Prostaglandins and mucosal defensive mechanisms

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ABSTRACT: The first line of mucosal defence includes the juxtamucosal unstirred layer/pH gradient and the apical surfaces of the luminal epithelial cells. Many damaging agents, including nonsteroidal anti-inflammatory drugs (NSAIDs), can overwhelm these defences and destroy extensive regions of the luminal epithelium. This damage is readily tolerated in the normal mucosa. Furthermore, a combination of increased mucosal bloodflow, epithelial migration, mucus release, and efflux of bicarbonate-rich fluid usually allows rapid recovery of mucosal integrity. In the presence of vascular damage and congestion, however, luminal acid can kill mucosal cells and destroy the substrate necessary for repair (by epithelial migration). Damage of this type results in the production of hemorrhagic erosions, which may then develop into chronic ulceroinflammatory disease if healing is prevented by excess luminal acid or by impaired mucosal immune response. Endogenous and exogenous prostaglandins could affect all aspects of the mucosal defensive responses, from the juxtamucosal unstirred layer/pH gradient (via effects on secretion of bicarbonate, acid and mucus, as well as stimulation of fluid efflux) to the function of the mucosal immune system. Protection against the acute damage produced by topically administered NSAIDs or concentrated ethanol can result from either administration of prostaglandins or topical application of 'mild irritants'. This is referred to as 'adaptive cytoprotection'. Parenterally administered NSAIDs can also produce mucosal erosions. Protection against this type of damage may depend on the effects of prostaglandins on neural and contractile elements in the mucosa. Studies on animal models also suggest that by preventing acute hemorrhagic erosions, prostaglandins may prevent the development of chronic ulcer in susceptible individuals. Can J Gastroenterol 1990;4(3):95-107

Key Words: Adaptive cytoprotection, Cytoprotection, Gastric ulcer, Microvasculature, Mucus-bicarbonate barrier, Prostaglandin, Unstirred water layer

Les prostaglandines et les mécanismes de défense de la muceuse

RESUME: La première ligne de défense de l'estomac est le gradient de pH de la couche juxtamucuse non perturbée et la surface externe des cellules épithéliales. De nombreux agents nocifs, parmi lesquels les anti-inflammatoires non stéroidiens (AINS), peuvent anéantir ces facteurs de protection et détruire l'ouverture de l'ulcère. La protection contre ce type de dommages peut dépendre de l'effet des prostaglandines sur les cellules nerveuses et l'élargissement des cellules du tissu mous. Les études sur des modèles animaux suggèrent que la prophylaxie des lésions hémorragiques par les prostaglandines peut prévenir le développement de l'ulcère chronique chez les animaux. Can J Gastroenterol 1990;4(3):95-107

Key Words: Protection adaptée, Protection, Ulcère gastrique, Microvasculature, Barrière muqueuse-bicarbonatée, Prostaglandine, Couche non agitée

Suppression of acid secretion by H2 receptor antagonists remains the basis for current therapy of peptic ulcer. The most obvious and well-defined therapeutic value of prostaglandins in the treatment of peptic ulcer results from their antisecretory properties (1). By decreasing the quantity and concentration of secreted acid, prostaglandins can promote healing of peptic ulcer and minimize the production of acute damage produced by agents such as aspirin. When viewed solely in terms of ability to suppress acid secretion, prostaglandins do not offer significant therapeutic advantages over the H2 receptor antagonists. It is becoming apparent, however, that prostaglandins have other potentially beneficial effects on the mucosa, independent of their effects on acid secretion, which may justify their use in certain clinical settings as alternatives to other therapies. In particular, prostaglandins have been shown to affect the function of several aspects of mucosal defence. Gastrointestinal mucosal defences are dynamic rather than static barriers. Successive levels of defences must be overcome during the transition from a broken mucosal barrier to acute erosion and finally chronic ulceroinflammatory disease (Figure 1). By defining these defences and identifying the agents and mediators which overcome or impair them, it is possible to identify sites and stages at which the mucosa is vulnerable and at which therapeutic inter-
des régions importantes du revêtement épithelial. Ce dommage est facilement toléré dans des conditions normales. De plus, l'augmentation de l'apport sanguin au niveau des muqueuses, la migration épithéliale, la production de mucus et l'afflux d'un liquide riche en bicarbonates conjuguent leur action pour permettre habituellement à la muqueuse de retrouver rapidement son intégrité. En présence de lésion vasculaire et de congestion, néanmoins, l'acide gastrique peut détruire les cellules de la muqueuse et le substrat nécessaire à la régénération épithéliale. Si l'excès d'acide gastrique ou une défaillance de la réponse immunitaire de la muqueuse empêche la cicatrisation, il se produit des érosions hémorragiques qui peuvent ensuite résulter en une affection ulcéro-inflammatoire. Les prostaglandines endogènes et exogènes pourraient bien affecter tous les aspects des réponses défensives de la muqueuse, à commencer par le gradient de pH de la couche jumataqueuse non perturbée (par leurs activités sur la sécrétion de bicarbonates, d'acide et de mucus, ainsi que la stimulation de l'afflux liquide) et jusqu'à la fonction du système immunitaire de la muqueuse. La prévention de tout dommage grave provoqué par les AINS ou l'alcool concentré peut s'obtenir en administrant soit des prostaglandines soit des 'irritants doux' - c'est à dire par 'cytoprotection adaptative.' Les AINS pris par voie parentérale peuvent aussi entrainer l'érosion de la muqueuse. La protection peut dans ce cas dépendre des effets des prostaglandines sur les éléments nerveux et contractiles de la muqueuse. Certaines études chez l'animal suggèrent également qu'en prévenant les érosions hémorragiques aiguës, les prostaglandines empêchent peut-être le développement de l'ulcère chronique chez les sujets sensibles.

