Myeloperoxidase and elastase are only expressed by neutrophils in normal and in inflammed liver

Ahmad Amanzada · Ihtzaz Ahmed Malik · Martin Nischwitz · Sadaf Sultan · Naila Naz · Giuliano Ramadori

Accepted: 28 January 2011 / Published online: 16 February 2011 © The Author(s) 2011. This article is published with open access at Springerlink.com

Abstract  Myeloperoxidase (MPO) is involved in acute and chronic inflammatory diseases. The source of MPO in acute liver diseases is still a matter of debate. Therefore, we analysed MPO-gene expression on sections from normal and acutely damaged [carbon tetrachloride-(CCl₄) or whole liver γ-Irradiation] rat liver by immunohistochemistry, real time PCR and Western blot analysis of total RNA and protein. Also total RNA and protein from isolated Kupffer cells, hepatic stellate cells, Hepatocytes, endothelial cells and neutrophil granulocytes (NG) was analysed by real time PCR and Western blot, respectively. Sections of acutely injured human liver were prepared for MPO and CD68 immunofluorescence double staining. In normal rat liver MPO was detected immunohistochemically and by immunofluorescence double staining only in single NG. No MPO was detected in isolated parenchymal and non-parenchymal cell populations of the normal rat liver. In acutely damaged rat liver mRNA of MPO increased 2.8-fold at 24 h after administration of CCl₄ and 3.3-fold at 3 h after γ-Irradiation and MPO was detected by immunofluorescence double staining only in elastase (NE) positive NGs but not in macrophages (ED1 or CD68 positive cells). Our results demonstrate that, increased expression of MPO in damaged rat and human liver is due to recruited elastase positive NGs.

Keywords  Myeloperoxidase · Kupffer cells · Liver · Neutrophile granulocytes

Introduction

Myeloperoxidase (MPO), a heme protein, is a major component of azurophilic granules of neutrophil granulocytes (NGs). Optimal oxygen-dependent microbicidal activity depends on MPO as the critical enzyme for the generation of hypochlorous acid and other toxic oxygen products (Nauseef et al. 1988). The native enzyme has a molecular weight of 120 kDa (Zgliczynski and Stelmaszynska 1975). The MPO is a tetrameric molecule consisting of a pair of heavy- (~59 kDa) and light chains (~13.5 kDa) and two iron atoms (Olsson et al. 1972). There are two more components separated by electrophoresis at 40 and 20 kDa which consist of the autocatalytic products of the MPO (Taylor et al. 1992). The peroxidase activity of the enzyme is dependent on this chlorine group (Nauseef 1988). Through various enzymatic reactions within the NGs and interactions with other activated NGs, the MPO produces reactive oxygen species. The MPO in the lysosome causes the destruction of ingested organisms by binding to the walls of the microorganisms and locally producing hypochlorous acid. In the extracellular and
intravascular spaces secreted MPO also bacteria, viruses, or even the body’s own tumor cells. The MPO is an important marker for differentiating acute myelogenous leukaemia (Bennett et al. 1976). Multiple forms of MPO have been isolated from human myeloid leukaemia HL-60 cells (Yamada et al. 1981; Morita et al. 1986). MPO detection also has been used as a marker of neutrophil infiltration into tissues (Haqqani et al. 1999; Mcconnico et al. 1999). Reactive oxygen species produced by MPO are involved in the processes resulting in tissue damage. Hypochlorous acid is known to oxidize sulphydryl and thioether groups of proteins (Winterbourn 1985; Arnhold et al. 1993). MPO products are also involved lipid peroxidation. Peroxidation by hypochlorous acid is favored by the presence of hydroperoxides previously accumulated in lipid material (Panasenko et al. 1997; Panasenko and Arnhold 1999). MPO is thought to be involved in the pathology of diseases such as atherosclerosis, cancer, multiple sclerosis, rheumatoid arthritis and Alzheimer’s disease (Podrez et al. 1999; Nagra et al. 1997; Reynolds et al. 1999; Jolivalt et al. 1996; Weiss 1989; Daugherty et al. 1994; Malle et al. 2000).

The expression of MPO in Kupffer cells (KCs) has been reported in a few publications (Brown et al. 2001; Nahon et al. 2009; Rensen et al. 2009) but remains a matter of debate. Other reports claim that monocytes express MPO. Again this observation is controversial. Owen et al. (1994) and Akiyama et al. (1983) reported the presence of two different monocyte populations, one with and without MPO expression. Brennan et al. (2001) have shown that murine macrophages in atherosclerotic lesions do not express immunoreactive MPO. Several investigators have reported that MPO is not detectable after the differentiation of monocytes into macrophages (Kumar et al. 2004; Cribb et al. 1990; Nakagawara et al. 1981).

The hepatocellular inflammatory reaction induced by CCl₄-administration and γ-Irradiation is characterized principally by monocytes/macrophages and to a lesser degree lymphocytes and granulocytes (Knittel et al. 1999; Imhof and Dunon 1995).

