We investigated the effects of the antibiotic ceftazidime (CAZ) on the cytolytic action of the neutrophil myeloperoxidase-hydrogen peroxide-chloride anion system (MPO/H2O2/Cl-). In this system, myeloperoxidase catalyses the conversion of H2O2 and Cl- to the cytotoxic agent HOCl. Stimulated neutrophils can release MPO into the extracellular environment and then may cause tissue injury through direct endothelial cells lysis. We showed that human umbilical vein endothelial cells (HUVEC) were capable of taking up active MPO. In presence of H2O2 (10^-4 M), this uptake was accompanied by cell lysis. The cytolyis was estimated by the release of 51Cr from HUVEC and expressed as an index of cytotoxicity (IC). Dose dependent protection was obtained for CAZ concentrations ranging from 10^-5 to 10^-3 M; this can be attributed to inactivation of HOCl by the drug. This protection is comparable to that obtained with methionine and histidine, both of which are known to neutralize HOCl. This protection by CAZ could also be attributed to inactivation of H2O2, but when cytolsis was achieved with H2O2 or O2- generating enzymatic systems, no protection by CAZ was observed. Moreover, the peroxidation activity of MPO (action on H2O2) was not affected by CAZ, while CAZ prevented the chlorination activity of MPO (chlorination of monochlorodimedon). So, we concluded that CAZ acts via HOCl inactivation. These antioxidant properties of CAZ may be clinically useful in pathological situations where excessive activation of neutrophils occurs, such as in sepsis.

**Key words:** Antibiotics, ceftazidime, endothelial cells, hypochlorous acid, myeloperoxidase, oxidant stress

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**Cytotoxicity towards human endothelial cells, induced by neutrophil myeloperoxidase: protection by ceftazidime**

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**Introduction**

Sepsis, septic shock, severe trauma, hypovolaemic shock and acute pancreatitis are examples of severe pathological situations accompanied by an intense inflammatory reaction involving activation of specific cells and release of mediators. This inflammatory reaction occasionally overwhems the organism’s defences, with excessive activation of polymorphonuclear leucocytes (PMN). This can lead to the systemic inflammatory response syndrome (SIRS) and eventually to multiple organ dysfunction syndrome (MODS).1-3 PMN activation produces activated oxygen species (mainly superoxide anion and hydrogen peroxide, H2O2), inflammatory mediators, and proteolytic and hydrolytic enzymes.4,5 PMN also release myeloperoxidase (MPO) from their granules; in the presence of H2O2 and chloride anion (Cl-), MPO generates hypochlorous acid (HOCl). HOCl reacts with amines to form chloramines, but also with H2O2 itself to produce singlet oxygen, an activated form of oxygen.5,7 These activated oxygen species and oxidant products of MPO activity normally play a beneficial role in host defence against invading microorganisms. However, their excessive production can be detrimental for tissues, especially for endothelial cells, such as, when PMN are trapped in capillaries and become strongly adherent to endothelium.8 They are also destructive for plasma proteins, especially those with thiol (-SH) functions, such as α2 macroglobulin, an essential plasmatic antiproteinase.8,9 In these situations associated with excessive activation of PMN, therapeutic agents capable of neutralizing these active oxygen species and protecting endothelial cells from oxidant stress would have potential clinical utility. Recent research has demonstrated that antibiotic molecules of the aminoglycoside and

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cephalosporin families can protect epithelial cells submitted to an oxidant stress, and are scavengers for HOCl molecules.12–14

We have demonstrated that the antibiotic ceftriaxone (CAZ), which belongs to the cephalosporin family, exerts an antioxidant effect in vitro. It inhibits percoll mediates lipoperoxidation by iron scavenging neutralizes HOCl, and quenches singlet oxygen.15–17 The purpose of this study was to demonstrate that CAZ is able to protect endothelial cells from the oxidant stress induced by the activity of MPO or by HOCl. In our study, CAZ was used as a model of broad spectrum β-lactamase resistant antibiotics, which are widely used in clinical situations of severe sepsis, as mentioned above; these are precisely the situations wherein intense PMN activation is seen.

