Mucosal adaptation to aspirin induced gastric damage in humans. Studies on blood flow, gastric mucosal growth, and neutrophil activation

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Abstract
The gastropathy associated with the ingestion of non-steroidal anti-inflammatory drugs (NSAIDs) such as aspirin is a common side effect of this class of drugs, but the precise mechanisms by which they cause mucosal damage have not been fully explained. During continued use of an injurious substance, such as aspirin, the extent of gastric mucosal damage decreases and this phenomenon is named gastric adaptation. To assess the extent of mucosal damage by aspirin and subsequent adaptation the effects of 14 days of continuous, oral administration of aspirin (2 g per day) to eight healthy male volunteers was studied. To estimate the rate of mucosal damage, gastroscopy was performed before (day 0) and at days 3, 7, 14 of aspirin treatment. Gastric micro-bleeding and gastric mucosal blood flow were measured using laser Doppler flowmeter and mucosal biopsy specimens were taken for the estimation of tissue DNA synthesis and RNA and DNA concentration. In addition, the activation of neutrophils in peripheral blood was assessed by measuring their ability to associate with platelets. Aspirin induced acute damage mainly in gastric corpus, reaching at day 3 about 3-5 on the endoscopic Lanza score but lessened to about 1-5 at day 14 pointing to the occurrence of gastric adaptation. Mucosal blood flow increased at day 3 by about 50% in the gastric corpus and by 88% in the antrum. The in vitro DNA synthesis and RNA concentration, an index of mucosal growth, were reduced at day 3 but then increased to reach about 150% of initial value at the end of aspirin treatment. The gastric micro-bleeding rate rose from about 0-38 ml/day at day 0 to about 7-7 ml/day at day 3 but then decreased significantly to virtually normal values at the end of the study. The neutrophil/platelet adherence showed significant increase during aspirin treatment. It is concluded that the treatment with aspirin in humans induces gastric adaptation to this agent, which entails the increase in mucosal blood flow, the rise in neutrophil activation, and the enhancement in mucosal growth.

The association between gastroduodenal injury and the use of non-steroidal anti-inflammatory drugs (NSAIDs) such as aspirin has been shown in experimental animals

1,2 How aspirin and other NSAIDs damage the gastroduodenal mucosa or delay its regeneration is not clearly understood. The deleterious effects of aspirin on gastroduodenal mucosa are generally attributed either to the direct damage of mucosal cells or to its ability to reduce the formation of prostaglandins.3 The inhibition of cyclo-oxygenase activity increases the susceptibility of gastric mucosa to injury while prostaglandins exhibit protective effects on the mucosa.4 Support for this mechanism is provided by the findings that NSAIDs affect various mucosal defence lines such as HCO3- secretion, mucus synthesis, and mucosal barrier.3,4 The effects of NSAIDs on mucosal blood flow, an important component of mucosal protection, remain controversial. In some studies mucosal blood flow was shown to be decreased by aspirin,5 whereas in others6 an increase in blood flow in the gastric mucosa was seen. On the other hand, aspirin and indomethacin have been shown to induce a microvascular damage7 and the extent of mucosal damage evoked by NSAIDs has not been closely correlated to the inhibition of mucosal cyclooxygenase activity.

Previous papers8-10 reported that the gastric mucosal injury in humans lessens or resolves despite the continued treatment with aspirin and other NSAIDs. This phenomenon of gastric mucosal adaptation associated with prolonged use of aspirin is well reported in animal studies11,12 but the mechanisms whereby the mucosa adapts to the damage remain uncertain.

Recently, it has been proposed that adherent leucocytes13 play an important part in the pathogenesis of NSAID induced gastropathy.14 This was supported by studies of Wallace et al15 and Lee et al16 who saw that the administration of monoclonal antibody against the leucocyte adhesion glycoprotein or anti-neutrophil serum largely prevented the microvascular injury caused by NSAIDs including aspirin in rats. Neutrophils release various mediators, such as proteases, and free radicals17 and their activation is mediated by newly recognised selectin P called platelet activation dependent granule external membrane protein (PADGEM).18 This selectin is responsible both for the platelet/neutrophil association and neutrophil/endothelium adherence. As activation of neutrophils seemed to be important in the development of gastric erosions after administration of indomethacin in rats,19 it has
become very interesting to assess the activity of neutrophils during the adaptation of the human stomach to aspirin.