**Figure 1**) Diagrammatic representation of gastric mucosal defences and the types of damage which may result when successive layers of defences are overcome. The locations of various defences on the diagram (corresponding to sites in or external to the surface epithelium and within the mucosa, submucosa and muscularis) indicate the approximate depth of damage as each successive set of defences is overcome. The components of each defensive response (such as the immunologic barrier or accelerated cell turnover) can occur throughout the depth of the gastric wall.

Prostaglandins have trophic and immunomodulatory effects. They can: stimulate secretion of mucus and bicarbonate; increase mucosal bloodflow; affect electrogenic ion transport; and both stimulate and inhibit gastrointestinal motility (2). Prostaglandins are also produced in concert with many other mediators of inflammation such as leukotrienes and platelet activating factor, and may affect their production or actions.

Thus, prostaglandins can affect all stages in the progression of injury and may not only prevent the initial development of acute injury, but also aid in healing and preventing relapse.

**PROSTAGLANDINS AND CYTOPROTECTION**

The demonstration that prostaglandins have both antisecretory and anti-ulcer activities in animal models (1,3), and that certain synthetic prostaglandins are much more potent antisecretory agents than natural prostaglandins (4), suggested that these agents may be of therapeutic value in the reduction of gastric acid secretion. A subsequent study demonstrated that 'cytoprotective' prostaglandins could also prevent the development of acute hemorrhagic erosions produced in the gastric mucosa by necrotizing agents such as concentrated ethanol and strong acids (5). This study provided further support for a potential therapeutic role of these compounds in the treatment of ulcero-inflamatory diseases of the upper gastrointestinal tract. Since the cytoprotective properties of prostaglandins were manifested at doses below those which were antisecretory, it seemed that a new therapeutic approach was feasible.

It was also apparent that many agents other than prostaglandins were cytoprotective. When applied to the gastric mucosa prior to necrotizing agents, so-called 'mild irritants' were shown to produce 'cytoprotection' equivalent to that provided by exogenous prostaglandins (6). This gave rise to the concept of 'adaptive cyto-
protection'. Adaptive cytoprotection was initially found to be abolished by the pretreatment of animals with inhibitors of cyclo-oxygenase activity. This was interpreted as evidence that protection depended on irritant-induced stimulation of prostaglandin synthesis. Elevated prostaglandin synthesis has also been invoked as an explanation for the cytoprotective actions of agents such as sucralfate and the aluminum-containing antacids (7,8).

It has proven difficult, however, to provide convincing explanations for the mechanisms which underlie the phenomenon of cytoprotection. Cytoprotection by prostaglandins in the gastric mucosa is clearly not dependent on the preservation of the integrity of the surface epithelium in the face of assault by barrier breakers, but rather results from the prevention of microvascular disruption and the maintenance of gastric mucosal blood flow (9). In this sense, most experiments have demonstrated indirect cytoprotection where preservation of mucosal integrity is a consequence of effects on the mucosal microvasculature. Furthermore, adaptive cytoprotection can occur even when prostaglandin synthesis is significantly inhibited by prior administration of indomethacin (10-12). It is also necessary to consider the means by which agents other than prostaglandins can prevent microvascular disruption.

There is evidence that prostaglandins exert limited direct cytoprotection on in vivo gastric mucosal preparations and gastric cells which are isolated or maintained in cell culture (13,14). Such protection has been demonstrated against nonsteroidal anti-inflammatory drugs (NSAIDs), ethanol and taurocholate, and could play a role in the survival of certain populations of mucosal cells following exposure to damaging concentrations of these necrotizing agents (13-15). Studies on in vivo systems have demonstrated that direct cytoprotection of the luminal epithelium of the stomach against concentrated ethanol can be achieved by prior exposure to a mild irritant (11) or to sucralfate (12). This may be due to a 'histodilutional' effect (16), whereby the presence of an irritant-induced fluid efflux and edema, or a protective covering of sucralfate and mucus, attenuates the concentration of ethanol which comes into contact initially with the mucosa. There is no reason to believe that the luminal epithelium is rendered resistant to concentrated ethanol.