Materials and Methods

Cell culture: Human umbilical vein endothelial cells (HUVEC) were isolated by collagenase treatment of umbilical cords according to Jaffe et al.8 HUVEC were cultured on 0.2% gelatine coated dishes in M199 medium supplemented with 10% heat-inactivated foetal calf serum (Gibco), 5% heat-inactivated human serum (Sigma), penicillin (100 U/ml), streptomycin (100 μg/ml), heparin (90 μg/ml) and endothelial cell growth factor (Boehringer–Mannheim) (20 μg/ml). Adherent HUVEC in six multiwell plates (passage 2) were used in all experiments.

Assay for MPO enzyme activity: MPO was purified from human polymorphonuclear neutrophils as described previously.19,20 Its enzyme activity was determined by spectrophotometric methods.

MPO peroxidative activity assay: MPO peroxidative activity was assayed by measuring the absorbance increase at 460 nm caused by the oxidation of ortho-dianisidine.21 MPO (dissolved in 50 μl) was added to 3 ml of 50 mM ortho-dianisidine in Sörensen buffer pH 5.5 and the reaction was started by the addition of H2O2 at a final concentration of 0.15 mM. The absorbance increase at 460 nm was followed for 1 min. One unit of activity was defined as the amount of MPO which produced an absorbance increase of 1 optical density unit per minute.

MPO-dependent chlorination activity: MPO-dependent chlorination activity was measured by following the conversion of monochlorodimedon to dichlorodimedon at 290 nm.22 Typical experimental conditions were as follows: MPO (5 μg) was incubated in 3 ml of monochlorodimedon (24 μM) in 100 mM phosphate buffer (pH 5.5) with NaCl (50 mM) added. The reaction was started by the addition of H2O2 (0.1 mM) and the decrease of absorbance at 290 nm was followed for 1 min. One unit of activity was defined as the amount of MPO which produced an absorbance decrease of 1 optical density unit per minute.

To determine the effects of CAZ on MPO enzyme activities, MPO was incubated with different concentrations of CAZ for 5 min at room temperature before the enzyme assays. The data represent arithmetic means ± S.D. of triplicates.

MPO uptake by HUVEC: Confluent cells in 1 ml of Hank’s Balanced Salt Solution (HBSS) were incubated at 37°C with increasing amounts of MPO (from 0 to 30 μg) for 3 h or with a fixed amount of MPO (5 μg) for periods of time ranging from 0 to 360 min. Each assay was done in triplicate and the entire experiment was repeated three times. After these incubations, cells were washed three times with HBSS, and the MPO content of the cells was estimated by measurement of MPO peroxidative activity (see above). Ortho-dianisidine (1 ml) was added to adherent cells and the reaction was started by the addition of H2O2 at a final concentration of 0.15 mM. After an incubation of 30 min, the reaction buffer was collected and the absorbance at 460 nm was read. The amount of MPO present in the cells was calculated from a standard curve for which known amounts of MPO were incubated under the same conditions.

Cytotoxicity assay: Cytotoxicity was assessed by measuring the release of previously incorporated 51Cr.23 Confluent HUVEC in six multiwell plates were labelled overnight by the addition of 20 μCi 51Cr in culture medium per well (sodium chromate, Amersham). HUVEC were washed in HBSS to remove unincorporated 51Cr and the cells were then submitted to oxidant stress as described below. Each assay was done in triplicate. At the end of the oxidant stress period, the supernatants were collected and the cells were washed three times with HBSS. Supernatant and washings were pooled and 51Cr release [R] was quantified by γ counting.