The expression of MPO in normal and isolated parenchymal and non-parenchymal cells of normal rat liver, and acutely injured rat liver induced by either CCl₄-administration or γ-Irradiation was assessed in the present investigation. MPO-expression was identified only in granulocytes.

### Materials and methods

#### Animals

Male Wistar rats weighing of about 170–200 g body weight were purchased from Harlan–Winkelmann (Brochen, Germany). They were maintained under standard conditions with 12-h light and dark cycles and allowed ad libitum access to fresh water and food pellets consisted with the university’s policies and guidelines for the care and use of laboratory animals. This research use of rats was reviewed, approved, and overseen by the local ethics committee of the University of Goettingen as well as the public authority on animal welfare. The Human liver sections were used as well for the immunohistochemical studies and were obtained with full I. R. B. approval.

#### Antibodies

Rabbit polyclonal antibody against human MPO, mouse monoclonal anti-human macrophage CD68, Swine Anti-Rabbit Immunoglobulin HRP and Goat Anti-Mouse Immunoglobulin horseradish peroxidase were purchased from DakoCytomation (Berlin, Germany). Mouse monoclonal antibody against human MPO was purchased from Labgen CTS where located. Rabbit polyclonal antibody against human neutrophil elastase (NE) was purchased from Calbiochem (Bad Soden, Germany). Mouse monoclonal antibody against ß-actin, Anti-Mouse IgG (FITC conjugate) and Goat Anti-Rabbit IgG (TRITC conjugate) were purchased from Sigma (Steinheim, Germany).

#### CCl₄-induced liver injury

Rats were administered by gastric gavage 300 µl of a mixture of CCl₄ and corn oil (1:1 by volume) per 100 g body weight. Control rats received the same volume of corn oil alone. Animals were sacrificed 0, 3, 6, 9, 12, 24 and 48 h after CCl₄ administration. Their livers were either immediately frozen in liquid nitrogen or perfused to isolate liver cells. Treated animals (three for each time point) and controls (three for each time point) were sacrificed at 1, 3, 6, 12, 24, and 48 h after CCl₄-administration.

#### Whole liver γ-Irradiation in vivo

These experiments were performed as described previously (Christiansen et al. 2006): planning computed tomography was performed with a scanner (Somatom Balance; Siemens Medical Solutions, Erlangen, Germany) to delineate the location of the livers of each animal. The margins of the liver were marked on the skin of the animals, and a dose distribution was calculated. The rats were anesthetized with 90 mg ketamine per kilogram of body weight (Intervet, Unterschleissheim, Germany) and 7.5 mg/kg of 2% xylazine (Serumwerk Bernburg, Bern-burg/Saale, Germany) administered intraperitoneally. The livers were irradiated
selectively with 6 mV photons (dose rate of 2.4 Gy/min) using a Varian Clinac 600C accelerator (Varian, Palo Alto, CA); 25 Gy was delivered via an anterior–posterior/posterior-anterior (ap/pa) treatment technique. Treated animals (three for each time point) and sham-irradiated controls (three for each time point) were sacrificed at 1, 3, 6, 12, 24, and 48 h after irradiation.

Human liver tissues

Fresh human liver was obtained at the time of liver transplantation for end-stage or fulminant disease or resection of a mass lesion and was immediately stored at −70°C. Archived blocks of diseased human tissues were obtained from the Department of Pathology, Georg August University. This component of study was approved by the Institutional Review Board of Georg August University reviewing Human studies.

Immunohistochemistry and immunofluorescence double staining procedures

Five-micrometer cryostat sections were obtained, airdried, fixed with acetone (−20°C, 10 min), and stored at −20°C. After inhibition of endogenous peroxidase by incubating the slides with PBS containing 180 mg glucose, 5 mg glucose oxidase and 100 μl of 1 M sodium azide, they were treated with fetal calf serum (FCS) for 30 min to minimize nonspecific staining. The sections were incubated in a humidified chamber with the first antibody directed against either MPO, NE, ED1 or CD68 diluted in PBS at 1:100 for 1 h at room temperature. Negative controls were obtained by incubating with isotype-specific mouse/rabbit/goat IgGs instead of the specific primary antibody. After washing, the slides were covered with peroxidase-conjugated anti-rabbit/anti-mouse/antigoat immunoglobulins preabsorbed with normal rat serum to avoid cross-reactivity. Slides were washed and incubated with PBS containing 3,3′-diaminobenzidine (0.5 mg/ml) and H2O2 (0.01%) for 10 min to visualize immune complexes. Nuclei were counterstained with Meyer’s hemalum solution before the slides and coverslips mounted.

