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Divergent patterns of matrix metalloproteinase activity during wound healing in ileum and colon of rats

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Abstract

Background—Uncontrolled and increased extracellular matrix degradation during early anastomotic repair in the intestine may reduce wound strength increasing the risk of anastomatic dehiscence.

Aims—To characterise the metalloproteinases present in intact and anastomosed ileum and colon to study their role in matrix degradation after surgery.

Subjects—Tissue extracts of uninjured, and of anastomosed rat ileum and colon at postoperative days 1, 2, 3, 7, and 90, were used.

Methods—Metalloproteinases were identified by gelatin and casein zymography. Aminophenylmercuric acetate (APMA) treatment was used to activate latent metalloproteinases.

Results—Both uninjured ileum and colon contained a 60 and 67 kDa activity, but a 54 and 72 kDa gelatinase was present in ileum only, and a 51 kDa activity in colon only. APMA treatment converted the 60 kDa protease to 54 and 51 kDa forms and the 72 kDa protease to the 67 kDa form. These gelatinases may correspond to latent and active forms of MMP 1 and MMP 2, respectively. Additional metalloproteinases were observed after anastomotic construction. Both ileum and colon contained 95 and 230 kDa gelatinases, which were converted to 83 and 76 kDa forms by APMA. They may be the latent and active forms of MMP 9, respectively. Gelatinolytic activities of 25 and 28 kDa were only found in anastomosed ileum. Caseinolytic activities were only found in ileum extracts and those were most prominent at day 1, 2, and 3 after surgery.

Conclusions—The metalloproteinase pattern in ileum and colon differ considerably suggesting that matrix degradation after anastomotic construction may also vary.

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Keywords: metalloproteinases, intestine, wound healing, rat, zymography, anastomosis.

Wound healing in the intestine is an area of ongoing research interest because its failure has potentially devastating consequences. Anastomotic dehiscence remains a major complication of gastrointestinal surgery with concomitant high morbidity and death rates. The integrity and mechanical strength of the intestinal wall is provided by a structural protein network in the submucosa. This connective tissue layer is largely composed of collagen and the turnover of this extracellular matrix component is therefore of considerable importance for anastomotic strength. Early anastomotic repair is mediated through close coordination of mesenchymal, epithelial, and endothelial cells, which together with rearrangement, growth, and differentiation are important for the tissue remodelling events during wound repair. These cells have to migrate through the existing connective tissue matrix and they achieve this by enzymatic solubilisation and partly degradation of collagen and other extracellular matrix components. In addition, the presence of protein degrading enzymes is necessary for removal of necrotic tissue. However, matrix degradation, although an intrinsic feature of the repair sequence, may at the same time constitute a hazard to wound strength because it may loosen the matrix that has to anchor the sutures. Identification of the enzymes normally present is a prerequisite for characterising abnormalities that arise – and should possibly be corrected – under adverse conditions. For instance, uncontrolled protease activity has been postulated as the mechanism responsible for reduced wound strength after anastomotic construction in an infected area. While we have shown an increased collagenolytic potential during early repair of anastomoses in the rat intestine, the matrix degrading enzymes present in the intact and anastomosed bowel wall have not yet been characterised.

The matrix metalloproteinases (MMPs) degrade a variety of extracellular matrix components and are considered to be the major enzymes responsible for the remodelling of tissue in a wound area. They include interstitial collagenase (MMP 1), gelatinases (MMP 2 and 9), and stromelysin (MMP 3). Increased MMP activities have been detected in epidermal wounds and in wound fluids of chronic ulcers. A sensitive method to identify MMP activity in tissues is by the use of zymography. It is based on the property of MMPs to degrade gelatin or casein gels. The advantage of this method is that both molecular weight and activity of the MMPs can be determined simultaneously: it has been used to characterise the numerous latent and active forms of MMPs in a variety of tissue extracts. In this study we used this technique to analyse the MMP profiles in intact ileum and colon, and in anastomoses at various times after operation.
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**Methods**

**Operative procedure**

Three month old male outbred Wistar/Cpb:WU rats, weighing 255 (8) g (mean (SD), n=18), were obtained from our own colony (Nijmegen, the Netherlands). The animals were housed in groups of two in Makrolon type 3 cages. Water and a standard laboratory chow (Diet AM II, Hope Farms, Woerden, the Netherlands) were supplied ad libitum. Body weight was recorded daily. The study was approved by the Animal Ethical Review Committee of the Faculty of Medicine, University of Nijmegen.