NSAIDs increase the rate of proliferation in endoscopically normal gastric epithelium of patients with arthritis. This may also be one of the mechanisms underlying gastric adaptation to repeated NSAID treatment because a mucosal cell turnover is an important factor for gastric regeneration and mucosal integrity. The aim of this study was to evaluate the ability of human gastric mucosa to adapt to prolonged aspirin treatment and to find out if mucosal adaptive changes induced by aspirin are related to changes in mucosal blood flow, neutrophil activation, and mucosal growth.

Methods

SUBJECTS

Studies were carried out on eight healthy male volunteers between the ages 20 and 24 and weighing 67–80 kg. The subjects were requested to refrain from taking alcohol and drugs for two weeks before and during the study. All of them were in good health without any previous or present gastrointestinal disease and with normal laboratory values for blood chemistry and haematology. The volunteers were not requested to follow any dietary restriction or to modify their lifestyle during the period of treatment. All subjects had an assessment before the study including gastroduodenoscopy and C13 urea breath test for Helicobacter pylori.

This study was approved by the appropriate institutional review committee, and all subjects gave informed consent.

STUDY DESIGN

After the pre-study assessment, each subject received aspirin treatment. Oral aspirin was taken three times daily; one tablet 500 mg unbuffered aspirin (Bayer, Germany), was taken after breakfast at 0800, one at 1600, and two before bedtime as the daily total dose was 2 g. Similar treatment was continued for 14 consecutive days.

ASSESSMENTS

Subjects had an endoscopy and were studied for microbleeding, gastric mucosal blood flow, salicylate plasma concentration, and granulocyte/platelet interaction determined 24 hours before the start of aspirin treatment. The tests were then repeated at days 3, 7, and 14 of aspirin treatment, and additionally twice namely: two and four days after aspirin treatment had ended. Venous blood samples were withdrawn to measure plasma salicylate concentration as described before. Blood was taken for the measurement of platelet adherence to neutrophils. For this last purpose the 'rosette' formation test was used according to Jungi et al. Polymorphonuclear leucocytes (PMN) were isolated with the Ficol paque technique according to Byoum. Attention was paid to minimise the contamination of platelets. Therefore blood was collected to sodium citrate (3-2%) and first centrifuged to obtain platelet rich plasma. Such platelet rich plasma was further used for isolation of platelets by gel filtration when the residual erythrocytes/PMN and monocyte fraction was mixed with 3% solution of dextran. After sedimentation the upper fraction contained PMN/monocyte cells and this fraction was used for the Ficol gradient isolation of PMN. The residual erythrocytes were removed with hypo-osmotic lysis. Both washed platelets and PMNs were coincubated in Tyrode solution for 20 minutes at 22°C. The expression of PADGEM in platelets was induced by the addition of subthreshold proaggregatory concentration of thrombin (20–50 mU). The PMN were examined in both stimulated and thrombin unstimulated platelets under water immersion by photomicroscope (Zeiss, Berlin, Germany).

The rate of gastric microbleeding was measured as follows: each volunteer swallowed a 16 French gauge orogastric tube. The stomach was rinsed of debris with 100 ml of distilled water, then 100 ml of test solution was instilled into the stomach for a 10 minute washing period. After five minutes of each period 2 ml of phenol red as a marker in 10 ml of water was introduced by the orogastric tube and dispersed in the stomach. After 10 minutes of each washing period the gastric content was siphoned out and collected. Three successive washouts were performed at 10 minute intervals. Test solution or phenol red was introduced into the stomach and the subjects performed a standard series of manoeuvres designed to ensure contact with the whole gastric mucosa and to permit the blood to accumulate in gastric washing. Blood rinsed from the stomach and accumulated in the gastric washing was quantified by the peroxidase activity of the haemoglobin. Gastric microbleeding during each 10 minute period was calculated after the correction for phenol red recovery. Mean gastric microbleeding for three 10 minute washing periods was calculated and expressed as bleeding rate in ml/day. Phenol red was measured spectrophotometrically at pH 10-5 and a wave length of 560 nm. The pH of gastric washing was determined using glass electrode.