Isolation of liver parenchymal and nonparenchymal cells, conditions of cell culture

Hepatocytes and hepatic stellate cells (HSC) from rat liver were isolated and cultured as described previously (Knittel et al. 1992; Neubauer et al. 1995). Endothelial cells and KC’s of normal and injured liver were isolated as described by Knook and Sleyster (1976) with modifications (Armbrust et al. 1993). Centrifugal elutriation was performed to separate different cell populations (Beckman centrifuge J2-21, J-6B rotor at 2,500 rpm) and cells were harvested at the following flow rates: Endothelial cells—12 ml/min, small KCs—28 ml/min and large KCs—55 ml/min. Cells were collected in the respective culture medium (for KCs: M199, 15% FCS), 100 U/ml penicillin, 100 mg/ml streptomycin; for endothelial cells: DMEM, 15% FCS, 1% L-glutamine, 100 U/ml penicillin, 100 mg/ml streptomycin, Seromed, Berlin, Germany) and plated, or further processed for RNA isolation.

Neutrophilic granulocytes were isolated and cultured as described earlier (Boyum 1968).

RNA isolation and quantitative using real time RT–PCR

Total RNA was extracted from cultured cells by guanidine isothiocyanate and purified by caesium chloride cushion ultracentrifugation (Chirgwin et al. 1979). For real-time PCR, reverse transcription of RNA samples was performed using the Superscript kit from Invitrogen (Groningen, Netherlands) and following the manufacturer’s instructions. Real-time PCR analysis of cDNA was performed at 60 to 95°C for 45 cycles in the Sequence Detection System of ABI Prism 7000 (Applied Biosystems, Darmstadt, Germany) following the manufacturers’ instructions and by using SYBR Green Reaction Master Mix (ABI Prism; Applied Biosystems) and the primers listed in Table 1. All primers were synthesized by Invitrogen. In every RNA sample, β-actin mRNA was measured as the housekeeping gene (Rat 18 s was also measured with very similar results). Values were then compared with those obtained by using the control-RNA obtained from each animal series. The results were normalized to the housekeeping gene and

| Primer          | Forward     | Reverse                  |
|-----------------|-------------|--------------------------|
| MPO             | 5′-ACCTACCCCACTACCAGATCC-3′ | 5′-AACTCTCCAGCTGGCAAAAA-3′ |
| NE              | 5′-CTTGTGAAACCGCTTTGAC-3′ | 5′-CACATTGAGCTCTTGAGCA-3′     |
| CD11b           | 5′-CACAGTACACCAGAACTGCA-3′ | 5′-ATCACAGCTGGCTTAGAC-3′     |
| β-actin         | 5′-ACCACATGTACCACAGGATT-3′ | 5′-CCACACAGATCTTGCGTCA-3′     |
| Rat 18 s        | 5′-AAACGGCTACCACATCCAAG-3′ | 5′-CCTCAATGGATCTCCTGTA-3′     |

---

Histochem Cell Biol (2011) 135:305–315 307 123
fold change expression was calculated by using threshold cycle (Ct) values. For calculation of the relative changes, gene expression measured in control animals sacrificed at the same time points was set at one.

Western blot analysis and SDS-PAGE

Lysates were prepared by homogenization at 4°C in 20 mM Tris–HCl (pH 8.0), 5 mM EDTA, 3 mM EGTA, 1 mM DTT, 1% SDS, 1 mM PMSF, and protease inhibitor cocktail (Sigma–Aldrich Inc.). Samples were cleared by centrifugation at 15,000 g for 15 min at 4°C, and the protein concentration was measured by BCA assay (Pierce, Rockford, IL, USA), using bovine serum albumin, as the standard. Protein lysates were separated on SDS-polyacrylamide gels, electrotransferred to polyvinylidene difluoride membranes (Invitrogen, USA), and probed with primary antibodies overnight. The appropriate peroxidase-conjugated secondary antibodies (DAKO, Glostrup, Denmark) in a dilution of 1:5,000 was then added and incubation continued for 1 h, at room temperature. Bound antibodies were visualized using chemiluminescent substrate (ECL; Amersham-Pharmacia, UK). Equal loading was previously controlled by transient Ponceau S staining.

Results

Expression of MPO in normal rat liver tissue

By indirect immunohistochemical labeling (antibodies against MPO, NE, and ED1), it was possible to detect only a few cells by MPO (Figs. 1a, 2b) and NE (Fig. 1c) and macrophage/KCs by ED1 (Figs. 1b, 2a) in normal rat liver. The MPO+ and NE+ cells were distributed diffusely. Significantly more ED1+ cells were detectable as MPO+ and NE+ cells. The MPO+ and NE+ cells were scattered, mostly round and had little cytoplasm. ED1+, macrophage/KCs had a stellate or spindle shape.