After a seven day pre-experimental period the rats were randomly divided into six groups. Three rats per group were used to check the reproducibility of the experiments. The animals in the control group were killed at day 0 to determine proteolytic activity in normal ileum and colon. The animals in the other groups were killed at 1, 2, 3, 7, and 90 days after surgery, respectively, to determine the activities in anastomotic tissue of both ileum and colon.

At the day of operation the rats were anaesthetised by an intraperitoneal injection of sodium pentobarbital (50 mg/kg). Surgery was performed using a Zeiss operation microscope. The abdominal skin was shaved, disinfected with 70% v/v ethanol, and opened by a median laparotomy of 4 cm. In each animal 1 cm of both ileum and colon was resected at approximately 15 cm proximal to the ileal-caecal junction and 3 cm proximal to the rectal-peritoneal reflection, respectively. An end to end anastomosis was constructed using eight single layer (8x0 Ethilon (Ethicon, Norderstedt, Germany) sutures. The first suture was placed at the resection edge, at the side where the intestine is connected to the mesentery. The second ligature was placed at 180° from the first one. Light tension was put on both sutures using small bulldog type haemostatic clamps resulting in an anterior and posterior anastomosis of equal length. At the anterior side three sutures were placed at 60° distance. To view the posterior side, the anastomosis was reversed pulling the second ligature behind the ligature at the mesenteric side. Both were put under tension again and the remaining three ligatures were placed at 60° distance. This way, all sutures were localised within 1 mm from the resection edge. After completion, the abdomen was closed using a 3x0 silk suture for the fascia and staples for the skin. The animals were killed by means of an overdose of sodium pentobarbital. A 5 mm sample containing the suture line was collected including the granulation tissue but also uninjured bowel wall.20 Resected segments were opened longitudinally, washed thoroughly with ice cold saline, weighed, and frozen in liquid nitrogen until further processing.

**Preparation of tissue extracts**

The resected segments remained frozen in liquid nitrogen during pulverisation in a Braun Microdisembrator. Approximately 25 mg of the pulverised tissue was suspended in one ml of 1% (w/v) Triton X-100, 0.5 M TRIS-HCl pH 7.6, 0.2 M NaCl, and 10 mM CaCl2 at 4°C. The suspension was frozen and thawed twice and centrifuged (40 000 g for 30 minutes at 4°C). The freeze-thaw cycle increased the amount of extracted protein of all samples with a factor 1.2-1.5 without activating the latent enzymes towards their active forms. The supernatant was collected and dialysed for 48 hours against an ice cold buffer containing 50 mM TRIS-HCl pH 7.6, 0.2 M NaCl, 5 mM CaCl2. The protein concentration of the extracts was measured using the bichinchoninic acid reagent.21 All extracts were stored at -80°C until needed for zymography. Before zymography part of the extracts was incubated with 1 mM p-aminophenylmercuric acetate (APMA) at 37°C for two hours to activate latent proteinases. The activation of the proteinases in the presence of APMA entails the conversion of the pro-enzyme form to its active lower molecular weight form by autoproteolytic cleavage of a peptide fragment from their amino-terminus.22

**Zymographic analysis of matrix metalloproteinases**

Gelatin (type A: from porcine skin; Sigma Chemical, St Louis, MO, USA) or casein (Sigma Chemical, St Louis, MO, USA) was added to standard Laemmli 7.5%-8% (w/v) sodium dodecyl sulphate (SDS)-polyacrylamide mixture23 at a final concentration of 1 mg/ml. The samples were diluted 1:1 in sample buffer consisting of 62.5 mM TRIS-HCl, pH 6.8, 10% glycerol, 2% SDS, and 0.05% bromphenol blue. After heating at 60°C for 20 minutes the samples (4 µg of protein per well) were loaded onto a 4% acrylamide stacking gel on a vertical mini gel (Bio-Rad Laboratories, Richmond, CA, USA). Molecular weight reference standards (Sigma, SDS-6H and SDS 7) were electrophoresed on each gel. The gels were cooled to 4°C during electrophoresis. They were run at 15 mA/gel while stacking and at 20 mA/gel during the separating phase until the bromphenol blue dye front had reached the bottom of the gel. The gels containing samples of the colon extracts or extracts of both colon and ileum (see Fig 4) were sometimes run 15 minutes longer to further separate the enzymes. After electrophoresis the gels were washed three times in 2.5% Triton X-100 (w/v) for 10 minutes at room temperature. After the gels were rinsed twice in a buffer containing 50 mM TRIS-HCl, 5 mM CaCl2, 0.1% Triton X-100, pH 7-8, they were incubated in the same buffer overnight at 35°C, stained with 0.1% Coomassie Brilliant Blue (w/v) in 40% methanol and 10% acetic acid for 30 minutes, and destained in the same solution without Coomassie Blue (destaining time 15 minutes, three changes). Proteolytic activities were visualised by clear zones indicating the lysis of gelatin or casein. Presence of true metalloproteinases was confirmed by adding 10 mM
Figure 1: (A) Time course of gelatinolytic activity after anastomotic construction in rat colon. Extracts (4 μg of total protein) of uninjured (U) and anastomotic tissue taken at 1, 2, 3, 7, and 90 days after operation were run on a gelatin zymogram. Note the increased activity of the 60 and 51 kDa bands in anastomosed tissue with respect to uninjured colon. The 230 and 95 kDa activities were only present during the first postoperative week (see results). Molecular weight standard markers are shown. (B) Activation of the colon proMMPs by APMA. The proMMPs of 230 and 95 kDa were partially or totally converted to forms of a lower molecular weight (see results). Molecular weight standard markers are shown.

