Inhibition of intestinal cell proliferation by villous cell extract

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SUMMARY In order to verify the hypothesis that intestinal cell proliferation is controlled by a mitotic inhibitor, extracts of villous epithelial cells from different species were analysed to study their effect on the proliferation of various intestinal cells. Villous extracts from rat and rabbit strongly and reversibly inhibited cell division and DNA synthesis in a rat intestinal epithelial cell line and a primary culture of rabbit intestinal epithelial cells. This non-cytotoxic, tissue specific but not species specific factor is present in both villous cells and crypt cells, with the highest concentrations occurring in the superficial epithelial cells. Assay of a partial purification of this factor showed that it has a molecular weight of approximately 190 000 daltons.

The mechanism controlling cellular proliferation in the small intestine has not yet been completely elucidated. The intestinal growth which occurs in man after intestinal damage, temporary ischaemia, irradiation, or in coeliac disease, termed type II intestinal growth response by Dowling, suggests that the functional zone of the intestine, the villi, controls the proliferative zone, the crypts. Furthermore, several experimental studies support the hypothesis that the digestive epithelium, like the liver, kidney, and haematopoietic tissue may synthesise a substance capable of interrupting cell proliferation during mitosis. Until now, however, studies aimed at showing the existence of such an inhibitor in the small intestine have been relatively uncontrolled. In many investigations, no attempt was made to show reversibility of action or absence of cytotoxicity. In several studies, the exact origin of cells providing the extract (crypt, villi, or epithelium) was not specified, and some authors have even used whole intestinal extracts. The best study to date was carried out on an intestinal epithelial cell line, but no data were provided concerning the effect of crypt cells on cell proliferation. Moreover, characterisation of the substance was incomplete, and species specificity was not discussed.

In our laboratory, the recent establishment of an intestinal cell line from rat fetus (IRD 98), whose characteristics readily prove its intestinal origin, provided us with the opportunity to study the control of intestinal epithelial cell proliferation.

Methods

ANIMALS AND RELATED TECHNIQUES Male Sprague-Dawley rats weighing 300–320 g were obtained from IFA CREDO (France). Male New Zealand albino rabbits weighing 2–2.5 kg from special breeding processes were also used. Culture products including medium, antibiotics and serum were purchased from GIBCO (Scotland). ＃H thymidine (44 Ci/mmol) and ＃C leucine (50 Ci/mmol) were obtained from the Commissariat à L’Energie Atomique (France). All other chemicals were obtained from commercial sources and were of the highest available purity. Several established cell lines, including rabbit kidney (RK 13), brain tumour (HeLa), dog kidney (MDCK), monkey kidney (BSC IL2), and mice embryo (FR3T3L1) were kindly provided by Dr Lefebvre (UER de Médecine, Nice, France); a human colon cancer cell line (HT 29) was provided by Dr J Marvaldi (Institut de Chimie Biologique, Marseille, France). A primary culture of rabbit epithelial intestinal cells (RIC) was isolated using the same method as for IRD-98. Alkaline phosphatase (EC 1.3.1.) activity was assayed as described by Louvard et al.

CELL CULTURE Cells were grown at 37 °C in a humidified 5 % CO2 atmosphere in the presence of Dulbecco’s modified Eagle’s medium supplemented with 10 % fetal calf serum, penicillin (50 U/ml) and streptomycin (50 μg/ml). The media were changed three times weekly. Cells were plated at 4.5 × 103 cells per cm2 in 60 mm...
plastic Petri dishes (Corning) for growth curve studies and in 35 mm dishes for \(^3\)H thymidine and \(^{14}\)C leucine incorporation. Cell counts were performed with a Coulter counter.