Fig. 1 Immunohistochemical detection of myeloperoxidase (MPO), neutrophil elastase (NE) and ED1 cells in rat liver 6 h after one single oral dose of CCl4-administration induced rat liver injury. MPO+ cells (a), ED1+ cells (b) and NE+ cells (c) in control rat liver. MPO+ cells (d), ED1+ cells (e) and NE+ cells (f) at 6 h after treatment. The lower panel shows MPO+ cells (g), ED1+ cells (h) and NE+ cells (i) at 24 h after treatment. Original magnification, ×100. G original magnification, ×200 173 × 116 mm (300 × 300 DPI)
Expression of MPO-gene in parenchymal and non-parenchymal liver cells

After immunohistochemical labelling the expression of MPO from isolated liver cell populations and NG at the level of RNA and protein by real time PCR and Western blot was assessed. By real time PCR (Fig. 3), the $C_t$ value of MPO in NG was 31.3. The $C_t$ value of MPO in small KCs was 36.7, in large KCs 34.8, in hepatocytes 34.6, in endothelial cells 35.7 and in HSC 36.8. Consistent with the results of real time PCR, expression of MPO by Western blot was evident only from NG (Fig. 4). The parenchymal and non-parenchymal cells of the liver did not express MPO.

Expression of MPO in CCl₄ and γ-Irradiation induced rat liver injury

Two different models of acutely induced liver injury with either CCl₄ or γ-Irradiation were utilized. Indirect immunodetection in liver sections after CCl₄ and γ-Irradiation

![Figure 2](image1.png)  
**Fig. 2** MPO$^+$ and ED1$^+$ cells in rat liver after γ-Irradiation induced rat liver injury. ED1$^+$ cells (a) and MPO$^+$ cells (b) in control rat liver. ED1$^+$ cells (c) and MPO$^+$ cells (d) at 3 h after γ-Irradiation. ED1$^+$ cells (e) and MPO$^+$ cells (f) at 24 h after γ-Irradiation. Original magnification, ×100 173 × 174 mm (300 × 300 DPI)

![Figure 3](image2.png)  
**Fig. 3** MPO-gene expression in different cells analyzed by amplification of total RNA extracted from isolated cell populations of normal rat liver. Comparison of $C_t$ values of MPO in NG, small (sKC) and large KC (lKC), hepatocytes (HC), endothelial cells (EC) and hepatic stellate cells (HSC). Results were obtained by real time PCR analysis of total RNA. Results represent three experiments (in duplicate) and mean ± SEM values are shown for each cell type 46 × 34 mm (300 × 300 DPI)
were carried out with the antibodies against MPO, NE and ED1 followed by peroxidase and immunofluorescence double staining.

In the acutely injured rat liver quantitatively more MPO\(^+\) and NE\(^+\) and ED1\(^+\) cells were detected than that present in normal rat liver.

In CCl\(_4\)-induced liver injury, a diffuse increase in the number of MPO\(^+\) (Fig. 1d, g) and NE\(^+\) (Fig. 1f, i) cells in the liver parenchyma by indirect immunohistochemical staining. The increase on MPO\(^+\) and NE\(^+\) cells was achieved at 24 h after CCl\(_4\) administration (Fig. 1g, i). An increase in the number of ED1\(^+\) cells was detectable at 6 and 24 h around the portal vein (Fig. 1e, h).

Immunofluorescence double staining with the same antibodies in CCl\(_4\) treated Animals also demonstrated an increase of MPO\(^+\) (Fig. 5b, d), NE\(^+\) (Fig. 5a, c) and ED1\(^+\) cells. But the ED1\(^+\) cells were not MPO positive. These results were confirmed utilizing real time PCR for MPO (Fig. 6a) and NE (Fig. 6b) gene expression at the RNA level and for MPO at the level of using protein by Western blot analysis (Fig. 7a). The real time PCR analysis showed an increased expression of MPO, 2.0-fold at 6 h and 2.7-fold at 24 h after CCl\(_4\)-administration (Fig. 6a). The gene expression of NE showed similar results after 6 h (2.9-fold change) and 24 h (2.0-fold change) after CCl\(_4\)-administration (Fig. 6b). An increase of MPO protein at 24 h after CCl\(_4\)-administration was confirmed by Western blot (Fig. 7a).

By indirect immunohistochemical labelling, it was possible to detect an early increase in the number of MPO\(^+\) and NE\(^+\) cells around portal vessels of the rat liver after \(\gamma\)-Irradiation. The accumulation of MPO\(^+\) (Fig. 2d) and NE\(^+\) (data not shown) cells reached a maximum at 3 h. The immunofluorescence double staining studies utilizing \(\gamma\)-irradiated rat liver showed an increase in the number of MPO\(^+\), NE\(^+\) and ED1\(^+\) cells. The increase of MPO\(^+\) (Fig. 8c, e) and NE\(^+\) (data not shown) cells was detectable principally in the portal area. ED1\(^+\) cells were not MPO\(^+\).

