Pancreatic polypeptide: a review of its involvement in neuro-endocrine reflexes, islet-acinar interactions and ethanol-evoked physiopathologic pancreatic gland changes

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Summary

This review was prompted by the unexpected experimental finding in canines that Tissucol-induced pancreatic ductal blockade elicits Pancreatic Polypeptide (PP) release and seems to be at the basis of the beneficial effects on taurocholate-induced acute pancreatitis (AP). In the release mechanism of this regulatory peptide secreted by PP cells located in the periphery of Langerhans islets and scattered in the ductal epithelium, two neuroendocrine reflexes (NER) are involved. The "short" NER is evoked from the duodenum by an unknown component of bile-pancreatic secretion. The "long" NER is triggered by a vagovagal reflex. PP induces a depression of the intrapancreatic cholinergic tone. On the one hand suppressing, hormonally, nervous impulses discharge from the vagal nuclear complex in the brainstem. On the other, interfering paracrinically on the cholinergic transmission by acting, presynaptically, on post-ganglionic cholinergic neurons. The resulting PP-evoked fall of the intrapancreatic cholinergic tone depresses the hormone induced (secretin, CCK) pancreon’s secretory response. PP, with other agents, contributes to the “fail-safe” system or pancreon’s brake that prevents, in pancreocytes, the evolving of a “supramaximal-ecbolic-stimulation” process. The PP involvement as a modulating agent of pancreon’s reactivity is reflected by the progressive increment of its plasma values in the first week of an evolving AP episode. In the AP associated to a large meal, an overpowering of the “pancreon’s brake might have a pivotal role. In experimental and clinical chronic alcoholism, a vagal neuropathy of the Pavlov inhibitory fibers that, as a consequence, impairs the pancreon’s brake through a depression of PP secretion is at the basis of an enhanced reactivity of the duodeno-pancreatic reflexes. The latter leads to intrapancreatic cholinergic hypertonus and to Vater papilla’s dysfunction. These changes, plus an enhanced pancreocyte’s response to CCK, are at the core of acinar cell "supramaximal stimulation" with the organelle disruption that process implies. The intrapancreatic cholinergic hypertonus, the enhanced exocrine cell reactivity to CCK stimulation, and the augmented resistance to the pancreatic secretion flow at Oddi sphincter, explain the aggravating influence of chronic alcoholism on an episode of acute biliary pancreatitis. As the PP secretion, normally elicited by secretin, CCK, food and insulin hypoglycemia, is depressed in the presence of an augmented number of PP cells, as it is in the cases of chronic alcoholics, cystic fibrosis patients and, also, in dogs with pancreatic fibrosis (ductal ligation), it has been inferred, besides our postulated impairment of the Pavlov inhibitory fibers in the vagus nerves, that the defect of PP release is localized to the common final pathway of the above stimuli, probably in or near the PP cell itself.

Key words. Pancreatic polypeptide, neuro-endocrine reflexes, islet-acinar interactions.
Polipéptido pancreático: revisión de su ingerencia en los reflejos neuro-endocrinos, en la interacción acino-islote y en cambios fisiopatológicos de la glándula pancreática provocados por el etanol