**PROSTAGLANDINS AND REPAIR OF SUPERFICIAL DAMAGE BY RESTITUTION**

Superficial damage to the stomach, of the type readily produced by barrier breakers such as acidified aspirin, sodium taurocholate, and concentrated ethanol may result in destruction of most of the luminal epithelium (Figures 2,3). Although this damage is irreversible at the cellular level, it is often restricted to luminal epithelial cells. If gastric mucosal microcirculation is not impaired, this type of damage is readily repaired or restored by cell migration from gastric glands (17). The restitution process is rapid and typically complete in about 1 h in both the stomach and small intestine.

Superficial injury is usually followed by increased gastric mucosal blood flow which, in combination with a cap of exfoliated cells and mucus, provides a localized alkaline microenvironment within which the influxing acid can be neutralized (17). Although it may be widespread, superficial damage to the epithelium must be considered a normal event and does not usually produce ulceration. Such injury is a result of the ready entry into cells of the undissociated lipophilic salicylate molecule, the detergent effects of bile salts, or osmotic damage in the case of hypertonic solutions. In the case of acetylsalicylic acid (ASA), once absorbed into the cells of the gastric epithelium, the molecule encounters a neutral pH and reverts to the ionized form, resulting in disruption of the intracellular ionic balance and localized accumulation of acid.

ASA also produces extensive and rapid damage to the intestine within minutes. In one study on humans, a
dose of ASA equivalent to two tablets in a glass of water damaged 25 to 30% of jejunal villi within 5 mins (18). Similar levels of damage have been produced in the proximal jejunum following exposure to intraluminal ethanol at peak concentrations of about 6% (19). As in the stomach, this type of damage was most frequent in the oldest cells (at the villus tips) and accompanied by blistering and rupture. Ivey et al (18) also found a 20% incidence of duodenal lesions by endoscopy in rheumatic disease patients receiving chronic ASA therapy. They noted that although enteric-coated ASA significantly reduced gastric mucosal damage, the incidence of duodenal erosions remained at about 20%.

Successful repair by restitution depends on either an absence of luminal acid or an intact mucosal microvascular supply which maintains a supply of bicarbonate-rich fluid to the overlying layer of damaged cells and released mucus. In the presence of vascular congestion, acid not only kills cells but also destroys the basal lamina (basement membrane), which is a substrate necessary for cell migration (17). Following exposure to prostaglandins, however, the mucosa is typically resistant to vascular damage and rapid repair can proceed. It is still not clear precisely how the administration of prostaglandins or mild irritants prevents necrotizing agents from causing vascular damage. It is clear, however, that the end result is the preservation of microvascular integrity.

ULTRASTRUCTURE OF GASTRIC MUCOSAL MICROCIRCULATION

The most revealing method for visualizing the architecture of the gastrointestinal microcirculation is that of scanning electron microscopy (SEM) of corrosion casts. With this technique, the microvasculature is perfused with saline followed by low viscosity plastic. The plastic is allowed to polymerize, the tissue digested, and the resulting cast viewed by SEM. The microvascular architecture of human and rat stomachs is basically the same (20,21). At the mucosal base, primary, secondary, and tertiary branches of mucosal arterioles give rise to an ascending capillary arcade (Figures 4, 5). The capillary arcade connects with the collecting venules in the upper regions of the mucosa (Figure 5). There are no lateral interconnections between the ascending capillaries and the collecting venules. However, the interconnections between the capillaries and the multiple connections of the capillaries with the collecting venules result in considerable redundancy (Figure 6).

An additional level of redundancy is introduced by the formation of large branches by many of the collecting venules. In the rat fundus, the authors found that almost half (44.8±3.4%) of the collecting venules were formed by fusion of major branches (Figure 7). More than 95% of these major branches...
Prostaglandins and mucosal defences

Figure 6) Scanning electron micrograph of luminal surface of corrosion cast of rat fundic mucosa. This view of the mucosal surface shows the honeycomb arrangement of the subepithelial capillaries and the interconnections between capillaries and a branched collecting venule. The four arrows indicate the major branches of the collecting venule and the direction of bloodflow (x170)

Figure 7) Scanning electron micrograph showing branching of collecting venules and venous plexus (vp) from rat fundic mucosa. This is a partial retrograde cast which was produced by infusion of plastic through the superior mesenteric vein. Perfusion was stopped before the capillary bed was filled (x100)

Figure 8) Scanning electron micrograph of venous plexus (vp). This is a view of the undersurface or serosal aspect of the mucosa. The vessels of the submucosa, muscularis and serosa were removed during preparation, and the image shows the interconnected nature of the venous plexus and the overlying capillary bed (x50)