At the same time, an increased of MPO and NE gene expression at the level of RNA and of MPO at the level of protein was detected by real time PCR analysis and Western blot, respectively. The real time PCR analysis

---

**Fig. 4** Western blot analysis of total protein of NG, small KC’s (sKC), large KC’s (lKC), hepatocytes (HC), endothelial cells (EC) and hepatic stellate cells (HSC). Cells were isolated and cultured for 24 h. Protein was then extracted, 20 g of total protein were separated by SDS–PAGE, and blotted onto PVDF-membranes. The membranes were subsequently incubated with the antibodies against MPO (upper panel) and s-actin (lower panel). The molecular weight of MPO is 59 and 13.5 kDa. In addition, autocatalytic products (AP) can be seen at 40 and 20 kDa (data not shown). The molecular weight of \(\beta\)-actin is 42 kDa 129 x 41 mm (300 x 300 DPI)

**Fig. 5** Double staining of liver sections with monoclonal antibodies directed against MPO or NE (red) and monoclonal antibody against ED1 (green) followed by fluorescence immunodetection in sections of rat liver at different time points after CCl\(_4\)-administration. a NE\(^+\) or ED1\(^+\) cells in CCl\(_4\)-induced liver injury at 0 h. b MPO\(^+\) or ED1\(^+\) cells in CCl\(_4\)-induced liver injury at 0 h. c NE\(^+\) or ED1\(^+\) cells in CCl\(_4\)-induced liver injury 24 h after administration. d MPO\(^+\) or ED1\(^+\) cells in CCl\(_4\)-induced liver injury 24 h after administration. Original magnification, \(\times\)100 173 x 130 mm (300 x 300 DPI)
showed an increased MPO gene expression 3.3-fold changes at 3 h and 2.5-fold change at 6 h after CCl₄-irradiation (Fig. 9a). NE gene expression increased 3.3-fold change at 3 h and 5.3-fold change at 6 h after CCl₄-irradiation (Fig. 9b). These results were confirmed by Western blot for MPO (Fig. 7b). CD11b-gene expression (Fig. 10), a marker for NG showed similar results.

Expression of MPO in acutely injured human liver

The results so far presented documented the expression of MPO in normal and acutely injured rat liver. To examine the expression of MPO in acutely injured human liver, indirect immunohistochemistry and immunofluorescence double staining studies were performed using histological sections of acute alcohol-associated injury. Indirect immunohistochemistry using antibodies against MPO (Fig. 11a) and NE (Fig. 11b) showed positive staining only within small, round cells with barely cytoplasm. What was striking was the presence of staining in the immediate vicinity of the cells, which probably represents extracellular secreted MPO and NE. Using immunofluorescence double staining with antibodies against MPO and CD-68, CD-68⁺ cells did not express MPO (Fig. 12a). This result was confirmed using antibodies against NE and CD-68. CD-68⁺ cells do not express NE (Fig. 12b). In Fig. 12 (sequential sections) there are MPO⁺ and CD-68⁺ cells (Fig. 12a) and NE⁺ and CD-68⁺ cells (Fig. 12b) can be seen.

Discussion

In this study, the expression of the MPO-gene in normal rat liver and isolated parenchymal and non-parenchymal rat liver cells, as well as in acutely induced rat liver either by CCl₄-administration or γ-irradiation was assessed. Human liver with acute alcohol induced injury was studied also. In normal rat liver immunohistochemically detectable MPO was detected only in single elastase positive NGs. MPO was not expressed in either parenchymal or non-parenchymal liver cells. Isolated mononuclear phagocytes did not express MPO, at either the level of RNA or at the level of protein. In contrast, Granulocytes were strongly positive for MPO expression to both RNA and protein.

During acute liver damage, induced by CCl₄-administration or single dose whole liver γ-Irradiation, an increase in the expression of MPO was observed that peaked 24 h after administration of CCl₄ and 3 h after γ-Irradiation. In acutely injured human liver, MPO expression was detected immunohistochemically only in granulocytes.

Fig. 6 Fold change of mRNA expression of MPO (a) and NE (b) after CCl₄-administration induced rat liver injury at different time points. Real time PCR was normalized by using two housekeeping genes: β-actin and 18 s RNA. Results represent mean ± SEM value of three experiments (in duplicate) compared with controls for each time point 173 × 62 mm (600 × 600 DPI)

Fig. 7 Western blot analysis of MPO of total protein in acute liver injury induced by CCl₄ (a) and γ-Irradiation (b). Protein was extracted from livers of control (Co) rats or 3, 6, 12, 24 and 48 h after one single oral dose of CCl₄-administration or γ-Irradiation induced rat liver injury. 20 μg of total protein were separated by SDS-PAGE, and blotted onto PVDF-membranes. The membranes were subsequently incubated with the antibodies against MPO (upper panel) and β-actin (lower panel). The molecular weight of the MPO is 59 and 13.5 kDa. In addition, autocatalytic products (AP) can be seen at 40 and 20 kDa (data not shown). The molecular weight of β-actin is 42 kDa 160 × 90 mm (300 × 300 DPI)
Immunofluorescence double staining of rat liver clearly showed that MPO was not detectable in ED1\(^+\) cells.

