Role of oxygen derived free radicals in platelet activating factor induced bowel necrosis

J P CUEVA AND WEI HSUEH

From the Department of Pathology, Children's Memorial Hospital, Northwestern University Medical School, Chicago, IL 60614, USA

SUMMARY The mechanism of tissue and cell injury in ischaemic bowel necrosis is unclear. The present study investigated the role of oxygen derived free radicals in the development of bowel necrosis using injections of platelet activating factor (PAF) into the mesenteric vasculature. Animals were pretreated with allopurinol or superoxide dismutase together with catalase, before administration of PAF. Superoxide dismutase/catalase markedly improved the PAF-induced lesions, indicating that most of the intestinal damage after PAF injection is because of the release of oxygen radicals. The major source of oxygen radicals is xanthine oxidase, as allopurinol ameliorated small bowel lesions. Pretreatment with allopurinol produced a significant (p<0.01) preventive effect on PAF induced hypotension. In contrast, superoxide dismutase/catalase did not alter PAF induced hypotension. Superoxide dismutase/catalase pretreatment improved PAF induced haemoconcentration and leucopenia, while allopurinol showed no effect.

Platelet activating factor (PAF), also named AGEPC or PAF acether, is an endogenous phospholipid mediator which is produced by a wide variety of cells and mediates a myriad of biological responses. Platelet activating factor is produced by ischaemic intestinal tissue and our previous work has shown that injection of PAF in rats produced necrosis of the intestinal tract. The lesions in our experimental model are morphologically indistinguishable from those of human ischaemic bowel necrosis or necrotising enterocolitis. The pathogenesis of bowel necrosis after PAF injection is probably not a result of thromboembolic phenomena but rather of release of secondary mediators, resulting in mesenteric vasoconstriction and subsequent ischaemia. The mechanism of tissue injury and cell necrosis, however, remains unclear. Recent evidence suggests that oxygen derived free radicals may be abundantly produced in ischaemic tissues, accounting for at least part of the damage observed. The major source of oxygen radicals in postischaemic tissues appears to be the enzyme xanthine oxidase, high levels being found in intestinal tissue. It is probable that oxygen radicals are generated during the postischaemic reperfusion period after PAF administration and play an important role in the development of bowel necrosis. In the present study, we investigated any attenuating effects of the ubiquitous superoxide dismutase/catalase and allopurinol (a xanthine oxidase inhibitor) on PAF induced bowel necrosis. Superoxide dismutase metabolises superoxide anions and catalase converts subsequently formed hydrogen peroxide to more innocuous substances.

Methods

RATS Male Sprague-Dawley rats weighing 275 (±50) g were anaesthetised with pentobarbital ip. The carotid artery and jugular vein were cannulated for blood pressure monitoring, drug administration, and blood sampling as previously described. PAF, (1-O-alkyl-2-O-acyl-sn-glycerol-3-phosphocholine, Calbiochem-Behring, La Jolla, CA, USA) was diluted from a stock solution in ethanol and stored at −70°C. Working solutions were prepared in 2.5 mg/ml bovine serum albumin saline solution. Four different pretreatment regimens were assigned to animal groups as follows: (a) pretreatment with 10 mg/kg superoxide dismutase and 10 mg/kg catalase...
(Sigma, St Louis, Mo, USA), dissolved in 0.9% saline, continuously infused at the rate of 0.01 ml/min, starting at 30 min before PAF administration, and continued for 150 minutes. (b) Infusion of vehicle (saline) only. (c) Intraperitoneal injection of 5 mg/kg allopurinol (Sigma), dissolved in dimethylsulphoxide (DMSO) (2.5 mg/ml), 30 min before PAF injection. (d) Injection of vehicle (DMSO) only. All animals received PAF (4 μg/kg) through aortic injection, proximal to the superior mesenteric artery, as previously described. The abdominal incision was covered with gauze moistened with warm saline, and the bowel was gently pulled out for examination with the aid of a Wild stereomicroscope. Blood samples were drawn immediately before PAF administration, and at 15, 45, and 105, and 150 minutes post-PAF injection.

At the termination of the experimental period the grossly discoloured bowel was examined and total circulatory supravital staining done by instillation of 2 ml 5% Evans blue. The length of non-stained bowel was recorded and sections were taken for histological examination to confirm the presence of necrosis. Histological assessment of intestinal injury utilised a 0 to 4 grading scale: 0 having normal histology, 1 showing epithelial cell loss injury at villus tips, 2 including loss of more than half of villi, 3 being damage extending to submucosa, and 4 having transmural necrosis.

