Staining procedures for the endocrine cells of the upper gastrointestinal mucosa: Light-electron microscopic correlation for the gastrin-producing cell

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SYNOPSIS Although histochemical, immunohistochemical, and electron microscopic methods have led to the identification of a large variety of endocrine cells in the upper gastrointestinal mucosa, no conventional light microscopic technique capable of the simultaneous identification of these cells has been reported. Such a staining method would be of considerable value to the pathologist as the malfunction of the endocrine cells of the gut, which produce numerous digestive hormones and biogenic amines, is closely related to a number of clinical conditions afflicting man. In this work, after testing three different polychrome staining methods, it has been concluded that a slightly modified Herlant's tetrachrome in tissues fixed in Zenker-formol is the procedure of choice. This method allows the distinction of several different cell types in the upper gastrointestinal mucosa of man and dog and permits the easy identification of the gastrin-producing cells on a routine basis. This identification has been confirmed in the case of two patients with gastrin cell hyperplasia, seen by both light and electron microscopy. Herlant's tetrachrome has proven valuable in the screening of human as well as experimental gastrointestinal tissues and it has been found to be very suitable for recognizing gastrin-producing cell hyperplasias. The usefulness of this method is expected to increase with the establishment of further correlations between the light and electron microscopy of the endocrine cells of the gut.

A large number and a remarkable variety of endocrine cells have been described in the upper gastrointestinal mucosa using histochemistry, immunohistochemistry, and electron microscopy (Capella, Solcia, and Vassallo, 1969; Forssmann, 1970; Forssmann, Orci, Pictet, Renold, and Rouiller, 1969a; Pearse, Coulling, Weavers, and Friesen, 1970; Sasagawa, Kobayshi, and Fujita, 1970; Vassallo, Solcia, and Capella, 1969, 1971). Most of these cells produce either polypeptide hormones (Grossman, 1968) or biogenic amines (Aures, Håkanson, and Owman, 1970; Håkanson, 1970). A nomenclature for the endocrine cells of the gut was adopted at Wiesbaden in 1969, chiefly on the basis of their electron microscopic appearance (Creutzfeldt, 1970). Seven basic types are currently recognized: A cells, purported to produce enteric glucagon; D cells, of unknown function; EC cells, the producers of serotonin; ECL cells, assumed to store some biogenic amines; G cells, the source of gastrin; S cells, said to produce secretin; and L cells, the function of which is still unsettled.

Studies on the endocrine cells of the gut by light microscopy have relied largely on histochemical and immunohistochemical methods each of which allows, as a rule, the identification of only one cell type (Bussolati and Pearse, 1970; Håkanson, 1970; Bussolati, Capella, Solcia, Vassallo, and Vezzadini, 1971; Håkanson, Owman, Sporrong, and Sandler, 1971; Lomský, Langr, and Vortel, 1971; Masson, 1914; McGuigan and Greider, 1971; Pearse, 1968; Polak, Bloom, Coulling, and Pearse, 1971a, b; Solcia, 1972; Solcia, Vassallo, and Capella, 1970a). After using these methods and/or electron microscopy quantitative as well as qualitative changes have been noted in the endocrine cells of the gut from patients with a number of clinical conditions. These include: (a) carcinoid tumours associated with typical (Williams and Sandler, 1963; Black, 1968; Soga and Tazawa, 1971) and atypical (Sandler and Snow, 1958; Campbell, Gowenlock, Platt, and Snow, 1963)
carcinoid syndrome, as well as peptic ulcers (Black and Haffner, 1968); (b) states of gastric acid hypersecretion, such as the Zollinger-Ellison syndrome with intestinal gastrinoma (Oberhelman, Nelson, Johnson, and Dragstedt, 1961; Thompson, Hirose, Lemmi, and Davidson, 1968), some patients with acromegaly (Pearse and Bussolati, 1970; Creutzfeldt, Arnold, Creutzfeldt, Feurle, and Ketterer, 1971), hyperparathyroidism (Pearse and Bussolati, 1970; Polak, Bussolati, and Pearse, 1971c) retained antrum (Solcia, 1972) and a percentage of peptic ulcers (Solcia, Capella, and Vassallo, 1970b; Lechago and Bencosme, 1973); and (c) states of gastric hypersecretion, such as pernicious anaemia (Rubin, 1969; Creutzfeldt et al, 1971; Polak, Coulling, Doe, and Pearse, 1971d). No light microscopic technique capable of demonstrating simultaneously the various endocrine cell types of the digestive mucosa has been reported so far. Such techniques should enable the pathologist to study biopsy and surgical specimens in more detail than has been possible heretofore. It would also be of considerable value to the researcher dealing with these endocrine cells in human as well as experimental subjects.