Resumen

El péptido regulador PP (polipéptido pancreático) es secretado por células insulinares PP, principalmente en islotes del "processus uncinatus" ("gancho") del páncreas (esbozo ventral de la glándula) y, también, por células "dispersas" en el epitelio glandular. En su mecanismo secretor se hallan involucrados dos reflejos neuroendocrinos (RNE): uno "cor-toc" (RNE-C) y otro "largo" (RNE-L). El primero es desencadenado desde el duodeno por un componente, aún no precisado, de la secreción biliopancreática. El del segundo, se centra en un arco reflejo vago-vagal que da lugar a una depresión del tono colinérgico intrapancreático. Éste, por vía humoral, frena la descarga de impulsos parasimpáticos por parte del "complejo nuclear dorsal del vago". Una consecuencia bien establecida de lo precedente es la caída del rol permissivo ejercido por el tono parasimpático sobre los efectos secretorios en los "pancreones" por parte, sobre todo, de la hormona secretina. El PP, probablemente en conjunción con otros agentes, ejerce un rol de "freno" que trata de prevenir en los pancreocitos acinares el desencadenamiento de un proceso de "estimulación ecbólica supramáxima". Su rol modulator, lo demuestra, a través del incremento de su nivel en plasma en la primera semana de un episodio de pancreatitis aguda. En esta última entidad, subsecuentemente a una "comida copiosa", el sobrepaso de la capacidad de "freno" de la glándula tendría una influencia fisiopatogénica clave. En el alcoholismo crónico, tanto experimental como clínico, una neuropatía vagal que compromete a las "fibras inhibitorias" de Pavlov es el fundamento que explica, teniendo por pivote la hormona secretina, CCK, alimentos y la hipoglucemia insulínica, en la condición antes esbozada se halla deprimida; ello, paradójicamente, en presencia de un aumento en el número de su célula endocrina. Esto se constata en pacientes alcohólicos crónicos, en aquellos con fibrosis quística y también en perros con ligadura ductal. Se ha inferido que en todas estas circunstancias aparte de un compromiso de las fibras inhibidoras vagales de Pavlov ("freno pancreal"), se acopla, sea próximo y/o en la propia célula efectora, un defecto en la vía final común de los estímulos antes descriptos. Una concatenación de modificaciones neuroendocrinas muy próximas a las precedentes es la que se comprueba en pacientes con insuficiencia renal crónica. Éstas logran ser mimeticizadas experimentalmente en la rata efectuando la exéresis de un 85% de la masa renal. Un hallazgo a enfatizar es el que se constata en diabéticos insulinodependientes. En esta entidad, como en el caso del alcoholismo crónico, cabe considerar una perturbada transmisión de impulsos nerviosos siguiendo el trayecto de las fibras inhibitorias vagales de Pavlov. El consecuente hipertono colinérgico intrapancreático, y quizás también hepático, este último favorecedor secretorio del factor HISS de Lautt, brindan una explicación coherente de las llamadas "curvas planas" que se aprecian en estos pacientes al efectuar una curva de tolerancia a la glucosa; también que se constaten, con cierta frecuencia, "crisis hipoglucémicas" tardías siguiendo a una ingesta de carbohidratos.

Palabras claves. Polipéptido pancreático, reflejo neuroendocrino, eje insulo-pancreal.
and of ostensible signs of an inflammatory response in both pancreatic segments: i.e., Tissucol-blockade lobe (retrogastric) and the uncinate process.

These observations induced us to postulate the hypothesis that the beneficial effects on experimental AP produced by the Tissucol filling of the ductal tree were not the consequence of a simple mechanical blockade but the result of a complex chain of reactions, consisting of the activation of neuro-endocrine reflexes (NER) and changes of the normal islet-acinar interactions. In this whole process, the regulatory peptide Pancreatic Polypeptide (PP) seemed to play a pivotal role. Indeed, in the last series of tests of the Torino group, a remarkable observation, detected at the level of the portal vein, was a six to ten fold increase of the PP plasma values. This, in contrast to the unmodified plasma values of bombesin, enkephalins and somatostatin. The foregoing findings prompted us to hypothesize that PP was at the center of both "long" and "short" NER, and that it played a significant role in the complex endocrine-exocrine interactions. Through both mechanisms, PP might be able to attenuate the AP lesions, the beneficial effects probable linked to a reduced pancreozymin secretory activity. The latter probably as the consequence of a PP-evoked depression of the intrapancreatic cholinergic tone blocking of CCK effects and inhibition of acinar cells enzymes exocytosis. (Figure 1, 2 and Table 1)

The adrenergic component of the autonomic nervous system also influences PP secretion. Thus, beta-blockade inhibits PP release. In contrast, alpha-blockade enhances PP secretion. The latter might be related to the raised PP plasma values observed after celiac ganglioneutectomy. The latter finding seems to provide a new support, as therapeutic measure, to the local anesthesia of this autonomic structure in cases of an AP episode.

