extremely electron-opaque secretory granules with two components: an eccentric spherical core and a granular mantle, the latter usually less dense (Fig. 3). The granule profile diameter varied considerably within a given cell and between different cells, and ranged from 250 to 600 nm; their size distribution is shown in Fig. 5. The mean value (nanometers±confidence limits 95%) of the profile diameter was 416±24.14. His-Pro DKP immunoreactive cells corresponded to the A cells. As shown in Fig. 3, there was another cell type with granules ~300–600 nm in diameter, but whose granules dis-
played a paracrystalline core and a broad halo. This cell type was referred to as B cells. As previously reported (19), the mean profile diameter (nanometers±confidence limits 95%) was 442±5.04. The D cells, in contrast, contained large granules (~300–950 nm in diameter) with clear contents surrounded by a broken limiting membrane (Fig. 3). The mean profile diameter (nanometers±confidence limits 95%) was 716±7.78. The B and D cells were not stained on semithin sections (Fig. 3).

**Ultrastructural localization of His-Pro DKP.** Immunocytochemical staining with His-Pro DKP antiserum gave a precise localization of the cyclic dipeptide in the cytoplasmic granules. The reaction product (electron-dense precipitates at least 30 nm in diameter) was deposited on electron-opaque secretory granules, leading to the conclusion that these cells corresponded to the A cells (Fig. 4). The reaction product usually stained the whole surface of the secretory granules (Fig. 4), as reported for glucagon (19).

**Discussion**

His-Pro DKP has been measured biochemically in the pancreas and, more specifically, in the islets of Langerhans, however its precise localization was unknown except that the cyclic dipeptide is not concentrated in streptozotocin-sensitive cells (7). As reported herein, its cellular location in the A cells revealed by immunocytochemistry may be taken as indicative of a potential relationship to other hormones and of functional relations between His-Pro DKP and the endocrine pancreas.

In the present paper, we have reported the presence of a strong His-Pro DKP immunostaining in the human fetal pancreas. Its cellular location has been assigned to an endocrine-type cell, predominantly confined to the periphery of islets. A general problem with immunocytochemical methods is the specificity of the technique, especially the possible existence of cross-reacting antibodies. This is a particularly important issue when using antisera raised to small peptides, such as the cyclic dipeptide His-Pro DKP. The radioimmunoassay and absorption tests suggest that each primary antiserum is not only specific to one peptide family but that, within its family, it is also specific to the peptide that has produced the antiserum.

Three techniques were used to characterize its localization: (a) consecutive semithin sections alternately stained with His-Pro DKP and other antisera; (b) superimposition of immunocytochemical and electron microscopic observations on the same cells; and (c) immunocytochemical staining on ultrathin sections.

![Figure 2](http://www.jci.org) Developmental pattern of His-Pro DKP/glucagon immunoreactive cells. Consecutive semithin sections stained with His-Pro DKP (A, C) and COOH-terminal-specific glucagon (B, D) antisera. (A, B) 12-week-old human fetus: the His-Pro DKP/glucagon cell reacting both to His-Pro DKP and COOH-terminal-specific glucagon antisera (arrows) is present in the islet cell clusters. ×1,500. (C, D) 19-week-old human fetus: the His-Pro DKP/glucagon cell stained by both antisera (arrows) is present in the peripheral part of the islets of Langerhans. ×1,100.
The first technique established whether different peptides were present in the same cells. His-Pro DKP immunoreactivity was seen exclusively in glucagon immunoreactive cells. Second, the superimposition technique permitted an excellent description of ultrastructure of individual cells of each type. With this method, we were able to describe the His-Pro DKP immunoreactive cells as cells with electron-dense granules ranging in size from 250 to 600 nm. This cell type was referred to as A cells (36, 37). With the third technique, more information was gained about the subcellular localization of His-Pro DKP. The immunoreactivity was found over all the secretory granules.

Morphological techniques have previously established the developmental profile of the different endocrine pancreatic cell populations and, more specifically, they have allowed identification of A cells in early fetal life (8 wk) (38). By radioimmunoassay, it has been documented that glucagon and glucagon-like immunoreactants are detected at the end of the 8th week with specific (COOH terminal) and nonspecific (NH₂ terminal)
glucagon antiserum, respectively (39). Using immunocytochemical techniques, it has been observed that glucagon-containing cells are present from 8 wk; the occurrence of glucagon immunoactivity is accompanied by glicentin (one glucagon-like immunoreactive; also see reference 35) immunoreactivity (30, 40). At present it is not known whether this fetal glucagon/glicentin cell is a cell type different from the His-Pro DKP cell producing glucagon/glicentin, or whether it represents a transitional stage in the differentiation of the pancreatic A cell. However, we have noted that the developmental pattern of His-Pro DKP is identical to that of glucagon/glicentin between 12 and 19 wk; moreover, His-Pro DKP is restricted to the lobe rich in glucagon/glicentin, suggesting that such a peculiar distribution could be related to an embryological origin from the dorsal primordium (28). Because true pancreatic-type glucagon has been traced to the glicentin-containing cells in the fetal (41, 42) and adult (43–45) human gastrointestinal tract, it will be interesting to check whether the intestinal L cell also contains His-Pro DKP.

The eventual implications of His-Pro DKP in the development of the pancreatic gland of the human fetus are open to discussion. It has been suggested that His-Pro DKP could be a metabolic breakdown product of TRH (8–11). Because the putative parent tripeptide TRH has been assigned to the pancreatic B cells in neonatal (46), streptozotocin-treated newborn (14), and hypothyroid (32) rat, and in the human fetus (19), we further hypothesize (12, 13) that the two peptides His-Pro DKP and TRH have independent origins. Recently the presence of His-Pro DKP immunoreactivity in the mammalian cerebrospinal fluid has been reported (47). This study also suggests that at least a certain portion of His-Pro DKP may be derived from sources independent of TRH. There is, however, one report of pancreatic TRH in some of the cells reactive to glucagon antiserum in developing rat pancreas (46). Unfortunately, absorption tests with the His-Pro DKP antigen were not carried out in that study. One report has also indicated a possible role in the modulation of islet hormonal responses to nutrients, and in the catecholaminergic control of islet secretory events (7). If the co-localization of His-Pro DKP and glucagon is confirmed by other techniques, His-Pro DKP may play a role in glucagon physiology. However, it is not inconceivable that other roles may exist, specifically related to the regulation of differentiation and/or growth of fetal cells.

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