To develop such a procedure we have tested a number of polychrome techniques known to be capable of distinguishing various polypeptide-producing cell types in other endocrine glands. Modifications of Masson’s trichrome and Gomori’s chrome-alum-haematoxylin have been successfully used to stain the cells of the pancreatic islets (Bencosme, 1952), while Herlant’s tetrachrome has proven to be adequate in discriminating among a large variety of cell types in the hypophys (Herlant, 1960; Kraicer, Herlant, and Duclos, 1967). In the present work an assessment of the value and limitations of these staining techniques will be made using samples of human and canine upper gastrointestinal mucosa. In addition, correlations will be attempted between the polychrome methods mentioned above and some silver impregnation procedures traditionally used to identify endocrine cells in the gut (Masson, 1914) and the pancreas (Hellerström and Hellman, 1960).

Among the endocrine cells of the gut the gastrin-producing cell has acquired increasing relevance in the last few years as gastrin appears to be a key factor in the pathogenesis of clinical conditions such as the Zollinger-Ellison syndrome (Gregory, Tracy, French, and Sircus, 1960; Friesen, Tracy, and Gregory, 1962, 1970; McGuigan and Trudeau, 1968; Becker and Seelig, 1969) and a significant percentage of peptic ulcers (Emås and Fyrö, 1964; Bonfils, Dubrasquet, Lewin, and Vatier, 1969; Byrnes, Young, Chisholm, and Lazarus, 1970; Emås, Borg, and Fyrö, 1971; Trudeau and McGuigan, 1971). Furthermore, the human gastrin-producing cell has been recently identified by electron microscopic immunohistochemistry (Greider, Steinberg, and McGuigan, 1972), while the canine gastrin cell has been identified by correlating immunofluorescence with electron microscopy observations (Pearse and Bussolati, 1972). In view of these developments an effort has been made here to correlate light and electron microscopic observations in order to identify this cell type by conventional light microscopy.

**Materials and Methods**

Human tissue samples were collected from the mucosa of the gastric corpus, pyloric antrum, and duodenal bulb of 10 subtotal gastrectomy specimens. Five of the patients had chronic peptic gastric ulcer, three had a chronic peptic duodenal ulcer, one had both duodenal and gastric ulcers, and one had an adenocarcinoma of the stomach. For the study of the uninvolved mucosa special care was exercised to avoid areas with ulcer or tumour. Tissues from canine fundic, pyloric, and duodenal mucosa were obtained from several anaesthetized animals.

Tissues were fixed for light microscopy in 10% buffered formalin, pH 7, for four to 10 days, Zenker-formol for 24 to 48 hours, and Bouin’s fixative for 24 hours. The tissues were then dehydrated in graded ethanols, embedded in paraffin, and sections 2 to 6 microns thick were mounted on glass slides. Some sections were stained with modified Masson’s trichrome, Gomori’s chrome-alum-haematoxylin (Bencosme, 1952), and Herlant’s tetrachrome (Herlant, 1960; Kraicer et al, 1967). Regarding the last technique, we have applied a modification of the variant developed by Kraicer and coworkers (1967). Since this was the method of choice and its execution is somewhat critical, it is described in detail here:

1. Fix in Zenker-formol for 24 to 48 hours.
2. Wash in tap water for 12 to 24 hours.
3. Place in 5% iodine in 90% ethanol for 12 hours.
4. Dehydrate, embed in paraffin, cut sections 2-4 μ thick, rehydrate.
5. Wash in running tap water for 30 minutes.
6. Stain in erythrosin solution for two hours. Erythrosin B, CI no 773 (Anachemia) 1 g; 0.2 M acetic buffer, pH 6.2 (Walpole) to 100 ml.
7. Rinse in distilled water.
8. Stain in Mallory Blue II for five minutes. Stock solution: Aniline Blue, water soluble, Michrome (E. Gurr) 0.5 g; Orange G, CI no. 27 (Anachemia) 2 g; distilled water to 100 ml; when dissolved, add 8 ml of glacial acetic acid. To stain dilute with an equal volume of distilled water.
9. Rinse in distilled water.
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10 Stain in acid Alizarine Blue solution for 10 minutes. Acid Alizarine Blue, BB, Microme no. 9 (E. Gurr) 0·5 g; aluminium sulphate or chloride 10 g; distilled water to 100 ml. Boil for three minutes until blue-purple, cool, and bring volume up to 100 ml with distilled water.
11 Rinse in distilled water.
12 Place into 5% aqueous phosphomolibdic acid for 15 minutes or more.
13 Transfer to 1% phosphomolibdic acid in 70% ethanol for four seconds.
14 Transfer to 1% phosphomolibdic acid in 90% ethanol for eight seconds.
15 Transfer to absolute alcohol (two changes), clear in toluol and mount.

Other sections were silver impregnated using the methods of Fontana-Masson for argentaffinity (Masson, 1914) and of Hellerström and Hellman (1960) for argyrophilia. Some representative blocks from different regions were serially sectioned to establish correlations between the results obtained with the different techniques, particularly with regard to the gastrin-producing cells.

For electron microscopy tissues were fixed for two hours in cold 3% glutaraldehyde in 0·1 M phosphate buffer pH 7·2. Then they were washed in the same buffer plus 10% sucrose for two hours and, subsequently, postfixed in 2% OsO₄ for one and a half hours. All reagents contained one drop of 1% CaCl₂ per each 10 ml of solution. The tissues were dehydrated with graded ethanolos and embedded in Epon 812. Sections obtained with an LKB ultratome were double-stained with uranyl acetate (Stempak and Ward, 1964) and lead citrate (Venable and Coggeshall, 1965) and photomicrographs were taken with a Hitachi HU 11C electron microscope.

Results

Bouin’s fixative and 10% buffered formalin were the fixatives of choice for the silver impregnation of Fontana-Masson while the three fixatives used were adequate for the method of Hellerström and Hellman. Zenker-formol, on the other hand, was the most suitable fixation for the subsequent application of the polychrome stains. In tissues fixed by the latter method, the most rewarding technique for the simultaneous demonstration of the various endocrine cell types of the upper gastrointestinal mucosa was Herlant’s tetrachrome. Masson’s trichrome was also suitable, while Gomori’s chrome-alum-haematoxylin was a distant third choice and will not be discussed further in this paper.

A synopsis of the endocrine cells seen in each of the three regions of the gut is presented in the table. Positive identification is possible only for the G and the enterochromaffin cells. Two more tentative identifications are made with regards to the A-like and D-like cells, while the remainder of the cells seen after the polychrome techniques will have to await the establishment of further correlations before they can be identified.

A conspicuous cell type, restricted to the antrum, has a pear-shaped cytoplasm which stains pale pink after Herlant’s tetrachrome (fig. 1). These cells stain pale brown after Masson’s trichrome. They are believed to be the gastrin-producing cells since they follow the same topographic distribution as those identified by several authors (Bussolati and Pearse, 1970; McGuigan and Greider, 1971; Pearse and Bussolati, 1972) in a number of species using immunohistochemistry. The gastrin cells of the dog are larger and their granules coarser than those of their human counterpart, stain brighter red after Herlant’s tetrachrome (fig 2) and darker brown after Masson’s trichrome.