**PREPARATION OF CELL EXTRACT**

Male Sprague-Dawley rats were killed by ether inhalation, and male New Zealand albino rabbits by cervical dislocation. The entire small intestine was quickly removed and washed thoroughly with cold saline containing 1 mM DTT. Cells were then collected by the Weiser method\(^4\) and alkaline phosphatase activity was assayed for each fraction. Fractions 1 to 4 were considered to be villous cells and 7 to 10, crypt cells; fractions from each animal were pooled and homogenised in cold, deionised water (30 strokes of a Potter homogeniser and ultrasonication). Homogenates from 10 rats or two rabbits were combined and centrifuged at 900 g for 30 minutes at 4 °C. The supernatant was ultracentrifuged at 105 000 g for 60 minutes at 4 °C. An aliquot of the supernatant from the villous and crypt cell extract was sterilised by passage through 0.22 μm Millex filters (Millipore) before biological assay. The high speed supernatant was then saturated to 40 % by addition of solid ammonium sulphate, and stirred overnight at 4 °C. The suspension was spun at 40 000 g for 30 minutes at 4 °C and the pellet discarded. Crystalline ammonium sulphate was then added slowly to the supernatant to yield an 80 % saturation, and stirred overnight at 4 °C. The 80 % saturated suspension was spun at 40 000 g as described above, and the supernatant was discarded. The pellet (40–80 % fraction) was dissolved in a small volume of PBS, then dialysed extensively against 10 mM NaCl, 10 mM K_2HPO_4 at pH 7.4 (starting buffer). The 40–80 % fraction was then applied to a 1.6 × 40 cm column of DEAE Sephacel (Pharmacia). After sample application, the column was washed with bed volumes of starting buffer; the material retained on the column was then eluted with a linear, 0–1 M NaCl gradient in the same buffer. Protein elution was monitored by ultraviolet spectroscopy (Varian) at 280 nm. The 1 M NaCl peak was then concentrated in B15 Minicon Concentrators (Amicon). This concentrated fraction was then extensively dialysed against PBS and sterilized by passage through 0.22 μm Millex filters before biological assay. An aliquot of this fraction was applied to a 30 cm × 7.5 mm TSK G 3000 SW HPLC column (Kontron) and eluted with PBS (1 ml/min); protein elution was monitored by UV spectroscopy (Kontron) and each peak was collected (Gilson). Each fraction was concentrated and sterilised, as described above, before biological assay.

**ASSAY OF \(^3\)H THYMIDINE INCORPORATION INTO DNA OF CELLS**

The cell medium was replaced with fresh medium (1 ml) containing PBS (control) or various protein fraction concentrations (100–400 μg/ml) for various periods of time (three to 72 h). One hour before the end of each period, 1 μCi of \(^3\)H thymidine was added. The cell monolayers were then washed twice with cold PBS, treated with 5 % TCA (10 min at 4 °C) and washed three times with alcohol. The cell layer of each dish was dissolved in 0.5 ml of 0.1 N NaOH; aliquots of this solution were used to assess the radioactivity incorporated into TCA-precipitable material by scintillation counting (Intertechnique SL 2000), and to assay cellular protein by the method of Lowry et al.\(^5\) using bovine serum albumin as the standard. Absolute amounts of incorporated radioactivity were expressed as counts per minute per microgram of cellular protein.

**ASSAY OF \(^{14}\)C-LEUCINE INCORPORATION INTO CELL PROTEIN**

Cells were preincubated for seven hours with medium containing either a maximally effective concentration of villous or crypt cell extract, or PBS (control). The cells were then incubated for one hour in fresh medium containing extract and \(^{14}\)C-leucine (0.5 μCi/ml). Thereafter, the cell monolayers were treated as described above for \(^3\)H thymidine incorporation.

**Results**

**EFFECT OF VILLOUS AND CRYPT CELL EXTRACT FROM RAT INTESTINE ON THE GROWTH OF IRD-98 CELLS**

Figure 1 shows the effect of increasing quantities of villous and crypt cell extract from rat intestine on the proliferation of IRD-98 cells after 72 hours of culture. Maximal inhibition was obtained with 400 μg/ml concentrations of both villous and crypt cell extract, although the latter extract had a lesser effect. This phenomenon was confirmed by observing the action of the maximally effective extract dose (400 μg/ml) on IRD-98 cell growth over eight days (Fig. 2). Proliferation at day 8 was decreased by 42 % for the villous extract and by 33 % for the crypt extract relative to the control culture.