**Figure 1. Schematic Representation of the Long Neuro-Endocrine Reflex.** a. Vagus Nerve with its afferent and efferent fibres. b. Inhibitory hemiloop of the reflex arc that blocks the vagal nucleus discharges.

**Figure 2. Schematic Representation of the Short-Neuro-Endocrine Reflex.** a. Triggering in the duodenum by an unknown bile-pancreatic secretion component of PP from the PP cells; b. Preganglionic fibre; c. Intrapancreatic cholinergic neuron; d. Pancreon unit; e. Blocking of Ach discharge by pancreatic polypeptide at presynaptic level.

**Table 1.** Main features of PP involvement in the physiology of exocrine pancreas.

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<thead>
<tr>
<th>Pancreatic Polypeptide (PP)</th>
<th>In the physiology of the Pancreatic Gland</th>
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<td>PP: Pivotal member of the “Fall-State” or “Brake” Neuro-Endocrine System that controls “Pancreon” Units Function</td>
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<td>&quot;Short&quot; and &quot;Long&quot; Neuro-Endocrine Reflexes</td>
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<td>Islet-Acinar Interactions (Insul-Pancreon-Axis)</td>
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Modulation of the Intrapancreatic cholinergic tone

Prevention of "Pancreon" supranormal Ectopic Stimulation

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PP is also involved in the interactions that evolve within the insulo-pancreo-axis interactions. Firstly, PP might exert a local trophic effect on exocrine pancreas, protecting the PP-rich lobe of the gland (uncinate process) from atrophy in diabetic patients. Secondly, following a meal, as the increasing insulin stimulates digestive enzyme synthesis in the peri-insular cells, PP, together with somatostatin, might inhibit immediate exocrine release of stored enzymes.

Preliminary observations, unpublished Torino findings in rats, of a significant drop in mortality rate of taurocholate-induced AP following the infusion of PP seems to warrant the performing of further tests in order to elucidate whether or not this regulatory peptide is, in fact, a valuable preventive and/or therapeutic agent in acute inflammatory episodes of the pancreatic gland.

**Main features of PP release and action**

PP is a 36-aminoacid polypeptide, structurally related to peptides that include peptide YY (PYY) and neuropeptide Y (NPY).

Pancreatic polypeptide is a hormone released from the "F" or "PP" cells. These are normal constituents of the Langerhans islets, precisely of those derived from the ventral pancreatic bud. Because of this singular embryonic derivation, expression of PP immunoreactivity has been proposed as a marker of the ventral pancreatic bud origin. The distribution of the PP cells in the pancreatic gland is inhomogeneous. They are located predominantly in the periphery of the pancreatic islets. A minority are found scattered among the cells of both the acinar and centroacinar-ductal segments of the pancreatic unit.

As we have already pointed out, PP cells develop solely in the ventral pancreatic bud during early embryogenesis. At week 5-7 post-gestation, complex morphologic events take place. The ventral bud rotates around the embryonic foregut to occupy its final position as the posterior pancreatic head and uncinate process. In the adult pancreas, over 90% of the PP cells are located on the posterior portion of the pancreatic head, an area that corresponds to only 15% of the entire pancreatic volume.

PP secretion is stimulated by "chew and spit" sham feeding. This response is abolished by vagotomy and atropine. Hypoglycemia, whether induced by insulin or tolbutamide, strongly stimulates PP secretion. But the most powerful stimulus for the release of this regulatory peptide is acetylcholine (Ach). It should also be pointed out that peptides, such as CCK, bombesin and neurotensin may activate receptors on nerve cells and/or fibers and stimulate the PP cells.