Results

Within a few minutes after PAF injection (4 μg/kg), a vasoconstriction response of the small mesenteric vessels in the mesentery itself and the vessels on the serosal surface of the bowel developed (Fig. 1). Five to 10 minutes after PAF administration, the normal mesenteric pulsations totally stopped and many small serosal vessels became undiscernible. As early as 45 minutes, the mesenteric vessels began to recover.

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Fig. 1  PAF induced an initial phase of vasoconstriction, followed by a recovery phase. (a) Before PAF injection. Note the diameters of normal mesenteric vessels. (b) 30 min after PAF injection (4 μg/kg). Note vasoconstriction of the same mesenteric artery. (c) 150 min after PAF injection. Note recovery of the vasomotor tone of the same mesenteric artery. Similar changes were observed in serosal vessels (not shown).
from this constricting response, and between 100 and 150 minutes, most vessels showed partial recovery (Fig. 1). Some visible arterial pulsation also returned. Thus we conclude that the bowel was reperfused after an initial vasoconstriction phase after PAF injection.

The systemic effects of PAF (4 µg/kg) include an immediate drop in mean arterial pressure. This severe initial hypotension was observed in all groups of animals (Fig. 2). The initial severe hypotensive episode (approximately 15–30 minutes) was followed by a gradual rise, which eventually (at 150 minutes) reached 48, 45, 61%, and 36% of the original blood pressures in groups (a), (b), (c), and (d) respectively. The group receiving allopurinol pretreatment showed significant (p<0.01) recuperation of blood pressure after PAF administration in comparison with the other groups. The WBC count decreased an average of 44% 15 minutes after PAF injection and remained low in groups (b), (c), and (d) thereafter (Fig. 3). Interestingly, the group receiving superoxide dismutase/catalase (a) had a significantly (p<0.05) milder leukocenia. All groups also developed marked increase in haematocrit (Hct) 15 minutes after PAF administration which remained high in groups (b), (c), and (d) (Fig. 4). Pretreatment with allopurinol had no effect on PAF induced haemoconcentration while superoxide dismutase/catalase pretreatment (group a) significantly ameliorated the haemoconcentration (Fig. 4). Intraperitoneal injection of penobarbitol on sham operated controls had no significant effects on blood pressure, haematocrit, WBC count, and intestinal perfusion.

Both superoxide dismutase/catalase and allopurinol significantly (p<0.01) ameliorated intestinal perfusion (as assessed by supravital staining) and attenuated the necrosis (Fig. 5). Only 40% of the small bowel showed perfusion when PAF was given alone, while pretreatment with superoxide dismutase/catalase or allopurinol improved perfusion to 70%. The degree of necrosis (histological score) also improved. In most animals injected with PAF alone, necrosis involved most part of mucosa or submucosa, while animals receiving PAF plus a pretreatment regimen (superoxide dismutase/catalase or allo-
purinol) developed no or mild necrosis limited to villus tips.

**Discussion**

Ischaemic bowel necrosis, or necrotising enterocolitis, is a life threatening condition often associated with shock or infection. We have developed an animal model of ischaemic bowel necrosis associated with shock by injecting PAF into the mesenteric vascular bed. Although evidence has suggested PAF induced bowel necrosis is caused by intestinal ischaemia, probably as a result of vasoconstricting mediators, the cellular mechanism of developing necrosis remains unclear. It has been shown that ischaemic injury in heart, kidney, liver and intestine is largely caused by superoxide radicals generated during the reperfusion period. Xanthine oxidase has been proposed by some investigators to play an important role in the reperfusion injury.