Brown et al. (2001) reported the expression of MPO in resident macrophages principally KCs, using immunohistochemical methods. In the present work, this finding could not be confirmed. In contrast, using an antibody against MPO only small round cells that are NG documented by NE staining were positive for MPO. ED1\(^+\) or CD68\(^+\) cells were negative for MPO. Analysis of the expression of MPO in parenchymal and non-parenchymal liver cells at the RNA level showed no expression of MPO.

The liver is known to be an important site for the clearance of NG (Peters et al. 1985; Farstad et al. 1991; Lovas et al. 1996). Various investigations have shown that endotoxemia selectively upregulates hepatic P-selectin and active phagocytosis of apoptotic circulating NG is limited...
to the liver. The complete absence of endothelial P-selectin results in neutrophilia (Frenette et al. 1996; Johnson et al. 1995; Mayadas et al. 1993). However, the phagocytotic capability of the KCs is diminished also (Shi et al. 1998).

In the damaged liver, adherent NG are metabolically active and transmigrate through the sinusoidal and microvascular endothelium following the release of extracellular matrix-degrading enzymes (Jaeschke and Farhood 1991; Jaeschke et al. 1991; Jaeschke and Smith 1997). The result is a pronounced inflammatory damage to the tissue. Activated NG generate reactive oxygen radicals and release proteases, which damage the endothelium and parenchymal cells directly (Jaeschke et al. 1990). NG emigrates quickly in inflamed tissue, engulf and destroy microorganisms and foreign antigens and ultimately undergo apoptotic death and phagocytoses by macrophage (Haslett 1992; Savill et al. 1989). Within the liver, two different populations of macrophages can be identified and functionally separated (Armbrust and Ramadori 1996) with newly immigrated mononuclear phagocytes manifesting MPO-positivity not present in terminal expended cells (Armbrust and Ramadori 1995). 24–48 h after administration of CCl₄, a strong immigration of inflammatory macrophages is observed in the pericentral area of the liver and these newly recruited monocytes are MPO-negative.

KCs represent approximately 10% of the liver cells and 80% of the tissue macrophage population (Armbrust et al. 1993; Hoedemakers et al. 1995) and play an important role in the defence mechanisms of the body. If KCs representing 10% of liver cells were to express MPO, either at the
RNA or protein level, they should have been immuno-stained positive for MPO. In this study no MPO expression was evident in KCs. The MPO protein within the liver is due to the presence of newly recruited NGs.

This increase of newly recruited MPO positive NGs at the site of liver injury underscores the important role of NGs in the inflammatory reaction that occurs during the multistep process of liver injury and repair.

Open Access This article is distributed under the terms of the Creative Commons Attribution Noncommercial License which permits any noncommercial use, distribution, and reproduction in any medium, provided the original author(s) and source are credited.

References

Akiyama Y, Miller PJ, Thurman GB, Neubauer RH, Oliver C, Favilla T, Beman JA, Oldham RK, Stevenson HC (1983) Characterization of human blood monocyte subset with low peroxidase activity. J Clin Invest 72:1093–1095

Armbrust T, Ramadori G (1996) Functional characterization of two different Kupffer cell populations of normal rat liver. J Hepatol 25:518–525

Armbrust T, Schwigler S, Zührens G, Ramadori G (1993) C1 esterase inhibitor gene expression in rat Kupffer cells, peritoneal macrophages and blood monocytes: modulation by interferon gamma. J Exp Med 178:373–380

Armbrust T, Ramadori G (1995) Mononuclear phagocytes of acutely injured rat liver abundantly synthesize and secrete fibronectin in contrast to Kupffer cells of normal liver. Biochem Biophys Res Commun 213:752–758

Arnhold J, Mueller S, Arnold K, Sonntag KJ (1993) Mechanisms of inhibition of chemiluminescence in the oxidation of luminol by sodium hypochlorite. Biol Chem 38:307–313

Bennett JM, Catoovsky D, Daniel MT, Flandrin G, Dalton DA, Granlindt HR, Sultan C (1976) Proposals for the classification of the acute leukaemias. French–American–British (FAB) cooperative group. Br J Haematol 33:451–458

Boyum A (1968) Isolation of mononuclear cells and granulocytes from human blood. Scand J Clin Lab Invest Suppl 97:77–89

Brennan ML, Anderson MM, Shih DM, Xu XD, Wang X, Mehta AC, Lim LL, Shi W, Hazen SL, Jacob JS, Crowley JR, Heinecke JW, Lusis AJ (2001) Increased atherosclerosis in myeloperoxidase-deficient mice. J Clin Invest 107:419–430

Brown KE, Brunt EM, Heinecke JW (2001) Immunohistochemical detection of myeloperoxidase and its oxidation products in Kupffer cells of human liver. Am J Pathol 159:2081–2088

Chirgwin JM, Przybyla AE, McDonald RJ, Rutter WJ (1979) Isolation of biologically active ribonucleic acid from sources enriched in ribonucleic acid. Biochemistry 18:5294–5299