A physiologic detail related to PP release from the digestive tract is that distention is an efficacious stimulus in man’s stomach and duodenum. Indeed, distending the gastric fundus raises PP plasma values. This reflex is abolished by the selective denervation of this stomach region (Figure 1).

A chemical stimulus is also operative to release PP. This has been shown by the presence of food in the gut and by the infusion of bile and pancreatic proteases into the duodenal lumen. The latter constitutes the basis of an "entero-pancreatic-inhibitory reflex" in which, according to Owyang et al PP plays a pivotal role. These authors have also stated that bursts of pancreatic and biliary secretion into the duodenum during late phase 2 and early phase 3 of the intestinal migrating complex and the associated increase in plasma PP are causally related. This contention gets support in their finding that perfusion of the duodenum with bile-pancreatic juice stimulates the release of PP into the circulation. This regulatory peptide might be responsible for the marked suppression of pancreatic and bile acid output during phase 4 of the intestinal migrating motor complex (Figure 2).

Owyang et al have concluded that an undefined factor in bile-pancreatic secretion, unrelated to pH, osmolarity or protein, is responsible of the PP release. According to their view, a cholinergic reflex is involved. Our group has also postulated, in rats, that the inhibition of bile-pancreatic secretion induced by its infusion into the duodenum was the result, at least partially, of a duodeno-pancreatic neural inhibitory reflex.

As regards to the intimate mechanism of the above reflex, Jung et al have provided evidence that PP, in rat pancreatic slices, inhibits potassium-stimulated amylase release but has no effect on the Ach or the octapeptide-elicited secretion of the enzyme. This suggests that PP exerts its action through neural elements, that it acts, essentially, on post-ganglionic cholinergic neurons. The site of inhibition of the cholinergic transmission might be pre-synaptic, between neuron and acinar cells. These effects of PP are, in fact, the first demonstration of a hormone suppressing pancreatic’s enzyme secretion through an interfering of the intrapancreatic cholinergic transmission.

The above-described PP-evoked changes are duplicated by pancreastatin, methionine-enkephalin and somatostatin. These agents provide a fail-safe system that prevents the pancreocytes’ "supramaximal ecbolic stimulation". Adler et al have stressed that somatostatin-28, the same as PP, belongs to what we have described as a pancreon’s brake or as a restraining rein on the physiologic interplay that modulate the exocrine pancreatic secretory process.
As shown later, somatostatin-14, released from the islet’s mantle D-cells, the same as PP, secreted from the “F” or “PP” cells in the Langerhans islets periphery, contribute to modulate the pancreon’s secretory process through influences exerted primarily upon the peri-insular pancreocytes (Figure 3).

During an episode of AP, the pancreatic gland tries to prevent overstimulation of the acinar cells. The testimony of this, in patients, is a progressive increment in the PP plasma values during the first week of an evolving acute inflammatory episode.20 (Table 2).

In the results reported by Torino et al1-3 in which the Tissucol filling of the pancreatic ducts, subsequent to the Cl2Ca-evoked of an AP, allowed the survival of dogs that otherwise might be dead within a 36hs period, the elevated PP plasma values probably disclose that a NER has been triggered. The Tissucol-evoked rising of the PP plasma levels probably occurred superimposed upon spontaneous-enhanced secretion of this regulatory peptide.4

Table 2. Involvement of PP in different types of acute pancreatitis.