**EFFECT OF VILLOUS AND CRYPT CELL EXTRACT FROM RAT INTESTINE ON \(^3\)H-THYMIDINE INCORPORATION INTO DNA OF IRD-98 CELLS**

As shown in Figure 3, DNA synthesis in IRD-98 cells after incorporation of \(^3\)H-thymidine was inhibited...
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![Graph 1: Effect of various cell extract concentrations on the growth of an intestinal epithelial cell line, IRD-98.](image)

**Fig. 1** Effect of various cell extract concentrations on the growth of an intestinal epithelial cell line, IRD-98. Cells were seeded at time 0 in Petri dishes (60 mm dia.) in standard medium in the absence (black column) or presence of the indicated concentrations of crypt (white columns) or villous extracts (hatched columns). The medium was changed every 24 hours. At 72 hours, cell numbers were determined in triplicate cultures from each group. Results represent the mean cell number; SE was less than 10% of the mean.

![Graph 2: Effect of intestinal cell extracts on the growth curve of the intestinal cell line, IRD-98.](image)

**Fig. 2** Effect of intestinal cell extracts on the growth curve of the intestinal cell line, IRD-98. Cells were seeded at time 0 in Petri dishes (60 mm dia.) in standard medium in the absence (*) or presence of 400 μg protein/ml of villous (○ ○ ○) or crypt (● ● ●) extracts. The medium was changed every 48 hours. At the times indicated, cell numbers were determined in triplicate cultures from each group. Symbols represent the mean cell number; SE was less than 10% of the mean.

after eight hours by the presence of villous or crypt cell extract. This inhibition was dose dependent, the maximum inhibition of 90% being obtained with 400 μg/ml of villous protein. An activating effect was observed, however, with 100 μg/ml of protein.

![Graph 3: Effect of various cell extract concentrations on ³H-thymidine incorporation into IRD-98 cells.](image)

**Fig. 3** Effect of various cell extract concentrations on ³H-thymidine incorporation into IRD-98 cells. Cells were preincubated for 8 h with medium containing the indicated concentration of crypt extract (white columns) or villous extract (hatched columns) and incorporation of ³H-thymidine (0.5 μCi/ml) into DNA was determined. Control cells (black column) were preincubated with medium containing an equal volume of PBS. Results are the means of duplicate determinations; SE was less than 10% of the mean.

**KINETICS OF INHIBITION OF ³H-THYMIDINE INCORPORATION INTO IRD-98 CELLS BY VILLOUS AND CRYPT CELL EXTRACT FROM RAT INTESTINE**

Figure 4 reveals that DNA synthesis was rapidly inhibited: after one hour of incubation with 400 μg/ml of villous cell extract, DNA synthesis was 50% lower than in the control culture. Maximum inhibition (95%) was reached at 12 hours; inhibition then decreased gradually, dropping to zero after 48 hours. When extract was then added, inhibition again occurred. In contrast, replacement with fresh medium (Fig. 5) resulted in complete return to normal.

**TOXICITY**

As shown in Figure 6, preincubation of growing IRD-98 cells with rat villous and crypt cell extracts produced the expected drop in ³H thymidine incorporation (90% reduction for rat villous extract, 80% for rat crypt extract, compared with control), but had a negligible effect on ¹⁴C leucine incorporation into protein (20% reduction compared with control for villous extract, 0% for crypt extract). Viability, as
judged by trypan blue exclusion, was estimated to be 90% during culture with extracts.

**SPECIFICITY**

*Tissue specificity*

The effect of rat villous extract was tested by evaluating thymidine incorporation in several established cell lines. Rat villous extract had no effect on RK13, HeLa, MDCK, BSC1L2, FR3T3L1 or HT29 cells. Under the same experimental conditions, this extract was also tested on a primary culture of rabbit intestinal cells (RIC) isolated under the same technical conditions as IRD-98. The inhibition of DNA synthesis in this model (74%) was similar to that observed for IRD-98 (80%, Table 1).