When adjacent sections from blocks fixed in Zenker-formol are stained with Herlant’s tetrachrome and the Hellerström-Hellman silver technique respectively, the gastrin-producing cells are silver negative. In tissues fixed in 10% formalin and in Bouin’s fixative these cells also appear to be silver negative with the same silver method. In contrast, cells with brown-yellow granules are silver-positive (argyrophilic).

Cells with granules stained golden-yellow after Herlant’s tetrachrome (fig 3) and brown-black with Masson’s trichrome are noted in small numbers in the human fundus and pylorus and in large numbers in the duodenal mucosa. In the dog these cells are relatively more numerous in the stomach and less conspicuous in the duodenum.

Fontana-Masson’s technique for argentaffinity yields small numbers of silver-positive cells in antral and fundic mucosa, whereas the duodenal crypts
exhibit the presence of significant numbers of these cells. No direct comparison is possible between the results obtained with Herlant's tetrachrome and with Fontana-Masson's silver technique as they require different fixations. Close inspection, however, strongly suggests that those cells containing black-brown granules after Masson's trichrome and brown-yellow (golden-yellow for human duodenum) granules after Herlant's technique correspond to the argentaffin cells on account of their number, shape, and location.

The other cells listed in the table are tentatively considered separate types on the basis of their different tinctorial affinities. The lack of correlation with other techniques at this point, however, renders their definitive identification a subject for further research.

The gastrin-producing cells were identified by electron microscopy in both dog and man. Canine gastrin cells are characterized by the presence of numerous round cytoplasmic granules, composed of a well defined limiting membrane which encloses a space either electronlucent or with scanty flocculent material. In man, cells were noted in the antrum identical to those recently identified as gastrin-producing cells by Greider and coworkers (1972) using electron microscopic immunohistochemistry. These cells are characterized by the presence of cytoplasmic granules between 120 and 200 nm in diameter, a somewhat irregular profile, and possessing a flocculent core of variable electron density (fig 4).

In two patients with chronic peptic ulcer (one gastric and one duodenal) there was marked hyper-
Fig 4  Human antrum. Electron microscopic picture of a gastrin-producing cell (G) containing cytoplasmic granules of variable electron density. Some of these are dark while others have a nearly 'empty' appearance. × 6400.
Inset  Close-up view of human G cell granules. × 26 400.

Fig 5  Human antrum from a patient showing G cell hyperplasia. Five gastrin-producing cells (G) are noted in a cross-section of an antral gland. Pale granules are numerous in these hyperplastic G cells. × 6000.
plasia of the gastrin-producing cells when observed under electron microscopy (fig 5). When antral tissues from these patients were stained with Herlant’s tetrachrome, it was seen that the pale pink cells, purported above to be the gastrin-producing cells, were also markedly increased in number (fig 6).

Discussion

The main morphological features leading to the identification of the endocrine cells of the gut by light microscopy are: (a) comparatively small size; (b) basal location with respect to the exocrine elements; and (c) the presence of specifically stainable cytoplasmic granules. Herlant’s tetrachrome modified by fixation with Zenker-formol has been the most adequate of the methods tried to characterize the various endocrine cell types of the digestive mucosa. It should be emphasized at this point that, in order to obtain satisfactory results with the techniques described here, care must be taken in the performance of the various steps. Polychrome techniques in general require careful ‘tailoring’ for optimal results with different tissues and even different species. Once these factors are taken into consideration, however, consistent results may be expected. Marked autolysis of the superficial digestive mucosa noted in necropsy specimens not reported here precludes in most cases the attainment of satisfactory results with these polychrome techniques. Unlike histochemical and immunohistochemical approaches, which often require sophisticated techniques, the polychrome methods tested here are essentially simple to perform and can be readily established on a routine basis. In addition, Masson’s trichrome and particularly Herlant’s tetrachrome, appear to be less controversial than Grimelius’ silver technique (Solcia, 1972) and more specific than lead haematoxylin (Pearse and Busolati, 1972) for the demonstration of the antral gastrin-producing cells. As these techniques allow the simultaneous demonstration of the various endocrine cell types scattered throughout the digestive mucosa, they facilitate the task of evaluating qualitative and/or quantitative changes in tissues from experimental animals and man.