Christiansen H, Batusic D, Saile B, Hermann RM, Dudas J, Rave-Hasert M, Hess CF, Schmidberger H, Ramadori G (2006) Identification of genes responsive to gamma radiation in rat liver and rat liver by cDNA array gene expression analysis. Radiat Res 165:318–325

Cribb AE, Miller M, Tesoro A, Spielberg SP (1990) Peroxidase-dependent oxidation of sulfonamides by monocytes and neutrophils from humans and dogs. Mol Pharmacol 38:744–751

Daugherty A, Dunn JL, Rateri DL, Heinecke JW (1994) Myeloperoxidase, a catalyst for lipoprotein oxidation, is expressed in human atherosclerotic lesions. J Clin Invest 94:437–444

Farstad BS, Sunderhagen E, Opdahl H, Benestad HB (1991) Pulmonary, hepatic and spleen sequestration of technetium-99m labelled autologous rabbit granulocytes: scintigraphic cell distributions after intravenous and intraarterial injections, exsanguination and intraarterial injection of cells passed through an intermediary host. Acta Physiol Scand 143:211–222

Frenette PS, Mayadas TN, Rayburn H, Hynes RO, Wagner DD (1996) Susceptibility to infection and altered hematopoiesis in mice deficient in both P- and E-selectins. Cell 84:563–574

Haqqani AS, Sandhu JK, Birnboim HC (1999) A myeloperoxidase-specific assay based upon bromide-dependent chemiluminescence of luminol. Anal Biochem 273:126–132

Haslett C (1992) Resolution of acute Inflammation and the role of apoptosis in the tissue fate of granulocytes. Clin Sci 83:639–648

Hoedemakers RMJ, Atsosorodjo-Briggs JE, Morselt HWM, Daemen T, Scherphof GL, Hardonk MJ (1995) Histochemical and electron microscopic characterization of hepatic macrophage subfractions isolated from normal and liposomal muramyl dipeptide treated rats. Liver 15:113–120

Imhof BA, Dunon D (1995) Leukocyte migration and adhesion. Adv Immunol 58:345–416

Jaeschke H, Farhood A (1991) Neutrophil and Kupffer cell-induced oxidant stress and ischemia-reperfusion injury in rat liver. Am J Physiol Gastrointest Liver Physiol 260:G355–G362

Jaeschke H, Smith CW (1997) Cell Adhesion and Migration. III. Leukocyte adhesion and transmigration in the liver vasculature. Am J Physiol Gastrointest Liver Physiol 273:G1169–G1173

Jaeschke H, Farhood A, Smith CW (1990) Neutrophils contribute to ischemia/reperfusion injury in rat liver in vivo. FASEB J 4: 3355–3359

Jaeschke H, Bautista AP, Spolarics Z, Spritzer JJ (1991) Superoxide generation by Kupffer cells and priming of neutrophils during reperfusion after hepatic ischemia. Free Radic Res 15:277–284

Johnson RC, Mayadas TN, Frenette PS, Mebius RE, Subramaniam M, Lacasce A, Hynes RO, Wagner DD (1995) Blood cell dynamics in P-selectin-deficient mice. Blood 86:1106–1114

Jolivalt C, Leininger-Muller B, Drozdz R, Naskalski JW, Siest G (1996) Apolipoprotein E is highly susceptible to oxidation by myeloperoxidase, an enzyme present in the brain. Neurosci Lett 210:61–64

Knittel T, Armbrust T, Schwegler S, Schuppan D, Ramadori G (1992) Distribution and cellular origin of undulin in rat liver. Lab Invest 67:779–787

Knittel T, Dinter C, Koboldt D, Neubauer K, Mehde M, Eichhorst S, Ramadori G (1999) Expression and regulation of cell adhesion molecules by hepatic stellate cells (HSC) of rat liver: involvement of HSC in recruitment of inflammatory cells during hepatic tissue repair. Am J Pathol 154:153–167

Knook DL, Sleyster EC (1976) Separation of Kupffer and endothelial cells of the rat liver by centrifugal elutriation. Exp Cell Res 99:444–449

Kumar AP, Piedrafita FJ, Reynolds WF (2004) Peroxisome proliferator-activated receptor gamma ligands regulate myeloperoxidase expression in macrophages by an estrogen-dependent mechanism involving the −463GA promoter polymorphism. J Biol Chem 279:8330–8335

Lovas K, Knudsen E, Iversen PO, Benestad HB (1996) Sequestration patterns of transfused rat neutrophilic granulocytes under normal and inflammatory conditions. Eur J Haematol 56:221–229

Malle E, Waeg G, Schreiber R, Grone EF, Sattler W, Grone HJ (2000) Immunohistochemical evidence for the myeloperoxidase/H2O2/halide system in human atherosclerotic lesions: colocalization of myeloperoxidase and hypochlorite-modified proteins. Eur J Biochem 267:4495–4503
Mayadas TN, Johnson RC, Rayburn H, Hynes RO, Wagner DD (1993) Leukozyte rolling and extravasation are severely compromised in P-selectin-deficient mice. Cell 74:541–554