Figure 9) Transmission electron micrograph of a subepithelial collecting venule after exposure of the mucosa for 10 mins to 16 mM ASA in 50 mM hydrochloric acid, and for a subsequent 5 mins to 50 mM hydrochloric acid. Note red blood cells (rbc), partially degranulated platelets (P), and fibrin (F) deposition (x8400)

Converged in the upper two-thirds of the mucosa. The collecting venules drain into an anastomosing collecting venule plexus which lies at the base of the mucosa, adjacent to the muscularis mucosa (Figures 7,8). The mucosal microcirculation can thus tolerate numerous localized sites of congestion or damage to capillaries or collecting venules. Cessation of bloodflow and local ischemia in the vulnerable, subepithelial region will only occur when all of the branches feeding a collecting venule have been damaged or congested. Similarly, the redundancy in the
underlying venular plexus ensures that drainage and perfusion will continue even when numerous individual collecting venules have become occluded.

**Sites of microvascular damage by topically applied aspirin:** When acidified aspirin is placed on the gastric mucosa it produces, within minutes, extensive subepithelial platelet thrombi in the collecting venules (Figure 9). Aggregated platelets actively synthesize thromboxane A2, which is a potent vasodilator and itself a promoter of further platelet aggregation. Whittle et al (22) showed that when local intravascular synthesis of thromboxane A2 was stimulated, exposure of the canine stomach to concentrations of acid and bile salt (100 mM hydrochloric acid and 5 mM sodium taurocholate), which were normally tolerated, resulted in extensive mucosal bleeding.

The impairment of hemostatic mechanisms, which is usually associated with aspirin ingestion, results from a systemic effect, via O-acetylation of platelet cyclo-oxygenase activity. However, in the gastric and upper intestinal mucosa, the ASA molecule need only penetrate a single layer of epithelium and a few microns of extracellular matrix before coming into contact with the endothelial cells of the subepithelial capillaries and collecting venules. ASA can therefore rapidly damage primarily the vascular endothelial cell and inhibit prostacyclin synthesis. Thrombus formation, congestion and local tissue damage, in the form of petechiae or punctate erosions, result.

**Effects of parenteral administration of NSAIDs and hypotension:** Parenteral administration of aspirin and indomethacin caused antral ulcers in the stomachs of animals when doses were used that gave plasma levels considered therapeutic in humans. The ulcers appeared in advance of signs of a broken barrier and were not related to the presence of salicylates in the gastric lumen, although the presence of luminal acid was necessary for lesion formation (23). Furthermore, Whittle (24) demonstrated that while neither 2 mM sodium taurocholate, nor any one of four subcutaneously administered NSAIDs (indomethacin, flurbiprofen, ASA, or naproxen), nor the vasoconstrictor noradrenaline, produced lesions, the combination of any of the NSAIDs or noradrenaline with acidified taurocholate was highly ulcerogenic. NSAIDs prevented the hyperemic response which normally follows topical damage. The result was the production of hemorrhagic erosions by a barrier breaker which would normally have been tolerated. The present authors found that when ASA (16 mM in 50 mM hydrochloric acid) was placed in the ex vivo gastric chamber for up to 30 mins, only small punctate erosions developed (unpublished data). Similarly, induction of hemorrhagic shock by withdrawal of blood to produce a rapid but transient decrease in mean blood pressure (to about 25 mmHg) produced few lesions and only temporary decrease in transmucosal potential difference. However, when ASA was placed in the chamber for 10 mins shortly after the induction of hemorrhagic shock, extensive ulcerations covering an average of 50% of the glandular mucosa resulted. These lesions first appeared as patches of intense focal pallor and were covered with a dense coagulum of extruded cells and mucus. This material covered regions of deep mucosal necrosis which ultimately became hemorrhagic in the presence of luminal acid. Such ulcers appear in the absence of diffuse damage and their genesis cannot readily be derived from consideration of the barrier-breaking hypothesis. They probably result from vascular blockade, perhaps due to vasconstriction at the mucosal base or in the submucosa.

**Effects of concentrated ethanol on the mucosal microcirculation:** The present morphological studies suggest that ethanol-induced hemorrhagic erosions resulted from congestion or occlusion at the point where the mucosal venular plexus is drained by veins which pass through the muscularis mucosa (Figure 10). This is a vulnerable site in the mucosal circulation. Blockade by compression or congestion of these draining veins would affect relatively large areas of the mucosa. After exposure to concentrated ethanol, the sites which were destined to become hemorrhagic erosions were clearly visible because of their congested blood vessels. Submucosal arteries, mucosal capillaries and collecting veins were typically congested at these sites (9), but veins in the submucosa below the exit from the muscularis mucosa were empty of blood.