Cells were lysed in NaOH (1 N) and the intracellular 51Cr [I] was counted. The percentage of 51Cr release [PR] was calculated for each test condition as follows:

\[
PR = \frac{R}{R + I} \times 100
\]

Where \( R + I \) is the total amount of 51Cr present in the cells before the application of the oxidant.
Ceftriaxone protects against oxidant stress. An index of cytotoxicity \( [IC] \) was calculated according to the following formula:

\[
IC = 100 \times \frac{[PR]_{\text{test}} - [PR]_{\text{cont}}}{100 - [PR]_{\text{cont}}}
\]

where \([PR]_{\text{cont}}\) represents the spontaneous release of \(^{51}\text{Cr} \) (in percent as above) by the control cells incubated in HBSS alone.

**Oxidant stress against HUVECs.** Four oxidant stressors were used against HUVECs. For each oxidant stress assay, controls were performed in which HUVECs were incubated under the same conditions in HBSS alone; each assay was done in triplicate.

**Cytotoxicity induced by the MPO/H\(_2\)O\(_2\)/Cl\(^-\) system.** HUVECs in 1 ml of HBSS were incubated in the presence of MPO (5 \( \mu \)g) at 37°C for 2 h. The Cl\(^-\) required for MPO enzyme activity was supplied by HBSS. H\(_2\)O\(_2\) (10\(^{-4}\)M) was added to initiate MPO enzyme activity and after a further 2 h incubation at 37°C, IC was determined as described before.

**Cytotoxicity induced by sodium hypochlorite (NaOCl).** NaOCl (10\(^{-3}\)M) was added to HUVECs in 1 ml of HBSS. After a 2 h incubation, IC was evaluated.

**Cytotoxicity induced by the glucose/glucose oxidase system.** HUVECs in 1 ml of HBSS (containing 5 mM of glucose) were incubated overnight with glucose oxidase (0.2 U) (Boehringer-Manheim) prior to IC evaluation.

**Cytotoxicity induced by the xanthine/xanthine oxidase system.** HUVECs in 1 ml of HBSS were treated with 2 mM xanthine/20 mU of xanthine oxidase (Boehringer-Manheim) for 2 h following which IC was determined.

### Protection from oxidant stress

The effects of CAZ on oxidant stress were compared to methionine, histidine and dabco (1,4-diazabicyclo[2.2.2]octane), which are well known HOCl and H\(_2\)O\(_2\) scavengers.\(^{24,25}\) \(^{51}\text{Cr} \) labelled HUVECs in 1 ml of HBSS supplemented with one of the putative protectors (CAZ, methionine, histidine or dabco) were incubated for 2 h prior to initiating the cytotoxic treatments described above. For the oxidant stress with MPO/H\(_2\)O\(_2\)/Cl\(^-\), the protector was added with MPO 2 h before the addition of H\(_2\)O\(_2\). After the stress, IC was determined, and the % protection was obtained according to the following formula:

\[
\text{% protection} = 100 \times \left(1 - \frac{I_{\text{eff}}}{I_{\text{stress}}} \right)
\]

where \(I_{\text{eff}}\) and \(I_{\text{stress}}\) were, respectively, the IC obtained in the presence or in the absence of the study drug.

### Results

**MPO uptake by HUVECs.** The results for the kinetics of MPO uptake are presented in Fig. 1. HUVECs were pre-incubated with MPO (5\( \mu \)g) and H\(_2\)O\(_2\) (10\(^{-4}\)M) added simultaneously, only a low cytotoxicity \((IC = 5.04 \pm 2.1)\) was found. No cytotoxicity was obtained with either MPO or H\(_2\)O\(_2\) alone. To permit the uptake of MPO prior to initiating the oxidant stress with H\(_2\)O\(_2\), the cells were pre-incubated with MPO for 2 h. This time of pre-
incubation was chosen because after 2h MPO uptake was near its maximal value, but still on the linear part of the curve of Fig. 1. After addition of H$_2$O$_2$ (10$^{-4}$ M), a further incubation of 2h with the complete cytotoxic system (MPO/ H$_2$O$_2$/Cl$^-$) was performed. After this time of incubation, convenient and reproducible IC were obtained. For shorter times of incubation, we measured low IC values, insufficient to allow reproducible assays of putative protective substances.