EDTA or 1 mM 1,10'-phenanthroline to the buffers used after electrophoresis.

Results
At each time point tissue extracts from three rats were available. Figures 1-4 show representative examples of zymography. Each lane shows the activities present in an extract from an individual rat. The pattern of activities was completely reproducible in the other two animals. Reduced staining indicates areas of gelatin or casein degradation and permits identification of, respectively, gelatinase and caseinase activities. The gels containing colon extracts were run 15 minutes longer than those of ileum extract (see methods) to attain a better separation of the 50 to 60 kDa gelatinases. This was possible because colonic extracts did not contain gelatinases with molecular weights lower than 50 kDa (see later).

Several areas of lysis were observed when colonic extracts from both uninjured and anastomotic rat colon were subjected to gelatin zymography (Fig 1A). In uninjured colon, one major activity was present with a molecular mass of 60 kDa. Two less prominent bands of 67 and 51 kDa could also be observed. Two additional activities of 95 and 230 kDa were present in colonic anastomoses at one, two, three, and seven days after anastomotic construction. It was also noted that the activity of the 60 and 51 kDa bands was increased with respect to uninjured colon. The activity of the 67 kDa gelatinase did not seem to change at any time after anastomotic construction. At postoperative day 90, the 95 and 230 kDa bands had disappeared.

After treatment of the same colonic extracts with APMA, the gelatinases were partially or totally converted to forms of a lower molecular size (Fig 1B). Compared with untreated extracts, the 230 kDa band had disappeared and the 95 kDa lysis activity was reduced. New lysis bands of a molecular mass of 83 and 76 kDa were observed, indicating a precursor-product relationship between the high molecular weight sizes of 230 and 95 kDa and those with lower sizes of 83 and 76 kDa. Furthermore, APMA treatment also decreased the activity of the 60 kDa of all colonic extracts. At all time points measured, a new activity was noted at 54 kDa and the intensity of the 51 kDa band increased, particularly at postoperative day 90.

Both uninjured and anastomotic ileum demonstrated high gelatinase activities at 72, 67, 60, and 54 kDa (Fig 2A). The 67 kDa band possessed the highest activity and was the most prominent one day after anastomotic construction. A minor band of 95 kDa was detected at postoperative days 1, 2, and 3. In contrast with colonic extracts, it had already

Figure 2: (A) Time course of gelatinolytic activity after anastomotic construction of rat ileum. Ileum extracts (4 μg of total protein) from uninjured (U) and anastomoses taken at one, two, three, seven, and 90 days after operation were run on a gelatin zymogram. With respect to uninjured ileum the 72, 67, 60, and 54 kDa activities were particularly increased at one day after anastomotic construction. Three minor additional activities of 95, 28, and 25 kDa were present at postoperative days 1, 2, and 3 (see results). Molecular weight standard markers are shown. (B) Conversion of the proMMPs in ileal extracts to lower molecular weight forms by APMA. Two major activities of 67 and 54 kDa remained after activation (see results). Molecular weight standard markers are shown.
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Discussion

This study purports to identify the various gelatin and casein degrading metalloproteinases extracted from intestinal tissue before and after anastomotic construction. The results show obvious differences between ileum and colon. Several gelatinases were identified in both uninjured and anastomotic tissue from ileum and colon. Caseinase activity was only present in extracts from ileum. All activities appeared to belong to the metalloproteinase family because EDTA and 1,10-phenanthroline were able to inhibit their activity completely. Most of the metalloproteinases were present in the latent form, which could be activated by APMA.