Oates and Hakkinen (25) have suggested a sequence of events which may lead to the production of ethanol-induced gastric lesions. They suggested that ethanol may act by causing mast cells to rapidly degranulate and release vasoactive mediators such as leukotriene (LT) C4 and histamine. These mediators could then induce venous constriction and gastric muscle contractions. They observed that arterial and arteriolar dilation occurred within 15 to 20 s of exposure of the stomach to ethanol, and that the combination of strong arterial/arteriolar dilation and venous constriction produced vascular engorgement and visible hyperemia within 1 min. Greatly elevated capillary pressure in the mucosa and enhanced permeability can produce mucosal edema with a resultant lifting and blistering of the epithelium. The authors' studies are in general agreement with these concepts (26,27). The primary roles for mast cells and the production of LTC4, however, remain in doubt, since ethanol caused significant mast cell degranulation in cytoprotected mucosa (28) and there was no correlation between the ability of various cytoprotective agents to in-
hibit LTC4 synthesis and their ability to protect the mucosa (29).

LOCAL AND REFERRED CYTOPROTECTION: PROSTAGLANDIN-INDEPENDENT AND PROSTAGLANDIN-DEPENDENT

A phenomenon which the authors have defined as 'referred cytoprotection' occurs at sites in the mucosa which are not in contact with the protective agent (30). In these studies, an ex vivo rat gastric chamber preparation was used (27). This preparation allows direct viewing of the mucosa and monitoring of physiological indicators of mucosal integrity during the periods in which protective agents are present and during the development of hemorrhagic erosions. By tilting the gastric chamber preparation at an angle of about 30°, it is possible to cover half of the mucosal surface with 1.0 mL of luminal solution. A 5 min exposure of half of the mucosa to the mild irritant - 0.25 M hydrochloric acid, for instance - produced referred cytoprotection against 40% ethanol. There was a highly significant reduction in the area of hemorrhagic erosions on both the side which was exposed to the irritant and the side which was not exposed. The authors have studied the effects of several cytoprotective agents, and some, such as 4% sodium chloride, produced only local protection against ethanol, while others produced referred protection (Williamson et al, manuscript submitted).

This model allowed the examination of changes which occurred in the half of the mucosa not exposed to irritant, and testing of the hypothesis that adaptive cytoprotection results from an increase in release of mucus or bicarbonate secretion. Consequently, neither of these changes occurred on the uncovered side which, although not directly exposed to the protective agents, was still protected against 40% ethanol. In addition, there was no increased mucus release and no change in the thickness of the juxtamucosal unstirred/alkaline layer.

Referred protection against ethanol was, however, blocked by parental in-
ethanol significantly reduced the damage produced by these agents (12). Pretreatment of animals with indomethacin abolished the protective effects of luminal stasis, but this protection was restored by sucralfate. These studies suggested that impaired prostaglandin synthesis, ie, 88% inhibition of synthesis of 6-keto-prostaglandin F1α, increased the susceptibility of the mucosa in the same manner as did stirring of luminal solutions.

**THE pH GRADIENT AND CYTOPROTECTION**

The authors have studied the effects of protective mild irritants and luminal stasis on the thickness and magnitude of the juxtamucosal pH gradient using an ex vivo rat gastric chamber model and antimony microelectrodes. The microelectrodes had tip diameters of 40 µm and were mounted on a micromanipulator which is capable of movement in three dimensions and advancement of the microelectrode in 10 µm increments. An indifferent electrode was also positioned in the chamber and changes in hydrogen ion concentrations were recorded in millivolts on a Metrohm Herisau E512 pH meter/voltmeter. This apparatus is accurate to 0.05 pH units. Microelectrodes were calibrated daily using standard solutions of known pH. The beginning of the pH gradient was defined as the point at which the reading increased by 2 mV or more above that of the bulk solution (equivalent to a change in pH of about 0.05). The thickness of the pH gradient was defined as the distance between the start of the gradient and the point at which the microelectrode touched the surface of the gastric mucosa. Placement of the electrode and contact with the mucosal surface were determined visually with a modified dissecting microscope. The change in pH across the gradient was defined as the magnitude of the pH gradient.

All of the results described in this section were obtained while bathing the gastric mucosa with 50 mM hydrochloric acid. The gastric chamber contained stirred (200 rpm) 50 mM hydrochloric acid during the first three 10 min periods. During the fourth 10 min period the luminal solution was either stirred or unstirred. At the end of the fourth period, the luminal solution was removed and replaced with 10 mL of 50 mM hydrochloric acid, and measurements of pH gradient depth and magnitude were obtained within 5 mins. Each experimental group contained five animals. Four measurements of pH gradient depth and magnitude were obtained from two sites on each of the dorsal and ventral surfaces of each stomach. The sites from which measurements were taken were located midway between the greater and lesser curvatures, and were equidistant from the anterior and posterior margins of the chambered mucosa.