McConnico RS, Weinstock D, Poston ME, Roberts MC (1999) Myeloperoxidase activity of the large intestine in an equine model of acute colitis. Am J Vet Res 60:807–813

Morita Y, Iwamoto H, Aibara S, Kobayashi T, Hasegawa E (1986) Crystallization and properties of myeloperoxidase from normal human leukocytes. J Biochem 99:761–770

Nagra RM, Becher B, Tourtellotte WW, Antel JP, Gold D, Paladin T, Smith RA, Nelson JR, Reynolds WF (1997) Immunohistochemical and genetic evidence of myeloperoxidase involvement in multiple sclerosis. J Neuroimmunol 78:97–107

Nahon P, Sutton A, Rufat P, Ziol M, Akouche H, Laguillier C, Chamaux N, Ganne-Carré N, Grando-Lemaire V et al (2009) Myeloperoxidase and superoxide dismutase 2 polymorphisms comodulate the risk of hepatocellular carcinoma and death in alcoholic cirrhosis. Hepatology 50:1484–1493

Nakagawara A, Nathan CF, Cohn ZA (1981) Hydrogen peroxide metabolism in human monocytes during differentiation in vitro. J Clin Invest 68:1243–1252

Nauseef WM (1988) Myeloperoxidase deficiency. Hematol Oncol Clin North Am 2:135–158

Nauseef WM, Olsson I, Arnljots K (1988) Biosynthesis and processing of myeloperoxidase a marker for myeloid cell differentiation. Eur J Haematol 40:97–110

Neubauer K, Knittel T, Armbrust T, Ramadori G (1995) Accumulation and cellular localization of fibrinogen/fibrin during short-term and long-term rat liver injury. Gastroenterology 108:1124–1135

Olsson I, Olofsson T, Odeberg H (1972) Myeloperoxidase-mediated iodination in granulocytes. Scand J Haematol 9:483–491

Owen CA, Campbell MA, Boukedes SS, Stockley RA, Campbell EJ (1994) A discrete subpopulation of human monocytes expresses a neutrophil-like proinflammatory (P) phenotype. Am J Physiol Lung Cell Mol Physiol 267:L775–L785

Panasenko OM, Arnhold J (1999) Linoleic acid hydroperoxide favours hypochlorite- and myeloperoxidase-induced lipid peroxidation. Free Radic Res 30:479–487

Panasenko OM, Arnhold J, Vladimirov YuA, Arnhold K, Sergienko VI (1997) Hypochlorite-induced peroxidation of egg yolk phosphatidylcholine is mediated by hydroperoxides. Free Radic Res 27:1–12

Peters AM, Saverymuttu SH, Bell RN, Lavender JP (1985) Quantification of the distribution of the marginating granulocyte pool in man. Scand J Haematol 34:111–120

Podrez EA, Schmitt D, Hoff HF, Hazen SL (1999) Myeloperoxidase-generated reactive nitrogen species convert LDL into an atherogenic form in vitro. J Clin Invest 103:1547–1560

Rensen SS, Slaats Y, Nijhuis J, Jans A, Bieghs V, Driessen A, Malle E, Greve JW, Buurman WA (2009) Increased hepatic myeloperoxidase activity in obese subjects with nonalcoholic steatohepatitis. Am J Pathol 175:1473–1482

Reynolds WF, Rhees J, Maciejewski D, Paladino T, Sieburg H, Maki RA, Masliah E (1999) Myeloperoxidase polymorphism is associated with gender specific risk for Alzheimer’s disease. Exp Neurol 155:31–41

Savill JS, Wyllie AH, Henson JE, Walport MJ, Henson PM, Haslett C (1989) Macrophage phagocytosis of aging neutrophils in inflammation. J Clin Invest 83:865–875

Shi J, Kokubo Y, Wake K (1998) Expression of P-selectin on hepatic endothelia and platelets promoting neutrophil removal by liver macrophages. Blood 92:520–528

Taylor KL, Pohl J, Kinkade JM (1992) Unique autocatalytic cleavage of human myeloperoxidase. Implications for the involvement of active site MET409. J Biol Chem 267:25282–25288

Weiss SJ (1989) Tissue destruction by neutrophils. N Engl J Med 320:365–376

Winterbourn CC (1985) Comparative reactivities of various biological compounds with myeloperoxidase-hydrogen peroxide-chloride, and similarity of the oxidant to hypochlorite. Biochim Biophys Acta 840:204–210

Yamada M, Mori M, Sugimura T (1981) Purification and characterization of small molecular weight myeloperoxidase from human promyelocytic leukemia HL-60 cells. Biochemistry 20:766–771

Zgliczynski JM, Stelmaszyńska T (1975) Chlorinating ability of human phagocytosing leucocytes. Eur J Biochem 56:157–162