A small intestine villous cell extract was prepared

![Graph](image-url)
Inhibition of intestinal cell proliferation by villous cell extract

Fig. 6 Effect of intestinal cell extracts on \(^3\)H-thymidine and \(^4\)C-leucine incorporation into IRD-98 cells. Cells were preincubated for 8 h with medium containing 400 \(\mu\)g protein/ml of crypt or villous extracts, and incorporation of \(^3\)H-thymidine (0.5 \(\mu\)Ci/ml) into DNA (hatched columns) and \(^4\)C-leucine (0.25 \(\mu\)Ci/ml) into protein (white columns) was determined. Control cells were preincubated with medium containing an equal volume of PBS. Results are the means of duplicate determinations; SE was less than 10% of the mean.

from rabbits, in the same manner as the rat extract, and was tested on the same cell lines. A similar decrease in \(^3\)H-thymidine incorporation (about 75%) was observed for IRD-98 and RIC. No inhibitory action was noted for RK13, HeLa, MDCK, BSC1L2, FR3T3L1 or HT29.

Organ specificity
The effect of extracts from various organs (liver, kidney, heart, lung), prepared under the same conditions, were tested on our two cell models; no significant inhibition was observed.

ATTEMPT TO PURIFY RAT VILLOUS CELL EXTRACT
Purification of the cytosolic villous extract from rat intestine was attempted. Essentially all of the inhibitory activity present in the high speed supernatant was recovered in the 40–80% ammonium sulphate fraction. Passage of this fraction on a DEAE column at a pH of 7.4 with a NaCl gradient from 0 to 1 M permitted us to isolate a fraction retained on the column and eluted with NaCl 1 M (Fig. 7). This fraction contained the inhibiting factor. After concentration and dialysis, the fraction was applied on a TSK 3000 HPLC column (Fig. 8). Elution isolated four peaks with molecular weights of 450 000, 190 000, 120 000, and 45 000 daltons. The inhibitory effect corresponded to the second peak, which had a
The factor we have isolated does not seem to be cytotoxic, as staining with trypan blue revealed conservation of cellular viability. Moreover, the action of this inhibitor is reversible; protein synthesis tested by incorporation of $^{14}$C-leucine was not modified during incubation in the presence of the factor.

This factor seems to be tissue specific; it inhibited proliferation of two intestinal epithelial cell cultures (the IRD-98 line originating in rat fetus small intestine and a primary culture of adult rabbit small intestine) but had no effect on tissue of extradigestive origin or on tissue from digestive tract tumours. No inhibitory activity was observed when cytosolic extracts of various other digestive and extradigestive tissues were tested on intestinal cell lines IRD-98 and RIC. By contrast, this factor does not seem to be species-specific: whether isolated from rat or rabbit intestine, it had a comparable effect on the proliferation of intestinal epithelial cells from either of these two species. It therefore satisfies the criteria for a chalone established by Bullough and described in several monographs.10,19,20

Purification of this factor from cytosolic villous cell extract showed that it was a negatively charged substance because it was retained to a great degree on DEAE. Its apparent molecular weight, as measured by gel filtration, is approximately 190,000. It has numerous similarities with the substance isolated by May et al.,10 which inhibits proliferation of IEC-6 cells.21 Whether this factor is related to this diamine-oxidase/ornithine-decarboxylase system has not been established.22 Diamine-oxidase is extremely active in mature villous tip cells, and might be a negative feedback inhibitor of crypt cell proliferation, but existing data are insufficient to permit firm conclusions on this point.

The identification of an endogenous inhibitor in this study supports the hypothesis of intrinsic control of crypt cell proliferation in the small intestine by a negative feedback mechanism. Rigorous purification and characterisation of this factor should provide better insight into the molecular basis of intestinal adaptation.

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