**Effects of luminal stasis on the pH gradient:***

Luminal stasis significantly (P<0.01) increased the thickness of the pH gradient from 410±20 µm when the solution was stirred to 820±30 µm (Figure 11). In the presence of stirring, the magnitude of the gradient was only 0.30±0.05 pH units. Luminal stasis increased the magnitude of the gradient to 1.10±0.05 pH units. However, despite this increase, the juxtamucosal pH was still less than 2.5 and the mucosa was protected in this instance without the production of an alkaline zone near the surfaces of the epithelial cells. Preliminary direct measurements of the thickness of the unstirred layer have been carried out, and it is likely that the present measurements of juxtamucosal pH gradient are also indicative of the thickness of the unstirred layer.

**Effects of indomethacin on the pH gradient:***

In earlier experiments the authors found that inhibition of prostaglandin synthesis by pretreatment with indomethacin resulted in the loss of the protection against taurocholate conferred by the 10 min period of luminal stasis (12). The authors' studies using the antimony microelectrodes demonstrated that indomethacin pretreatment significantly (P<0.01) reduced the thickness of the pH gradient by more than 50% to a mean of 360±20 µm (Figure 11). Indomethacin also reduced the magnitude of the gradient to 0.40±0.05 pH units.

These results indicate that impaired prostaglandin synthesis and the resulting increased susceptibility of mucosa to damage are at least partially due to a decrease in thickness of the unstirred layer/pH gradient.

**Effects of inhibiting bicarbonate synthesis on the juxtamucosal pH gradient and on mucosal susceptibility to acute damage:** The authors tested the effects of the carbonic anhydrase inhibitor, acetazolamide (100 mg/kg subcutaneously) on the chambered gastric mucosa. Acetazolamide pretreatment abolished the protective effect of luminal stasis against taurocholate and significantly reduced both thickness (to 270±50 µm) and magnitude (to 0.25±0.05 pH units) of the juxtamucosal pH gradient. These findings suggest that bicarbonate secretion is important in mucosal defence and that it is impaired by NSAIDs. They also suggest, however, that the beneficial effects of bicarbonate secretion may be due to effects on the unstirred layer as well as any effects resulting from elevation of pH at the mucosal surface since, even in the presence of unimpaired bicarbonate secretion, the pH at the mucosal surface is less than 2.5 (Figure 11).

**TROPHIC EFFECTS OF PROSTAGLANDINS**

Prostaglandins have marked trophic effects on the gastric mucosa. Administration of misoprostol to dogs for 11 weeks produced a 36% increase in stomach weight, primarily due to increases in length of the upper or foveolar region of the gland, and reflecting a significant increase in the gland cell production rate (33). Similar increases in thickness of the gastric mucosa, particularly in the foveolar regions, were also seen in humans after administration of 15(R)-15-methyl prostaglandin E2 for four months (34). The hyperplasia produced by prolonged treatment with prostaglandins resulted in increases in epithelial cells which secrete bicarbonate and mucus. This is the population which responds to luminal irritants and is responsible for mucosal defensive responses against...
agents such as bile salts and excess luminal acid. Although it can be hypothesized that the changes resulting from prolonged administration of prostaglandins should render the stomach more resistant to acute damage, the authors are unaware of any published results of such experiments.

PROSTAGLANDINS, MUCOSAL DEFENSES, AND DEVELOPMENT OF CHRONIC ULCER

About four to five million North Americans suffer from peptic ulcer disease at any given time, and as much as 5 to 10% of the population may be affected during an individual lifetime (35). There are associations between depressed levels of prostaglandin synthesis and the presence of chronic ulcer (36-38). The highest incidence of gastric ulcer is found among persons who ingest large quantities of NSAIDs, particularly the elderly (39,40). There
is little evidence for increased incidence of gastric ulcer in alcoholics even though acute exposure to high concentrations of ethanol produced superficial gastric damage in humans (41). Ethanol ingestion has, however, been associated with an increased incidence of duodenal ulcer (42). Moreover, the authors have shown that acute exposure to ethanol produced extensive, superficial damage to the duodenum and proximal jejunum (19). Cigarette smoking is associated with the presence of both duodenal and gastric ulcers (42,43), and there is convincing evidence that smoking produces a transient (less than 12 h) decrease in prostaglandin synthesis in the gastric mucosa (44). The effects of smoking on duodenal prostaglandin synthesis are less clear, although the deleterious effects of smoking on healing time and relapse rate are well established (45,46).

Several studies suggest that replacement or supplementation of endogenous synthesis of prostaglandins with exogenous sources may be beneficial in situations where high doses of NSAIDs are being used. The prostaglandin E1 analogue misoprostol has been shown to produce a significant reduction in NSAID-induced gastric lesions, a benefit which has not been demonstrated by H2 receptor antagonists (47). In addition, the reduction by misoprostol of indomethacin-induced increases in intestinal mucosal permeability in humans (48) raises the interesting possibility that exogenous prostaglandins may be beneficial in the treatment of NSAID-induced small intestinal inflammation.