<table>
<thead>
<tr>
<th>PP in the Pathology of the Pancreatic Gland</th>
<th>Activation of the “Fail-Safe” or “Brake” System to Prevent “Pancreon’s” Supranormal Ecbolic Stimulation</th>
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<td>1- Biliary Acute Pancreatitis (BAP)</td>
<td>Increasing PP plasma values during the first week</td>
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<td>2- Acute Pancreatitis Post-Large meal</td>
<td>Overpower of “Pancreon’s” “Brake” by excess stimulation of its “trigger” zone</td>
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<tr>
<td>3- Acute Pancreatitis Superimposed on chronic alcoholic Pancreatitis</td>
<td>Hyper-response of the Duodeno-Pancreatic Reflexes due to the ethanol-induced failure of the “Fail-Safe” or “Brake” System</td>
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The pathways of the long neuroendocrine reflex

Having outlined the main features of the short NER, in fact equivalent to our previously described negative duodeno-pancreatic-reflex,8, 21, 22 and the one postulated by Owyang et al16 as an entero-pancreatic-inhibitory reflex in the feedback control of the exocrine pancreatic secretion, it remains to be analyzed the long NER (Figure 1 and 2). The latter consists of two distinctly hemi-loops. The first one runs from the vagal-nuclear-complex in the brain-stem up to the PP cells which are located in the periphery of the Langerhans islets and scattered in the exocrine pancreas. The hemi-loop that completes the feedback circle pivots around the circulation of PP from the pancreatic gland up to the vagal-nuclear-complex. The nerve impulses that flow the former hemi-loop run, initially, in the gastric branches of both vagal trunks. Subsequently, through what we have conceived as a neural-plexual-freeway, that has as a background the gastro-duodenal myenteric and submucosal neural networks.23 After traversing the pylorus or bypassing it following the pathway of the hepatic branch of the left vagus, reaching afterward the neural plexus of the enteric “freeway” at the first duodenal segment, the neural vagal impulses follow the nerve fibers that jump, in rich density, the duodeno-pancreatic cleft. This duodeno-pancreatic linking takes place primarily in the region comprised between the pylorus and the peri-Vaterian region.24-34 Confirmation that this is precisely the pathway followed by the first hemi-loop of the long NER is provided by the experimental observation, in dogs, that duodeno-pancreatic disconnection or duodenectomy29 diminishes, on the one hand, the hypoglycemia-induced PP release and, on the other, the hormonal PP response to a meal.

As for the second hemi-loop of the long NER, the one that pivots around the blood stream PP flow from the pancreatic gland up to the vagal-nuclear-complex in the brain-stem, the group of Taylor29 has given evidence that this regulatory peptide does indeed enter in contact with the vagal-nuclear-complex. The latter is achieved at the level of the area postrema and the tractus solitarius nucleus, two regions of the central nervous system that lack a blood barrier. Once in the vagal-nuclear-complex binds to specific receptors and inhibits the neuron discharge of the dorsal motor nucleus. The consequence is a depression of exocrine pancreatic secretion due to a fall of the intrapancreatic cholinergic tone. The latter, as we have already pointed out, normally exerts an intricate interaction with the hormonal system upon the pancreon’s secretory process.5 (Figure 1).

From all the preceding facts, it is evident that the long as well as the short NER are important components of a fail-safe system or, in other words, of a brake mechanism that, under physiologic circumstances prevents the noxious consequences upon the acinar pancreocytes which are attached to abrupt episodes of supramaximal-ecbolic-stimulation.33-34 (Figure 1, 2 and Table 1). When the above-depicted fail-safe or brake system is overpowered, as for example following a large meal
rich in proteins, fats and alcohol, an episode of AP can be evoked.

In the sixties, we have pinpointed this clinical entity as an AP due to an over activation of the “trigger” zone (PV-D) of the exocrine pancreas (“pancreon” units). Indeed, intoxication might explain the remarkable divergent changes the ethanol-evoked impairment of the pancreon’s “brake”, primarily of PP must play a crucial role.

A remarkable subsequent observation was that chronic ethanol administration induced a reversal of the exocrine pancreatic secretion inhibitory response evoked, during the nonalcoholic stage, by an acute ethanol administration. This finding was interpreted by us as the consequence of alcohol-elicited impairment of the normal physiologic “brake” system in the control of exocrine pancreatic secretion by the pancreon units.

