Uric acid disrupts hypochlorous acid production and the bactericidal activity of HL-60 cells

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1. Introduction

Neutrophils are the first cells to be recruited in the defense against microbial invasion in mammals. Microbe recognition by specific receptors at neutrophil surface induces phagocytosis, release of cytoplasmic granules into the phagosome and generation of oxidants by a process called neutrophil oxidative burst. Granule proteins discharged into the phagosome digest microorganisms and act synergistically with oxidants to kill them [1–4]. The neutrophil oxidative burst starts by the reduction of oxygen to the anion radical superoxide (NOX) [5]. Superoxide dismutates to hydrogen peroxide in the acidic intraphagosomal environment [6] and this is the main source of hydrogen peroxide, that in turn oxidizes chloride to hypochlorous acid (HOCl), the strongest microbicide in neutrophils [2,7,8]. Oxidation of hydrogen peroxide, that in turn oxidizes chloride to hypochlorous acid

Abstract

Uric acid is the end product of purine metabolism in humans and is an alternative physiological substrate for myeloperoxidase. Oxidation of uric acid by this enzyme generates uric acid free radical and urate hydroperoxide, a strong oxidant and potentially bactericidal agent. In this study, we investigated whether the oxidation of uric acid and production of urate hydroperoxide would affect the killing activity of HL-60 cells differentiated into neutrophil-like cells (dHL-60) against a highly virulent strain (PA14) of the opportunistic pathogen Pseudomonas aeruginosa. While bacterial cell counts decrease due to dHL-60 killing, incubation with uric acid inhibits this activity, also decreasing the release of the inflammatory cytokines interleukin-1β (IL-1β) and tumor necrosis factor-α (TNF-α). In a myeloperoxidase/Cl-/H2O2 cell-free system, uric acid inhibited the production of HOCl and bacterial killing. Fluorescence microscopy showed that uric acid also decreased the levels of HOCl produced by dHL-60 cells, while significantly increased superoxide production. Uric acid did not alter the overall oxidative status of dHL-60 cells as measured by the ratio of reduced (GSH) and oxidized (GSSG) glutathione. Our data show that uric acid impairs the killing activity of dHL-60 cells likely by competing with chloride by myeloperoxidase catalysis, decreasing HOCl production. Despite diminishing HOCl, uric acid probably stimulates the formation of other oxidants, maintaining the overall oxidative status of the cells. Altogether, our results demonstrated that HOCl is, indeed, the main relevant oxidant against bacteria and deviation of myeloperoxidase activity to produce other oxidants hampers dHL-60 killing activity.

Keywords: Uric acid Myeloperoxidase Hypochlorous acid dHL-60 Pseudomonas aeruginosa Microbicidal
with both myeloperoxidase-Compound I and Compound II make it a relevant substrate for myeloperoxidase [15]. Uric acid has been extensively studied by its antioxidant properties (one-electron reduction potential = 0.59 V, pH 7.0) [16], being able to react with hydroxyl radical, singlet oxygen, hypochlorous acid, to chelate transition metal ions and to repair free radicals in proteins [17-20]. Despite its antioxidant ability, uric acid and its monoanion urate increase oxidative damage and inactivate enzymes sensitive to oxidative stress [21-24]. Products of uric acid oxidation react with glutathione, methionine and thiol-peroxidases [25,26]. Therefore, the predominant effect of uric acid as pro or antioxidant is still a matter of debate [27].

The oxidation of uric acid by myeloperoxidase and other peroxides generates the unstable urate free radical, which reacts at diffusion-controlled rate with superoxide [28] to form urate hydroperoxide [15,25,29]. This reaction is of particular relevance into the phagosome where there are plentiful amounts of superoxide, hydrogen peroxide and myeloperoxidase. Urate hydroperoxide is a strong oxidant [25,26] and potentially bactericidal. Therefore, the oxidation of uric acid and production of urate hydroperoxide in neutrophils could contribute to the bactericidal effect of these cells.