**Cytoprotection and the transition from acute to chronic ulcer:** In considering the possible mechanisms by which prostaglandins may affect the defensive responses of the upper gastrointestinal tract, it is necessary to consider the type of damage which is under study. The majority of studies aimed at gaining an understanding of the etiology of chronic gastric ulcer have actually concentrated on the study of acute mucosal erosions, which are readily produced in animal models. These are prevented by prior administration of prostaglandins. Protection is provided in both humans and animals against a variety of damaging agents such as ASA and ethanol.

The inherent assumption in these studies has been that chronic exposure to the factors which produce acute erosions ('barrier breakers', acid, hypovolemia) will result in the development of a chronic ulcer. If prostaglandins act to prevent this acute damage, it is suggested that they will also prevent the development of chronic ulcer. In fact, it is now apparent that repeated exposure to agents or conditions which produce acute erosions does not necessarily result in chronic ulceration. Instead, the gastric mucosa may develop resistance or 'tolerance' to the ulcerogenic agents. This occurs in both humans and animals and has been demonstrated for ASA as well as for ethanol (49-52).

Only under exceptional circumstances does prolonged exposure to ulcerogenic agents produce, in animals, pathologies which bear some resemblance to human chronic ulcer. There is neither a naturally occurring analogue of human peptic ulcer nor a widely accepted animal model for chronic gastric ulcer, which can be used for studies of ulcer persistence and healing. The closest approximation to a chronic model is the acetic acid model in which the serosal application of concentrated acetic acid produces a long-lasting wound on the mucosal surface (53). Although it has no etiopathologic relevance to human gastric ulcer, the acetic acid model has proven useful in some studies of the effects of prolonged administration of prostaglandins and NSAIDs on mucosal healing (54,55).

**Mechanisms of development of chronic ulcer:** In the authors' studies on animal models of chronic ulcer—inflammatory disease, the following hypotheses to explain the development of chronic ulcer have been invoked.

- Chronic ulcer is initiated by acute damage, with consequent development of acute inflammation and increased mucosal permeability.
- Chronic inflammation depends on entry into the lamina propria of a luminal antigen which is not adequately cleared by the mucosal immune system.
- Conventional therapy may produce transient healing but recurrence of acute injury or enhanced permeability in the presence of the inciting antigen will result in relapse.

Products of microbial infections such as *Campylobacter pylori* (Figure 12) are a potential source of luminal antigens in the upper gastrointestinal tract, whereas acute erosions can have many causes, including the ingestion of NSAIDs. In addition, recent studies have demonstrated that NSAIDs produced increased permeability to macromolecules in the intestine and that prostaglandins have prevented this increased permeability (48). These effects occurred at sites in the bowel where decreased levels of luminal acid, due to the antisecretory effects of prostaglandins, would not have had a role.

Can cytoprotective prostaglandins prevent the development of chronic ulcer? The hypotheses outlined above predict that agents which prevent the development of acute hemorrhagic erosions should prevent the otherwise inevitable development of chronic ulcer—inflammatory disease. The abilities of misoprostol to prevent the development of NSAID gastropathy in humans and to reverse NSAID-induced changes in intestinal permeability are consistent with these concepts.

The authors have examined the ability of prostaglandins to prevent the development of chronic ulceration of the stomach and colon in two animal models which they have developed. They found that oro gastric intubation of fasted (18 h) female Sprague-Dawley rats with 1 mL of 40 or 50% (v/v) ethanol containing 50 mg of the hapten trinitrobenzene sulfonic acid (ethanol/TNB), produced one or more chronic type gastric ulcers (56). It was not necessary to use prior sensitization to produce these chronic lesions. In this model the concentrated ethanol produced acute damage and the hapten TNB, when complexed with tissue proteins, provided a poorly cleared antigen. Administration of either ethanol or
ethanol/TNB produced, within 1 h, extensive, linear, hemorrhagic erosions in the glandular mucosa of all animals. There was no difference at either 1 h or one day in the severity or appearance of the lesions produced by ethanol or ethanol/TNB. The acute damage produced by ethanol alone was totally healed, however, by about five days after administration, whereas 80% of the animals which received ethanol/TNB had one or more chronic type ulcers in the antral mucosa of the lesser curvature at least five to 10 days after administration (Figure 13). The ulcers were ovoid or circular and ranged in diameter from 2 to 5 mm. Linear, fissuring ulcers were also present in the corpus of some animals. The antral ulcers were associated with erythematous regions of gastritis, and open ulcers persisted for up to 15 days.