About 30 minutes after completion of gastric washing, the standard, unsedated, upper gastrointestinal endoscopy was performed by one investigator using an Olympus GIF Q panendoscope to evaluate the mucosal damage using the Lanza score system. Grading score was from 0 – normal to 4 – large area of submucosal haemorrhage with active bleeding or widespread involvement in the stomach. During endoscopy, samples of gastric juice were obtained by aspiration through the suction channel of the gastroscope into a trap. Care was taken to ensure the channel was clear of any contaminating fluid. About 5 ml aliquots of gastric juice were obtained from each subject, mixed with aprotinin (Trasylol,
Bayer, Germany) to prevent epidermal growth factor degradation and frozen at −70°C until radioimmunoassay using the technique described previously.27 Subjects were instructed not to swallow during and after gastric washings with all saliva being expectorated.

During endoscopy, the gastric blood flow was measured by the laser Doppler flowmetry technique using a novel generation of blood perfusion monitor (Laserflow, model BPM2, Vasamedics, St Paul, MN, USA).28 The flow probe (PR-436) was inserted down the biopsy channel of Olympus GIF Q panendoscope and after five minutes of stabilisation the blood flow was measured under direct vision in the gastric fundus and antrum. The values of the flow (ml/min/100 g) displayed on the digital panel meter were means of six measurements in the antrum and six in the corpus of the stomach, each measurement was made over a 10 second period required for the stabilisation of the recorded gastric blood flow. Two mucosal pinch biopsy specimens were taken from the gastric antrum and the corpus for the estimation of the DNA synthesis and RNA and DNA concentration in the gastric mucosa. The rates of DNA synthesis in the mucosal biopsy specimens were measured by incubating the tissue at 37°C for 30 minutes in Eagle’s minimal essential culture medium containing 2 μCi/ml [3H] thymidine (5 Ci/mmol Amersham, England). Tissue was gassed continuously with 95% oxygen-5% carbon dioxide during the incubation. The action was stopped with 0.4 N perchloric acid containing carrier thymidine at 5 mM. Samples were removed by being hydrolysed in 0.3 N KOH for 90 minutes at 37°C. DNA was precipitated with 10% perchloric acid. The RNA content of the supernatant was determined using the orcinol reaction.29 After standing on ice for 10 minutes, the DNA containing tubes were centrifuged, and the supernatant was discarded. DNA in the residual pellet was solubilised in 10% perchloric acid, heated to 70°C for 20 minutes. The DNA content of the samples was determined by the Burton procedure,30 as modified by Giles and Myers.31 The incorporation of [3H] thymidine into DNA was determined by counting 0.5 ml DNA containing filtrate in a Beckman liquid scintillation system. DNA and RNA content were expressed as micrograms per 100 mg mucosa and DNA synthesis was expressed as disintegrations per minute (dpm) per μg DNA.

**STATISTICS**

Results are expressed as mean (SEM). The significance of the difference between means was evaluated using analysis of variance followed by Duncan’s test with a confidence value at p<0.05.

**Results**

Eight volunteers entered the study and completed the aspirin treatment. The pre-treatment assessment of gastric microbleeding was about 0.38 (0.14) ml/day (Fig 1). After three days of treatment with aspirin, gastric microbleeding increased significantly (p<0.001) in all subjects reaching about 7.68 (0.52) ml/day. After seven days of treatment the microbleeding remained at a value that was not significantly different from that recorded after three days of treatment. Continuation of aspirin treatment for 14 days resulted in a significant attenuation of the rate of gastric microbleeding by about 50%. Two days after aspirin ingestion had stopped gastric microbleeding tended to return towards the pre-treatment value but was still significantly higher than the initial value.