The presumption was that an ethanol-induced vagal neuropathy was responsible of this change. The latter preferentially impinging upon the Pavlov fibers (inhibitory), precisely the ones resulting in the release of PP and perhaps other members of the pancreon’s fail-safe system, like beta-endorphins, substance P and somatostatin.

The release of PP and/or beta-endorphin evoked during the nonalcoholic stage by an episode of acute ethanol intoxication might explain the remarkable divergent changes of gastric and exocrine pancreatic secretion. Indeed, under the above circumstances, there is an enhancement of gastric acid secretion and a simultaneous inhibition of exocrine pancreatic secretion. Suggestively, similar secretory responses, as the above outlined, are observed with the administration of either beta-endorphin or PP. As to the postulation of an alcohol-evoked impairment of the Pavlov fibers, the vagal neuropathy described in humans by Duncan et al and the degeneration of intrapancreatic nerve fibers shown by Berger and Feher in mice, provide a suggestive support to this contention. Moreover, the secretory studies, in dogs, by Schmidt et al add further confirmation to our previously stated assumption of an ethanol-elicited gradual disappearance of an exocrine pancreatic secretion inhibitory factor.

The Hajnal et al report, in humans chronic alcoholics, of a lack of PP response following an acute ethanol or wine administration superimposed to a test meal, undoubtedly offers an additional support to our hypothesis above outlined.

The loss of involvement of the Pavlov vagal inhibitory fibers induces the phenomenon of neural decentralization and the consequent enhanced reactivity of the peripheral antro-duodeno-pancreatic reflexes, and as a result of that, an increased intrapancreatic cholinergic tone. In fact, the idea of an evolving supranormal-ecbolic-stimulation of the pancreon units secondary to the loss of the negative component of pancreas innervation (PP, endorphins, etc) is at the core of our postulation concerning the physiopathogenesis of alcoholic pancreatitis. (Table 2).

That chronic alcoholism is indeed associated with an increased reactivity of the entero-pancreatic reflexes, is supported by Brugge et al findings. These authors have observed, in man, that: a) basal duodenal trypsin output and the interdigestive duodenal contraction rate are higher in chronic alcoholics that in controls; b) chronic alcoholics have an increased post-prandial trypsin secretion compared to nonalcoholics; c) alcohol-fed and nonalcoholics show similar post-prandial increments in plasma levels of gastrin and CCK; d) when compared to nonalcoholics, chronic alcoholics display a remarkable lack of PP increments in respect to a test meal or test solutions; e.g., glucose, ethanol, wine.

Yamasaki et al have reported, in monkeys, that chronic ethanol administration induces papillary dysfunction and exocrine pancreas hyper secretion. Both might play a role in the observed raised pancreatic ductal pressure and the microscopic signs of acinar pancreocytes hyper function.

It is our postulation that in the above-described changes the ethanol-evoked impairment of the pancreon’s brake, primarily of PP must play a crucial role.

A feature to emphasize is that a similar sequence of events develops in uremia. Indeed. Owyang et al have shown, in patients with chronic renal insufficiency, that both intraduodenal mannitol and i.v. CCK evoke an ecbolic hyperresponse of the exocrine pancreas. A closely similar finding was reported in rats in which the renal mass had been 85% surgically reduced. This unexpected exocrine pancreatic secretion change response was attributed to an autonomic nervous imbalance characterized by a predominant cholinergic prevalence.

In the above clinical and experimental setting, it seems logical to presume an impairment of the pancreon’s brake with a failure centered in the synthesis and/or release of PP.
Another observation to be stressed is that the same fall of PP secretion that we have commented upon in chronic alcoholic patients have been shown in patients with cystic fibrosis and in dogs with pancreatic fibrosis following ductal ligation. 1, 2, 4, 9

A feature to emphasize in all these entities is that the depression of PP release occurs in the presence of an augmented number of PP cells in both the islets and the exocrine parenchyma.