The enterochromaffin cells have been traditionally demonstrated with Fontana-Masson’s method for argentaffinity. Unfortunately, this technique cannot be applied to tissues fixed in Zenker-formol, therefore precluding the use of serial sections to compare the results obtained with argentaffin and polychrome techniques. Nevertheless, it appears that on the basis of number and topographical distribution, the argentaffin cells correspond to the brown-yellow (golden-yellow in human duodenum) cells after Herlant’s tetrachrome. This correlation is further strengthened by the observation of the gastrointestinal mucosa of dogs treated with reserpine (Lechago and Bencosme, 1973). After 24 hours of treatment the argentaffin cells after Fontana-Masson’s method and the brown-yellow cells after Herlant’s tetrachrome have completely disappeared.

Comparison of serial sections of tissues fixed with Zenker-formol and stained with the Hellerström-Hellman silver technique and with Herlant’s tetrachrome respectively shows that the brown-yellow cells are argyrophilic. The gastrin-producing cells, on the other hand, are silver negative after all fixations used. These observations are consistent with those of Mcguigan and Greider (1971), who found that the gastrin-producing cells as identified by immunofluorescence were silver negative with a variety of impregnation techniques, including that of Hellerström and Hellman. Lomsky and co-authors (1971), on the contrary, reported that the gastrin-producing cells they identified by immunofluorescence in antral as well as duodenal mucosa were silver positive when re-stained with the Hellerström-Hellman technique. One explanation for this discrepancy is that these different authors introduced modifications in their silver impregnation techniques.

Fig 6 Light microscopic picture of the same antrum described above showing numerous gastrin-producing cells (G). Herlant’s tetrachrome. × 400.
which are not apparent from reading their methodology. Or, alternatively, that their immunohistochemical techniques demonstrate two different types of gastrin-producing cells: one of them would be silver-negative while the other would be silver-positive. The latter possibility seems to find some support from the observations of Bussolati and Pearse (1970). These authors reported that while most of the antral gastrin-producing cells are positive with Grimelius’ silver technique for argyrophilia, another group of cells, displaying weaker fluorescence, were silver-negative with the mentioned technique.

The electron microscopic identification of the human antral gastrin-producing cells has been greatly facilitated by the recent work of Greider and co-workers (1972), who showed these cells after immunoperoxidase labelling techniques. Light-electron microscopic correlation in human subjects has been confirmed by the observations reported here of tissues from gastric mucosa with hyperplasia of the gastrin cells seen by both light and electron microscopy.

In the case of the dog, the gastrin cells have been recently identified by electron microscopy in tissues previously treated with immunofluorescence by light microscopy (Pearse and Bussolati, 1972). Reports by others that these cells change their appearance following alcohol infusion (Forssmann and Orci, 1969) or after feeding (Forssmann, Orci, Forssmann, and Rouiller, 1969b) had already suggested their identity. Similarities in topographical distribution when comparing light with electron microscopy of canine antra indicate that the antral red cells after Herlant’s threchome are the gastrin-producing cells in this species.

In conclusion, Herlant’s threchome as modified in this work is offered as the routine method of choice for the simultaneous recognition of a variety of endocrine cells in the upper gastrointestinal mucosa. This technique is particularly suitable for the identification of the gastrin-producing cell in man as well as in experimental animals. Although the small number of gastrectomy specimens studied in his work precludes the drawing of any meaningful conclusion from the statistical standpoint, it seems apparent from our results that marked gastrin cell hyperplasias can be readily recognized. Quantitative studies of these cells will undoubtedly be relevant to the pathologist, when correlated with pertinent physiological and biochemical data, for a better assessment of patients with disturbed gastric acid secretion. We believe that the introduction of the staining procedures reported here will render this quantitative approach a more easily attainable objective.

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