Figure 3 plots the dose--response effect of MPO on HUVEC cytotoxicity: a plateau was reached at 5μg MPO (with a maximal IC = 39.92 ± 0.64).

Protection of HUVEC from MPO/H$_2$O$_2$/Cl$^-$ or NaOCl oxidant stresses: To test the effects of putative protectors (CAZ, methionine, histidine, or dabco), the protector was added at different concentrations on HUVEC, together with 5μg of MPO, and the cells were incubated for 2h prior to addition of H$_2$O$_2$ (Fig. 4). We controlled the effects of CAZ on MPO uptake by endothelial cells and found neither inhibiting nor enhancing effects of the antibiotic: without CAZ, we found an uptake of 0.476 ± 0.044μg MPO, and with CAZ (10$^{-3}$ M), an uptake of 0.469 ± 0.011μg (no statistical difference). After the stress, IC was calculated and the results were expressed as percent protection according to the formula defined in Materials and Methods: a protection of 100% corresponds to a total inhibition of stress-induced cytotoxicity.

The protection by CAZ was dose-dependent and particularly effective at the concentration of 10$^{-3}$ M (percent protection = 92.4 ± 1.7). This protection was confirmed by light microscope observations, which indicated a similar cellular aspect for control cells (without stress) and cells...
Stressed in the presence of CAZ $10^{-3}$M, while the cells stressed in the absence of CAZ appeared strongly damaged (loss of most of the cells and strong morphologic changes). At a CAZ concentration of $10^{-3}$M, a protective effect was still observed (percent protection = 26.0 ± 0.9). Methionine and CAZ demonstrated comparable protective effects (no statistically significant difference between the two compounds), but histidine was less effective ($p < 0.05$ for each concentration compared to CAZ and methionine). In contrast, no protection was obtained with dabco (percent protection = 6.6 ± 9.0).

From these results, we concluded that CAZ was capable of neutralizing cytotoxicity induced by the MPO/H$_2$O$_2$/Cl$^-$ system by inactivating the HOCI generated by MPO. To confirm the direct effect of CAZ on NaOCl, $^{51}$Cr labelled HUVEC were exposed to NaOCl stress in presence of CAZ, methionine, or histidine; the cells were incubated at 37°C for 2h in the presence of putative protector (at a concentration of $10^{-3}$M). NaOCl ($10^{-3}$M) was then added and the IC was determined after a further 2h incubation. The IC in the absence of protector was 14.86 ± 3.15 and the percent protection, calculated as previously described, are presented in Table 1. Marked protection was seen with CAZ and methionine, while histidine was weakly active.

**Absence of a protective effect of CAZ on oxidant stress generating H$_2$O$_2$:** To test the effect of CAZ on HUVEC cytotoxicity induced by H$_2$O$_2$ or O$_2^-$, we incubated the cells in the presence of enzymatic systems generating this reactive oxygen species: glucose/glucose oxidase or xanthine/xanthine oxidase systems.

For the two stresses, no statistically significant difference was observed (two-tailed Student’s test) between the cytotoxicity obtained in the absence or in the presence of CAZ ($10^{-3}$M). The IC values obtained for glucose/glucose oxidase stress were respectively 25.02 ± 3.91 without CAZ and 25.93 ± 5.26 with CAZ. For xanthine/xanthine oxidase stress the IC values were 76.98 ± 0.49 without CAZ and 78.37 ± 2.31 with CAZ.

**Effects of CAZ on MPO enzyme activities:** Even at concentrations of CAZ ($10^{-3}$M) which were highly protective for HUVEC submitted to MPO/H$_2$O$_2$/Cl$^-$ stress, only weak inhibition of MPO peroxidative activity was observed (90.0 ± 0.7% of control activity) (Fig. 5). In contrast, MPO-dependent chlorination of monochlorimedon was very sensitive to CAZ: no chlorination was observed in the presence of $5 \times 10^{-4}$M and only $61.0 \pm 4.9\%$ of activity was conserved in the presence of $2.5 \times 10^{-3}$M CAZ.