The pH of gastric content was 5.8 (0.1) before the beginning of aspirin treatment, dropped down to 3.8 (0.1) at day 3 of aspirin treatment, and did not change significantly from this value during the rest of aspirin treatment.

After three days of treatment with aspirin (2 g per day) plasma salicylate concentration was 10.3 (1.3) μg/ml and remained close to this value in all volunteers up to the end of the aspirin ingestion.
blood flow in both, the gastric corpus by about 51% and gastric antrum by about 88%. Mucosal blood flow slightly and insignificantly dropped at day 14 (by about 23% in the corpus and by 24% in the antrum) but still remained significantly raised compared with the initial values. Blood flow, measured two days after aspirin ingestion was finished, was back to control values.

The concentration of epidermal growth factor in gastric juice measured before the start of aspirin treatment averaged 1·8 (0·2) ng/ml (Fig 4). After three days of aspirin treatment the concentration of epidermal growth factor was almost doubled (3·4 (0·6) ng/ml and reached the peak (7·2 (1·4) ng/ml) after seven days of treatment. After 14 days of treatment, epidermal growth factor concentration was still significantly raised (4·3 (1·8) ng/ml), but four days after aspirin administration had finished the concentration of epidermal growth factor in the gastric content was not significantly different from the initial value (2·1 (0·4) ng/ml).

Aspirin treatment caused significant changes in the incorporation of 3H thymidine into DNA (Fig 4). After three and seven days of aspirin ingestion DNA synthesis dropped by about 50% in both the gastric corpus and antral mucosa and was followed by a significant increase at day 14 with the highest value being reached two days after aspirin treatment ended. DNA and RNA concentrations tended to decrease at day 3 of aspirin treatment but this was not statistically significant (Table). At day 7 of aspirin treatment, RNA concentration returned to the pre-treatment value to reach a significant increase in both the gastric corpus and antral mucosa at day 14. The DNA concentrations were not significantly affected throughout the period of aspirin treatment.

In the venous blood drawn before the start of aspirin treatment the coincubation of platelets and PMN without addition of thrombin resulted in the formation of 15 (0·6) rosettes per vision field showing only a slight association of platelets with leucocytes during the isolation procedure (Fig 5). Addition of thrombin resulted in the increase of the rosette number up to 25·0 (0·7). After three days of aspirin treatment there was a significant increase of the rosette number in both unstimulated (22·0 (0·9)) and thrombin stimulated platelets (33·0 (1·5)). The number of leucocytes with rosettes was increasing during the whole aspirin treatment period and reached the highest value at day 14 (26·2 (1·6) thrombin unstimulated and 41·3 (2·2) thrombin stimulated platelets). Four days after aspirin treatment the rosette number did not differ from the initial value in both thrombin unstimulated and thrombin stimulated platelets.

Discussion
This study confirms earlier results that aspirin is harmful to the gastric mucosa \(^8\text{-}10\) but that the gastric mucosa shows a remarkable...
ability to adapt to repeated exposures to the injurious action of this drug so that microscopic damage inflicted on the first contact is minimised on repeated challenge.8–10

In our study continuous administration of aspirin for 14 days in healthy volunteers resulted in a statistically significant reduction in the extent of gastric damage as determined by endoscopic score and microbleeding from the stomach. Graham et al8 showed that aspirin in humans induced maximal gastric injury within three days of treatment but then the lesions tended to resolve despite continued administration of the drug. We also saw the most pronounced endoscopic damage at the third day of aspirin treatment and this was well correlated with a considerably increased rate of gastric microbleeding. The increase in microbleeding seen by us was greater than that reported previously34,35 but this could be explained simply by the higher dose of aspirin used in our study. This is in keeping with the finding of Pierson et al36 who found direct correlation between the dose of aspirin and the rate of gastric microbleeding.