Gelatinase patterns observed in healthy colon and ileum were only partly the same. Both contained a 67 and a 60 kDa gelatinase activity. However, a 72 and 54 kDa gelatinase was present in ileum only, and a 51 kDa activity in colon only. Furthermore, specific gelatinolytic activity in ileal extracts seemed to be higher than in colonic extracts as indicated by the increased intensity of the lysis bands. Previous results on basal collagenolytic activities in uninjured intestine, using fibrillar collagen as a substrate, have also shown a higher activity in ileum than in colon.7 These results show that the potential of both tissues to solubilise and degrade collagen or other extracellular matrix components could be different.

Ileum and colon may react differently to injury. Our data show that most of the pre-existing gelatinolytic activities were considerably increased in both ileum and colon during the first postoperative days. Raised MMP activities may loosen the extracellular matrix around sutures and thus contribute to the loss of anastomotic strength seen during this period.24 In addition, new activities were detected with respect to uninjured intestine. However, the latter seemed more pronounced in colon than in ileum during the first postoperative days and, furthermore, they persisted longer in colonic anastomoses. These results support the hypothesis that localised extracellular matrix degradation in the early postoperative period may be higher in colon than in ileum.25 Previous results have also shown that the increase in postoperative collagen synthesis disappeared seven days after operation. Extracts of days 1, 2, and 3 also contained two minor activities with a molecular mass of approximately 28 and 25 kDa. The bands were not present at days 7 and 90.

Incubation of the ileum extracts with APMA (Fig 2B) showed two major bands at 67 and 51 kDa, indicating the conversion of the 72 and 60 kDa to lower sizes. In anastomotic tissues at day 1, 2, 3, and 7 minor activity was observed at 83 kDa. The 95 kDa lysis activity was reduced and the 28 and 25 kDa activities remained unaltered with respect to untreated extracts.

To confirm the apparent differences between ileum and colon, extracts from one day old anastomotic tissue from both bowel parts were run on the same gel (Fig 3). In general, the gelatinase activities in ileum extracts were higher than in colonic extracts. At one day after anastomotic construction ileum extracts included very high activities of gelatinases between 60 and 51 kDa with respect to colonic extracts (Fig 3). Furthermore, very active 72 and 67 kDa bands were present in ileum but not in colon extracts.

Two major bands of lytic activity with molecular masses of 77 and 67 kDa were found when ileum extracts were subject to casein zymography (Fig 4). Both activities increased between one and three days after anastomotic construction but had decreased by seven days after injury. Caseinolytic activity was not observed with colonic extracts (not shown).

Addition of EDTA, as well as 1,10-phenanthroline to the zymography incubation buffer resulted in a complete inhibition of all activities detected in tissues of both ileum and colon.

Figure 3: A direct comparison between colonic and ileal anastomoses. Extracts of anastomotic colon (C1) and ileum (I1) at 1 day after surgery; C1+ and I1+ after APMA activation. Note the presence of the 72 kDa activity in ileal extracts and its absence in colonic extracts. The specific enzyme activities on protein basis were higher in ileal extracts than in colonic extracts. Molecular weight standard markers are shown.

Figure 4: Time course of caseinolytic activity after anastomotic construction in rat ileum. Extracts (4 µg of total proteins) of uninjured (U) and anastomotic tissue taken at one, two, three, seven, and 90 days after operation were run on a casein zymogram. Note the increased activity of the 77 and 67 kDa at postoperative days 1, 2, and 3 (see results). Caseinolytic activity was not observed with colon extracts. Molecular weight standard markers are shown.
is delayed in colon with respect to ileum. Together, these findings indicate that the postoperative phase characterised by localised collagen catabolism is possible longer lasting in colon than in ileum. If so, this may be part of the reason why large bowel anastomoses seem to be more prone to leakage than anastomoses in the small bowel.