In PAF induced intestinal ischaemia, a recovery from the initial vasoconstriction of the small mesenteric arteries and the intestinal arteries was observed, suggesting reperfusion of the intestinal tissue after the initial vasoconstriction phase. Thus, it is very likely that oxygen radicals are also involved in the pathogenesis of bowel necrosis in our model. The present study has indicated: (a) an important factor of tissue damage and cell necrosis in our model could be accounted for by release of oxygen radicals, as superoxide dismutase/catalase pretreatment markedly ameliorated the necrotic lesions. (b) The major source of oxygen radicals in this model is xanthine oxidase (probably within the intestinal tissue), as allopurinol, a competitive inhibitor of xanthine oxidase, significantly ameliorated PAF induced lesions. (c) The allopurinol group showed significant
recuperation of blood pressure post-PAF administration. This recuperation is probably caused by inhibition of cardiac xanthine oxidase and the observed severe hypotension is partially attributed to PAF-induced cardiac depression.\(^6\) Xanthine oxidase has been found in heart tissue, the source appears to be endothelial cells of capillary and small vessels.\(^7\) (d) Pretreatment with superoxide dismutase/catalase ameliorated PAF induced haemoconcentration, suggesting that oxygen radicals are involved in the development of systemic increase in vascular permeability. (e) Superoxide dismutase/catalase pretreatment also improved PAF induced leucopenia which presumably resulted from lung polymorphonuclear leucocytes sequestration. This observation suggests that oxygen radicals may also be involved in this process, although the detailed mechanism remains unclear. (f) Interestingly, allopurinol had no attenuating effects on PAF-induced haemoconcentration or leucopenia. This observation is different from those of the ischaemia reperfusion models reported by other investigators.\(^8\) This difference is probably because of the systemic effects of PAF in our model. Platelet activating factor has been shown to activate polymorphonuclear neutrophils (PMN) and induce superoxide release by these cells.\(^9\) The released radicals and other PMN products subsequently cause endothelial cell damage and an increase in vascular permeability. It may be that systemic effects such as leucopenia and increased vascular permeability in our model are partly accounted for by direct activation of circulating leucocytes by PAF, while much of the intestinal injury observed after PAF administration was a consequence of local formation of xanthine oxidase.

It has been shown that treatment with either allopurinol or superoxide dismutase prevents ischaemia-reperfusion induced neutrophil infiltration in the intestine,\(^1\) and depletion of granulocytes with neutrophil antisera or prevention of granulocyte adherence with a monoclonal antibody afforded protection against ischaemia reperfusion injury in the intestine.\(^2\) It is possible that part of the protective effect of allopurinol and superoxide dismutase was the result of prevention of oxygen radical mediated neutrophil infiltration. We do not believe, however, that neutrophils play an essential role in PAF induced bowel necrosis. This is because (a) there was a marked sequestration of neutrophils in the lung and a severe systemic leucopenia after PAF injection,\(^3\) and (b) preliminary experiments using Mustargen induced leucopenic rats failed to show any protective effect against PAF induced bowel injury (unpublished observation).

Our previous studies have shown that PAF injection resulted in development of bowel necrosis and this action was potentiated by endotoxin.\(^4\) The pathogenesis of bowel necrosis seems to involve release of vasoconstrictive mediators, such as peptide leukotrienes and noradrenaline,\(^5\) which results in prolonged intestinal ischaemia. Furthermore, we have recently reported that PAF was formed within bowel tissue in response to lipopolysaccharide.\(^6\)

Based on these previous observations and our present findings, we hypothesise on the pathogenesis of ischaemic bowel necrosis as follows: PAF is formed endogenously as a response to infection or release of endotoxin. The produced PAF releases peptide leukotrienes and other vasoconstrictive mediators in the bowel, which result in prolonged ischaemia and initial small bowel necrosis. The intestine is one of the richest sources of xanthine oxidase in the body\(^13\)\(^14\) and exists as a NAD\(^+\) reducing, xanthine dehydrogenase. This high activity is found primarily in the mucosal layer, with an increasing gradient of activity from villus base to tip.\(^15\) This phenomenon may explain our observation that bowel necrosis as a rule begins at the tip of the villus. It has been well established that during ischaemia, xanthine dehydrogenase is converted to xanthine oxidase\(^16\)\(^17\) and ATP is catabolised to hypoxanthine, which then accumulates locally in the tissue. We have also reported that vasodilating prostaglandins (PG) are also released after PAF administration.\(^18\) The released PGs and other vasodilators resulting in intestinal reperfusion. Reperfusion provides molecular \(O_2\) to react with the accumulated tissue hypoxanthine to generate oxygen metabolites. This large burst of oxygen radicals further damages tissue and causes more necrosis.

Our present studies indicate that most of the tissue damage in the bowel is a result of reperfusion injury caused by release of oxygen radicals. PAF, whether endogenously produced or exogenously administered, also acts on leucocytes to produce oxygen radicals. These toxic oxygen metabolites damage endothelial cells and produce increased vascular permeability. Our observations also suggest that oxygen radicals also play a role in PAF induced leucopenia and haemoconcentration. The detailed mechanism, however, awaits further study.

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