The effect of aspirin on the mucosal blood flow, an important component of the mucosal defence system, remains controversial. Main and Whittle37 saw a reduction in mucosal blood flow in rats treated with indomethacin at the dose that inhibited prostaglandin formation. This could be consistent with a prostaglandin mediated mechanism, as prostaglandins are known to enhance mucosal blood flow.38 Kauffman39 found a decrease in basal gastric mucosal blood flow in conscious dogs given indomethacin. Both indomethacin and aspirin were also shown to induce rapid microvascular damage and subsequently to reduce mucosal blood flow so the fall in the gastric circulation could result from the widespread haemorrhagic necrosis.7 On the other hand, Ashley et al6 have shown that aspirin caused the reduction in gastric mucosal blood flow only at the site of damage, with enhancement of blood flow elsewhere with an overall increase in gastric blood flow. This is consistent with our study in which the flow measured in the microscopically unchanged mucosa was significantly increased during the whole aspirin treatment period. On the other hand Shorrock and Ree40 using a similar technique of blood flow measurement to ours but giving indomethacin to healthy volunteers saw a significant reduction in gastric mucosal blood flow after 24 hours administration of this agent and this decrease was closely correlated with maximal mucosal damage. Indomethacin, however, differs from aspirin and exerts a stronger influence on the gastric blood flow than aspirin both in animals and in humans.41

In this study, aspirin was used at a dose usually used in arthritic patients. Using this dose, the most profound injury was seen after three days, and in contrast with the study with indomethacin40 there was a significant increase in mucosal blood flow. The mechanism of this rather unusual finding of increased blood flow combined with visible mucosal lesions after aspirin in humans is unknown and suggests a difference between indomethacin and aspirin in the action on gastric circulation that is not dependent upon the inhibition of prostaglandin biosynthesis. Studies in rats42 showed that aspirin can damage both superficial and deep mucosal microvessels with a widespread destruction of endothelial cells within 15 minutes of injury. In humans, a single dose of aspirin caused evident gastric haemorrhages within one hour followed by widespread petechial haemorrhage and erosions several hours later.43 We measured the gastric blood flow before and then three days after the start of aspirin treatment so we could have missed an early reduction in gastric blood flow, which might have occurred after the first aspirin ingestion. The increase in the gastric blood flow seen after three days of aspirin treatment might be an early sign of gastric adaptation to this drug rather than the effect of mucosal damage.

This study confirmed and extended the finding that the human gastric mucosa adapts to continuous treatment with aspirin.8–10 The

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<td>420 (19)</td>
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Values are mean (SEM) of eight observations in eight subjects.
* = Significant changes compared with control before aspirin treatment.

Figure 5: The platelet/neutrophil adherence in eight subjects treated with aspirin for 14 days. Data expressed as mean (SEM). * Shows significant (p<0.05) change compared with control values.
mechanisms of this phenomenon are not fully understood but it has been suggested that an increase in the rate of proliferation in the gastric mucosa associated with the rapid renewal of the damaged epithelium play an important part. The rise in epithelial cell proliferation measured by 3H-thymidine uptake in response to four weeks of treatment with aspirin was previously reported in rat fundic mucosa. In our study on humans the early damaging effect of aspirin was accompanied by some reduction in DNA synthesis without significant changes in RNA or DNA concentrations but later on, despite the continuation of aspirin challenge, an appreciable increase in DNA synthesis and RNA contents was seen. Thus our study shows that the exposure of healthy human gastric mucosa to aspirin leads to initial widespread mucosal damage accompanied by a decrease in cell proliferation seen during the first days of aspirin treatment but then the mucosal regeneration starts. This process is remarkably impressive after 14 days of repeated treatment with aspirin as evidenced by the in vitro [3H]thymidine incorporation into DNA. DPA of gastric biopsy samples accompanied by the increase of RNA concentration in both corpus and antral mucosa. The fact that DNA synthesis increases but total DNA content does not suggests that there is no accumulation of cells probably because of simultaneous cell exfoliation. The increase in RNA content suggests that the cells that are present synthesise more protein and eventually other components, which may play a part in adaptation. Our results are in good agreement with a recent report of Levi et al. who found that NSAIDs given long term stimulate the mitotic activity of mucus in the patients with arthritis and misoprostol does not reverse this effect. It was proposed that the increase in mitotic activity in gastric glands seen after two weeks of NSAID treatment could be one of the mechanisms underlying gastric adaptation to NSAIDs but no attempts were made to measure the proliferation before the start and after the treatment with NSAIDs in the same subjects.