Of the gelatinases found in the tissue extracts, the 95 and 230 kDa gelatinases were induced by anastomotic construction. They may correspond to the propeptide form of the MMP 9 (proMMP 9; 92-95 kDa) or may represent a multimer of the proMMP 9 (approximately 235 kDa), respectively. This protease (gelatinase B) is produced by neutrophils, macrophages, and some other cell types. The proMMP 9 can be converted by self cleavage or APMA activation to the active MMP 9 with a molecular mass of 83 kDa. It has also been reported that treatment with excess APMA may result in further cleavage to a 72-76 kDa gelatinase. Both the 83 kDa and the 76 kDa form were seen in our APMA treated colon extracts.

In contrast with MMP 9, which was only found in extracts from wound tissue, the other enzyme activities were already present in uninjured intestine, though with a far lesser specific activity. The 67 kDa band was the most prominent activity extracted from ileum, both before and after APMA treatment. This band probably corresponds with the active form of MMP 2. Its precursor, proMMP 2, with a molecular weight of 72 kDa was also present. It has been proposed that MMP 2 (gelatinase A or type IV collagenase) maintains collagen homeostasis in tissues. Wound fibroblasts of rat, but also granulation tissue of skin wounds synthesise MMP 2, but inflammatory cells do not.

Zymograms with highly purified preparations of collagenase from human synovial fibroblasts showed that these MMPs have a molecular weight of about 55 kDa (proMMP 1) and 35 kDa (MMP 1). Others showed that a porcine MMP of about 52 kDa, activated with APMA, was able to degrade collagen. Our data suggest that ileal and colonic gelatinases in the range of 51-60 kDa may correspond to various forms of this collagenase. Most of these proteases was present in a 60 and 54, and a 60 and 51 kDa form in ileum and colon, respectively. The activity of the 60 kDa was lowered by APMA treatment resulting in an increase of 54 and 51 kDa activities. However, we did not find the activated form of MMP 1. It may be that the conversion is prevented by tissue inhibitor of metalloproteinases (see later).

Potentially, there may be also other MMPs active with a molecular weight in the range of proMMP 1. ProMMP 3 (pro-stromelysin) has a molecular weight of about 57 kDa and is also very active in gelatin zymograms. However, when casein, a preferred substrate of (pro)stromelysin, was incorporated into the zymogram gels, no lysis bands of this molecular weight were detected. Thus, uninjured and anastomotic ileum and colon seem not to express stromelysin. On the other hand, two caseinase activities were found with molecular masses of 67 and 77 kDa in ileum at postoperative days 1, 2, and 3, but not in colon. It may be that the activated forms of MMP 2 and 9 have also a little caseinolytic activity, which could be seen if present on the gel in high concentrations.

Of the gelatinases extracted from ileum detected on the substrate gels, the 25-28 kDa activity remains unidentified. It might be the activated form of MMP 1, although it has a smaller molecular weight than reported earlier. Furthermore, the activity was expressed in extracts from anastomotic tissue of ileum only and not in extracts of colon. If it indeed represents MMP 1, the activity should also be expected in colonic anastomoses because proMMP 1 seems to be present in high concentrations here. Another possibility is that these MMPs correspond to MMP 7 or PUMP-1. This member of the MMP family with a molecular weight of 28 kDa has been found in rat uterus. Whether these intestinal gelatinases participate in extracellular matrix degradation cannot be answered at present. The fact that the activity of these proteases in ileum is higher in anastomotic tissue implies that they may be involved in matrix remodelling.

Gelatin zymography is useful in characterising the activation of most latent metalloproteinases. APMA treatment induced the conversion of 230, 95, 72, and 60 kDa forms to lower molecular weight forms. However, the transformation of two of them was incomplete. The 95 and 60 kDa activities still partly exist after three hours of incubation. It has been suggested that some tissue metalloproteinase activity (inter or intramolecular) is needed for a complete transformation. Also, the addition of exogenous tissue inhibitor of metalloproteinases (TIMP) to culture medium of chondrocytes resulted in the inhibition of most of the APMA induced conversions to lower molecular weights. It may be that endogenous TIMP is still associated with the proteases in our extracts, and thus limit the effect of the APMA treatment.

The findings of this study show that rat intestine contains MMPs that exist largely in an inactive form. Anastomotic construction increases their presence and additional MMPs are observed. Preliminary results in our laboratory on human rectum and colon showed a MMP pattern similar to that in rat colon suggesting the existence of an interspecies MMP system in the intestine. It remains to be established if patterns of anastomotic MMP activity change under conditions where the development of anastomotic strength is impaired and the risk for anastomotic failure increases. Such studies are currently under way. If this proves to be true, interventions aimed at counteracting activation of specific MMPs may possibly be derived to try to prevent anastomotic insufficiency.

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