**Discussion**

In this study we demonstrate that endothelial cells are capable of binding and taking up myeloperoxidase. This uptake shows steep time dependency at 37°C, and is maximal at 3h incubation. The yield of the uptake process is low, with a maximum incorporation of 11% when 5µg of MPO are added to the culture medium (Fig. 1). Increasing the amount of MPO added to the medium does not increase uptake, which peaks at 7.5µg (62.4nM) added MPO. At this level, 7.8% of the added dose is incorporated after 3h incubation. These results agree with those of Zabucchi et al. who show that 75% of endothelial cells have a positive cytochemical peroxidase reaction after preincubation with high concentrations of MPO (up to 420nM). These workers also showed that the enzyme was

Table 1. CAZ protection of HUVEC from NaOCl stress. Comparison with methionine and histidine

<table>
<thead>
<tr>
<th>Protector</th>
<th>% protection</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAZ ($10^{-3}$M)</td>
<td>77.90 ± 11.50</td>
</tr>
<tr>
<td>Methionine ($10^{-3}$M)</td>
<td>91.38 ± 5.20</td>
</tr>
<tr>
<td>Histidine ($10^{-3}$M)</td>
<td>20.18 ± 12.00</td>
</tr>
</tbody>
</table>

The data represent mean ± S.D. of three experiments performed in triplicate.

FIG. 5. Effects of CAZ on MPO enzyme activities. MPO was preincubated with different concentrations of CAZ for 5 min at room temperature. , peroxidative activity of MPO; , chlorination activity of MPO. MPO enzyme activities obtained in the absence of CAZ were taken as the reference (100%). Each point represents the mean ± S.D. of two experiments performed in triplicate. *No statistical difference ($p > 0.05$) and **statistical difference ($p < 0.05$) vs. control when analysed with the two-tailed Student’s t-test.
present on the cell membrane and in the cytoplasm. Using the same cytochemical reaction, we showed uptake of MPO by endothelial cells, and its presence in intracellular granules (results not shown). Spectrophotometric measurement of intracellular enzymatic activity confirmed uptake and showed that the internalized enzyme retained its activity. We used lower concentrations of MPO than Zabucchi et al., but in our study, we used a 3 h incubation (as opposed to the 10 min used by Zabucchi’s group). This duration of incubation did not alter the enzyme, because at the end of the period, the sum of enzyme left in the supernatant and that incorporated into the cells exactly equalled the total initially added.

In the presence of H$_2$O$_2$, MPO produces a cytolytic oxidative stress on endothelial cells, which we have expressed as a cytotoxicity index. This IC increases linearly with the dose of MPO with a plateau at 5 btg per 4 x 10$^5$ cells. We chose lower concentrations of MPO than Zabucchi et al., but in our study, we used a 3 h incubation (as opposed to the 10 min used by Zabucchi’s group). This duration of incubation did not alter the enzyme, because at the end of the period, the sum of enzyme left in the supernatant and that incorporated into the cells exactly equalled the total initially added.

We thus see that CAZ, in addition to its antibiotic activity possesses antioxidant properties.
These could be useful in situations where excessive PMN activation, production of activated species of oxygen, and liberation of MPO are seen. Sepsis and septic shock are two such situations. In our model, the lowest concentration associated with a protective effect ($10^{-5} \text{M} = 5.46 \mu g/ml$, 26% protection) is inferior to the serum concentrations attained in patients treated with this antibiotic: following an intravenous infusion of 2g CAZ in healthy volunteers and patients, the serum concentration peaks at 59 to 83 $\mu g/ml$. The combination of an antioxidant activity with those of an antibiotic in one molecule allows simultaneous treatment with a well characterized, widely used drug, of both the infectious aspect of the disease process, as well as the consequences of excessive PMN activation.

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