graphic brush cytology has proved negative by using a fine spring-loaded Tru-cut needle Biopsy gun (Bipty TM, Radiplast, Uppsala), percutaneously under ultrasound guidance. The use improved metal stents was alluded to in the leading article, and a primary indication would be those patients with non-resectable hilar cholangiocarcinoma in whom good survival might be expected. There is no evidence that the optimum route of insertion for these metal stents is via the transhepatic route, and there are many advantages in placing metal stents endoscopically.

Low structures
Messrs Russell and Rees failed to reference Roux en Y cholecdochojunostomy as the preferred procedure for biliary drainage, for there is none except surgical history. In fact, conventional cholecdochooduodenostomy is an adequate bypass and this has the added advantage of easing endoscopic procedures later if nodes or tumour growth around the porta hepatis obscure the bile duct.

We agree that there are a few figures suggesting an improved survival after radical resection, but numbers are small and very few centres achieve a 5% mortality, and cures are rare. The improved survival may merely indicate the improved selection.

We agree with Professor Cotton that difficulties do arise in the management of a few patients for whom resective surgery may be the correct approach. The patients are best cared for in a unit which has a specialist team of interventional endoscopists, radiologists, and a pancreatobiliary surgeon.

In the article routine stenting was not advocated; in fact, it was clearly pointed out that clinicians now have to take a mature approach in their decision making, and have to balance the patient's condition and likelihood of survival before deciding on a surgical or an endoscopic approach to palliation. Clearly, there are going to be some patients who look fit and well, and in whom some form of surgical palliation may be appropriate to reduce the need for admissions, which would be necessary in the patient stented endoscopically. At the other end of the spectrum there are those patients over the age of 70 years in whom a simple stenting procedure seems a very suitable alternative to any form of surgical palliation, particularly, when we know that most patients die jaundice free with their original stent in situ.

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Ultrastuctural demonstration of histamine in human enterochromaffin like cell granules

Str—We read with great interest the paper by Lönroth et al.1 by using immunohistochemical methods in light microscopy in normal volunteers, the authors showed that in addition to mast cells some gastric endocrine cells contained histamine. These cells were located exclusively in the fundus, constituted 44% of the total number of endocrine cells in the oxyntic mucosa, and stored neither 5 hydroxy-

tryptamine nor somatostatin. These endocrine cells were supposed to be enterochromaffin like cells. To study this hypothesis, we tried to demonstrate histamine (HA) in the enterochromaffin like cells by an immunocytochemical method in electron microscopy. Ultrastuctural analysis of fundic sections, indeed, allow enterochromaffin like cells to be distinguished from their typical secretory granules, from the other gastric endocrine cells.2 This study was done in a patient with pernicious anaemia, hypergastrinaemia (>1000 pg/ml), and microglandular hyperplasia of gastric (Gri-mellus argyrophil technique). Fundic mucosal biopsy specimens were obtained during gastroscope, fixed in 4% glutaraldehyde in phosphate buffer at 20°C, dehydrated in ethanol, and embedded in Epon 812. Ultrathin sections were cut and mounted on gold grids. They were incubated for four hours in 1/200 diluted polyclonal guinea pig anti-HA antibodies (Peninsula ref. 61069) at room temperature and rinsed in phosphate buffer and in distilled water. The grids were then incubated for one hour in a 5 nm gold particle conjugated antiguinea pig immunoglobulin (Biocell EMGAGS). 1/20 diluted in phosphate buffer solution. Finally, the grids were rinsed in water, dried, and contrasted with uranyl acetate in ethanol.

The efficiency and the specificity of the immunocytochemical study were first checked on rat peritoneal mast cell (PMC) sections. Normal rat PMC granules were shown to contain HA while, after in vitro incubation in a poly-L-lysine (secretagogue) solution,3 the PMC granules no longer contained HA.

In the human biopsy sections few mast cells were present around the gland. They were shown to contain HA in their uniformly electron dense granules. The enterochromaffin like cells were identified ultrastructurally. Most of these cells contained granules positively marked by the anti-HA immunocytochemical reaction (Figure). This reaction was reproduced several times on many sections of the same biopsy specimen with concordant results. The control sections were incubated either with anti-HA neutralised by HA followed by the immunogold reaction, or with immunogold antibodies alone. They were both negative except a light aspecific background. The results of this study are in agreement with Lönroth’s conclusions: the histamine-containing endocrine cells of the human fundus are enterochromaffin like cells. The role of these cells in the physiology of histamine mediated acid secretion should be explored. Further studies should determine if the cell granules release histamine under the influence of gastrin.

Ultrastuctural histamine demonstration in enterochromaffin like cells secretory granules on human fundus biopsy section. The 5 nm gold particles are located in the electron dense granules and inside a typical vaculated granule (original magnification x60,10).

Reply

Sm,—Dr Delwaide and colleagues have with this report further confirmed that the enterochromaffin like cells of the human fundus indeed contain histamine. In the human gastric mucosa these cells are confined to the oxyntic gland area, which also presents a higher histidine decarboxylase activity than the non-acid producing pyloric gland region.3 Patients with hypergastrinaemia of different origin also have a higher histidine decarboxylase activity together with an increased density of enterochromaffin like cells in the oxyntic gland area.4 In addition, pentagastrin infusion is followed by a release of histamine and by a substantial increase in histidine decarboxylase activity in the oxyntic gland mucosa of healthy volunteers. These findings, in induced and non-induced patients do not occur in the pyloric gland region...2

In conclusion, all this circumstantial evidence together with the results presented by Dr Delwaide and colleagues favour the view that the enterochromaffin like cells of the human stomach store histamine, release the amine on proper stimulation, and have the capacity to synthesise histamine.

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BOOK REVIEWS


Although secretory diarrhoea is not everyone’s cup of tea, I approached this book with some