In one set of experiments, the authors examined whether pretreatment of rats with cytoprotective agents would prevent the development of the chronic type gastric ulcers. Two groups of 10 rats each were fasted for 18 h and then given 1 mL of either saline or saline containing 5 mg/kg of 16,16-dimethyl prostaglandin E2 (dmPGE2), followed 20 mins later by 50 mg of TNB in 1 mL 100% ethanol. After seven days the animals were sacrificed, and the number and size of chronic type ulcers in the gastric mucosa were determined. Pretreatment with dmPGE2 reduced the mean lesion number from 4.8 per animal to zero. In a similar experiment in which the effects of sucralfate (100 mg) and the PGE1 analogue rioprostil (100 µg/kg) were tested for their ability to prevent gastric ulcers, only rioprostil produced a significant decrease at day 5 after intubation with 2 mL 100% ethanol containing 50 mg TNB (Figure 14). In experiments on a model for chronic, transmural inflammation of the colon, the authors found that rioprostil, when given 20 mins before intracolonic infusion with ethanol/TNB, also significantly reduced the severity of the resulting chronic inflammation (57). In these models, the protective effects of the prostaglandins probably reflected their ability to prevent acute damage, and thus to decrease entry of the reactive hapten (TNB) into the lamina propria (58).

CONCLUSION

It is becoming apparent that the gastrointestinal mucosa relies for its defence on a series of dynamic responses. The beneficial effects of prostaglandins cannot be entirely explained with reference to their effects on only one aspect of this defensive repertoire. The abilities of prostaglandins to suppress acid secretion and modify NSAID-induced changes in mucosal permeability could aid in both the healing of chronic ulcer and prevention of relapse. Effects on mucosal bloodflow, mucus release and alkaline secretion could explain resistance to the development of acute erosions and accelerated recovery from acute damage. Similarly, protection against acute damage can also result from trophic effects and prevention by prostaglandins of the high amplitude gastric contractions which result from exposure to various ulcerogenic procedures and agents. These 'cytoprotective' effects may, by preventing acute injury, also prevent the initial stage necessary for the production of chronic ulcer or for relapse of healed ulcer.

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REFERENCES

70. Figure 14) Effects of pretreatment with a prostaglandin E analogue, with sucralfate and with saline vehicle, on the incidence of chronic type gastric ulcers present in rat stomachs five days after orogastric intubation with 50 mg of trinitrobenzene sulphonic acid in absolute ethanol. ** P<0.01


11. MacNaughton WK, Williamson TE, Morris GP. Adaptive cytoprotection by 0.25 M HCl is truly 'cytoprotective' and may not depend upon elevated levels of prostaglandin synthesis. Am J Physiol 1986;250:1075-81.


Lee CE, Menzies IS, Levi AJ. Miso-
prosrol reduces indomethacin-induced 
changes in human small intestinal per-

49. Graham DY, Smith JL, Spjut HJ, 
Torres E. Gastric adaptation. Studies 
in humans during continuous aspirin 
administration. Gastroenterology 

50. Tepperman BL, Soper BD. Prostaglandin 
E2 binding sites in porcine oxyntic 
mucosa: Effects of salicylates. Can J 

51. St John DJB, Yeomans ND, 
McDermott FT, de Boer WGRM. 
Adaptation of the gastric mucosa to 
repeated administration of aspirin in 

52. Ivey KJ, Tamawski A, Stachura J, 
Werner H, Mach T, Burks M. The 
induction of gastric mucosal tolerance 
to alcohol by chronic administration. 

53. Okabe S, Pfeiffer CJ. The acetic acid 
ulcer model. A procedure for chronic 
duodenal or gastric ulcer. In: Pfeiffer 

54. Okabe S, Takeuchi K, Honda K, Takagi 
K. Effects of acetylsalicylic acid (ASA), 
ASA plus L-glutamine and L-glutamine 
on healing of chronic gastric ulcer in the 

55. Wang JY, Yamasaki S, Takeuchi K, 
Okabe S. Delayed healing of acetic 
acid-induced gastric ulcers in rats by 
indomethacin. Gastroenterology 
1989;96:393-402.

56. Morris GP, Rebeiro L, Herridge MS, 
Szewczuk M, Depew WT. An animal 
model for chronic granulomatous 
inflammation of the stomach and 
colon. Gastroenterology 
1984;86:A1188. (Abst)

57. Wallace JL, MacNaughton WK, 
Morris GP, Beck PL. Inhibition of 
leukotriene synthesis markedly 
accelerates healing in a rat model of 
inflammatory bowel disease. 

58. Allgayer H, Deschryver K, Stenson 
WF. Treatment with 16,16′-dimethyl 
prostaglandin E2 before and after 
induction of colitis with trinitroben-
zenesulfonic acid in rats decreases 
inflammation. Gastroenterology 
1989;96:1290-300.