The above observation prompts the speculation if in the chronically inflamed gland the PP cells produce less hormone than normally or, contrariwise, release less in response to different stimuli.

As the absolute secretion normally elicited by secretin, CCK, food and insulin hypoglycemia is similarly affected, one is tempted to assume that the defect of PP release in chronic pancreatitis is localized to the common final pathway of these stimuli, probably in or near the PP cell itself.

The above discussed PP secretion failure is probably at the basis of the enhanced intrapancreatic cholinergic tone characteristic of chronic alcoholics. In these patients it might explain the supranormal secretagogue pancreatic response following the intraduodenal infusion of a CCK releaser (oleic acid) and, besides, of its normalization by previous atropinization. 30

The above findings justify our postulation that chronic alcoholism sensitizes the trigger zone (PV-D) of the pancreatic gland. 22, 31, 33, 51 This notion helps to explain why these patients are more prone to develop AP episodes especially following the ingestion of large meals, rich in protein and fat (CCK releasers), phenomenon that is potentiated when associated to alcohol ingestion (Table 2).

**Adrenergic innervation and PP secretion**

Besides the already discussed interactions that evolve between PP and the pancreatic cholinergic innervation, it is worth considering those that relates to the adrenergic component of the autonomic nervous system. In that sense, several studies have shown that beta-blockade inhibits PP release, while, contrariwise, alpha-blockade enhances PP secretion.

The raised PP plasma values observed after celiac ganglionectomy might be related to the interruption of alpha adrenergic influences; according to Larson et al42 celiacectomy may be considered a sort of alpha-adrenergic-blockade.

In relation with the fore mentioned innervation features, it should be pointed out that celiac ganglionectomy does indeed affect the nerve terminals and the catecholamine content of the pancreas: norepinephrine drops by 90%, while epinephrine and dopamine fall by 50 to 70%. 7, 9, 52, 53 Blocking of the celiac ganglia by local anesthetics might trigger a rising of the PP plasma values. If this is confirmed, it could mean a new support to justify this procedure in cases of AP. In this clinical setting, celiac ganglion anesthesia would constitute a therapeutic measure not only to relieve pain but also an efficacious method to reduce the intrapancreatic cholinergic tone.

**PP Involvement in the pancreatic trophic changes**

When the NER are disrupted or when a diabetic syndrome has fully developed, islet-acinar (insulo-pancreo-axis) alterations are induced that, themselves, evoke trophic changes in the pancreatic gland.

The former situation is represented by our results in rats subjected to either a transection and reanastomosis of the peri-Vaterian region or to a resection of supra-Vaterian duodenal segment. 24 The pancreatic head weight increase observed under the above circumstances is probably related to an enhanced PP concentration in the pancreatic gland. This probably as a result of the interruption of the cholinergic impulses flowing through the hypothetical of the network of the enteric autonomic nervous system, characterized by us as a sort of neural-plexual-freeway. 23 The main support of this contention is the Raher et al’s report. These authors have speculated that PP cell, located predominantly (90%) in the dorsal region of the pancreas may exert a local trophic effect. This property might protect the PP-rich lobe of the gland from atrophy in diabetic patients. These statements of Raher et al13 are based on their observations at an autopsy level. Indeed, in insulin-dependent diabetics (IDO) it was remarkable that the pancreas was markedly decreased but at the expense of an almost selective atrophy of the lobe poor in PP cells. A similar finding, although of a lesser magnitude, was appreciated in non-insulin dependent diabetics (NIDD) patients. These findings are in keeping with those of Greenberg et al54 observations in rats in the sense that PP increase DNA synthesis in pancreatic acinar cells. They are also coherent with the microscopic detail that PP cells present long cytoplasmatic processes in contact with the exocrine cells. The above features fit coherently with the suggestive Oria’s55 observations that the pancreatic hook segment, the one, located around the mesenteric vessels, precisely the richest one in PP cells, is frequently spared in the severe episodes of necrotizing AP.