The factors participating in the enhanced mucosal proliferation during adaptation to NSAIDs have not been identified but Wright et al. reported that the damage of the human gastrointestinal tract may induce the development of novel cell lineages (from stem cells) that secrete locally epidermal growth factor and this epidermal growth factor could participate in this proliferation of mucosal cells after damage. Although there are no reports showing mRNA for epidermal growth factor in the normal mucosa, increased epidermal growth factor content in the gastric mucosa has been shown in patients with gastritis suggesting that epidermal growth factor mRNA is present although in small amounts in the intact mucosa. It is probable that epidermal growth factor may be produced locally in larger amounts after the mucosal damage. Indeed a considerable increase in the luminal content of epidermal growth factor was seen in our subjects treated with aspirin and this could contribute to the stimulation of DNA synthesis and mucosal cell proliferation. Although the salivary origin of this epidermal growth factor was largely excluded by aspiration of gastric luminal content, the possibility exists that the salivary epidermal growth factor entering the stomach was taken up by the mucosa and released into the gastric lumen upon the mucosal damage by aspirin. Although epidermal growth factor receptors are localised predominantly to the basolateral aspect of the cells in the proliferative zone, and after mucosal damage induced by aspirin, luminal epidermal growth factor may have access to these receptors to initiate cell proliferation and mucosal regeneration. This is supported in our study by the fact that the increase in epidermal growth factor production came before the rise in DNA synthesis and RNA content in the mucosa exposed to aspirin.

It has been reported recently that adhesion of platelets to neutrophils is selectively mediated by the newly recognised selectin P, called PADGEM. Expression of PADGEM on the endothelial cell membrane initiates the adhesion of neutrophils to endothelium and their migration. Wallace et al. and recently Lee et al. suggested that neutrophils play a crucial part in experimental damage in rats including the mucosal injury caused by NSAIDs. In our study in humans it was not possible to measure the adherence of leucocytes to the endothelium in the gastric mucosa. Therefore, we studied the activation of mucus and mucus cell enzyme. The rise in DNA synthesis and mucosal damage occurred but then the mucosal damage by aspirin was accompanied by aspiration of gastric luminal content, the possibility exists that the salivary epidermal growth factor entering the stomach was taken up by the mucosa and released into the gastric lumen upon the mucosal damage by aspirin. Although epidermal growth factor receptors are localised predominantly to the basolateral aspect of the cells in the proliferative zone, and after mucosal damage induced by aspirin, luminal epidermal growth factor may have access to these receptors to initiate cell proliferation and mucosal regeneration. This is supported in our study by the fact that the increase in epidermal growth factor production came before the rise in DNA synthesis and RNA content in the mucosa exposed to aspirin.
unstimulated and thrombin-stimulated platelets. As the adhesion of leukocytes to the endothelium is a prerequisite for their contribution to the blood into tissues the observed activation of leukocytes could contribute to the early aspirin-induced gastro-
titis but may also play a part in subsequent gastric adaptation to aspirin. Support for this idea is our finding that the peak increase in granulocytes activation occurred when the full adaptation to aspirin (decrease of endoscopic score and microbleeding) was seen. Additionally, the support for the implication of neutrophils in aspirin induced mucosal damage originates from the study of Wallace et al. who saw that administration of monoclonal antibodies directed against the leucocyte adhesion glycoprotein largely prevents the epithelial cell and microvascular injury caused by aspirin or indomethacin. Aspirin seems to enhance the adherence of neutrophils to the vascular wall by increasing the release of proinflammatory mediators including leukotriene B4 platelet activating factor, and others.14 16

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41 Ruppin H, Person B, Robert A, Domshke W. Gastric cyto-
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