Concerning diabetics, Will et al56 have reported that these patients with signs of autonomic neuropathy reveal an impaired PP plasma response following insulin-induced hypoglycemia. That is to say a clinic-pathological situation closely resembling that of chronic alcoholism.
Other features of the pp involvement in the insulo-pancreonaxis interaction

The Raher et al. findings bring into focus the interesting phenomenon of the islet-acinar-axis interactions. Some of them are unequivocally exerted by the PP cells located in the periphery of the Langerhans islets.

Nakagawa et al. have recently concluded that the insulin resistance that has been demonstrated in chronic pancreatitis or the Whipple procedure (pancreatogenic or type 3 diabetes), a deficiency of PP may play an important role. This speculation is supported by the well known procedure of PP administration ameliorates the hepatic insulin resistance.

The above speculation is supported by the well known and accepted histologic phenomenon centered on the presence of peri-insular halo that has been accepted as an expression of more synthesis and less secretion of enzymes in the peri-insular region.

Immediately after a meal, the increased insulin could stimulate digestive enzyme synthesis in the peri-insular cells, whereas local somatotatin and/or PP provide means to rapidly regulate the suppression of enzyme release from the acinar cells.

Chronic pancreatitis and pp

In this review it seems appropriate to make reference to the fact that PP administration ameliorates the hepatic insulin resistance that has been demonstrated in chronic pancreatitis. Seymour et al. have reported that PP secretion is impaired in patients with severe chronic pancreatitis and that a strong correlation exists between test meals demonstrated PP deficiency and hepatic insulin resistance. They have also shown that chronic PP treatment increases insulin receptor concentration in hepatocytes membranes from chronic pancreatitis rats and also improves glucose tolerance after enteral administration of dextrose.

According to Brunicardi, in the diabetic syndrome associated with chronic pancreatitis or the Whipple procedure (pancreatogenic or type 3 diabetes), a deficiency of PP may play an important role. This speculation should be coupled to that of a glucagon deficit originally suggested by Bank. The impairment of both PP and glucagon it should be added that of a beta-cell hypo function. The latter would be consequence of a previous period of cholinergic-elicted beta-cell hyper stimulation that has been put in evidence by Patto et al. in chronic alcoholic patients.

Berger and Feher have shown, in mice, after alcohol feeding, the degeneration of the intrapancreatic nerves which are an important source and locus of action of PP. These immunohistochemistry findings provide support to our contention that alcohol feeding does indeed impair the normal physiologic brake that modulates exocrine pancreatic function. This change, as we have already pointed out, might be at the basis for the increased reactivity of the trigger (PV-D) and of the pancreon units that characterize the earlier stages of alcohol-evoked chronic pancreatitis.

Final comments

All the forementioned data provide substantial basis to justify our earlier contention that the Tissucol-evoked beneficial effects upon the dog’s Cl2Ca-elicited AP lesions are related to the enhancement of normal islet-acinar cells interactions and triggering of the long NER from the intrapancreatic ductal tree.

The latter assumption seems to get support in the results observed with the exogenous PP administration in rats subjected to taurocholate-induced AP. Indeed, when this regulatory peptide was subcutaneously injected every 8 h preceding the AP episode, the animal mortality rate dropped from 100 to 50%. When PP was injected for 7 days following the taurocholate-induced pancreatic inflammatory lesion, animal mortality rate was of 52%, but when the above-described approaches were associated, the mortality rate fell significantly to a level of 15%.

From all the above gathered information and discussion, it seems logically warranted to perform tests with PP in order to elucidate whether or not this regulatory peptide is, in fact, a valuable preventive and/or therapeutic agent when an acute inflammatory episode is suspected.

Referencias

Polipéptido pancreático: revisión de su ingerencia provocadas por el etanol
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