In addition to the pro-oxidant effect, uric acid has been widely described by its pro-inflammatory properties. It can precipitate as monosodium urate crystals and activate the NLRP3 inflammasome to release IL-1β [30,31]. Soluble uric acid released from dying cells initiates inflammatory response by recruiting neutrophils [32,33]. Uric acid activates NADPH oxidase to produce superoxide [34], activates MAP kinases and the nuclear factor kappa-B (NF-kB) increasing the expression of monocyte chemoattractant protein-1 (MCP-1) [35,36], primes mononuclear immune cells enhancing the pro-inflammatory effect of toll-like receptors ligands [37] and induces formation of neutrophils extracellular traps (NETs) [38]. All this pro-inflammatory capacity could also stimulate immune cells and increase their bactericidal effect.

Therefore, we designed this study to address the question of whether uric acid would indeed improve the killing activity of neutrophil-like cells. We differentiated HL-60 cells into neutrophils (dHL-60) and incubated them with the opportunistic Gram-negative pathogen Pseudomonas aeruginosa, a major cause of hospital acquired infections, microbial keratitis and chronic lung infections in cystic fibrosis [39-41]. Surprisingly, we found that uric acid decreased the killing activity of dHL-60 cells. This effect was likely related to the disruption in HOCl production by uric acid. Therefore, the decrease of HOCl levels is severely detrimental to the killing potential of dHL-60 cells and the production of other oxidants like urate free radical and urate hydroperoxide is not sufficient to overcome HOCl deprivation.

2. Material and methods

2.1. Materials

Human leukocyte MPO (EC 1.7.1.11) was purchased from Planta Natural Products (Vienna, Austria). Fetal bovine serum was from VitroCell (Campinas, Brazil). R19-S probe was from FutureChem (Seoul, Korea). Xanthine oxidase was purchased from Calbiochem – Merck Millipore (Darmstadt, Germany). dihydroorhadamine 123 (DHR) was purchased from Life Technologies (Carlsbad, CA, USA). All other reagents including cell culture materials RPMI 1648, penicillin and streptomycin, uric acid, 5,5′-dithiobis-(2-nitrobenzoic acid) (DTNB), taurine, catalase, cytochrome c, propidium iodide (PI), 4′,6-diamidino-2-phenylindole dye (DAPI), phorbol myristate acetate (PMA), staurosporin and acetaldehyde were purchased from Sigma-Aldrich (Darmstadt, Germany).

2.2. Human cell culture and differentiation

Human leukemic cell line (HL-60) (BCRJ, RJ, Brazil) was maintained in RPMI 1648 growth medium supplemented with 20% fetal bovine serum (FBS), streptomycin (100 µg/mL) and penicillin (100 U/mL) at 5% CO2 and 37 °C. HL-60 cells were differentiated into neutrophils (dHL-60) by the presence of 1.3% dimethyl sulfoxide (DMSO) in growth medium supplemented with 10% FBS. After five days, dHL-60 were centrifuged at 350g for 10 min and washed twice with sterile saline (0.9% NaCl). Cell pellet was then resuspended into phosphate buffer saline supplemented with glucose (PBS glucose: 10 mM NaH2PO4; 2 mM KH2PO4; 137 mM NaCl, 1 mM CaCl2, 0.5 mM MgCl2 and 1 g/L glucose) or in growth medium without antibiotics. The differentiation was confirmed by superoxide production using the cytochrome c assay, dHL-60 (1 × 106 cells) were incubated with taurine (5 mM) and cytochrome c (40 µM) in 300 µL PBS/glucose. Cells were activated with PMA (100 ng/mL) and samples were red at 550 nm in a Synergy H1 Hybrid Reader (Biotek, USA). A flux of 2–4 µM/min was expected in differentiated cells.

2.3. Bacterial culture

Pseudomonas aeruginosa UCBPP-PA14 (PA14) [42] was grown in Luria-Bertani medium (LB) overnight under shaking at 200 rpm and 37 °C. The day after, cells were diluted to OD600 nm = 0.1 and incubated again until OD600 nm = 0.4 or OD600 nm = 2 for initial or late exponential phase of growth, respectively.

2.4. Bacterial killing assay

dHL-60 cells (2 × 105) were challenged with PA14, OD600 nm = 2, multiplicity of infection (MOI) 1:10 in absence or presence of 0.2, 0.5 or 2 mM uric acid and incubated at 37 °C for 1, 2 or 3 h in antibiotic-free RPMI medium. After incubation, samples were collected and diluted in series so they contained 105–108 bacteria/mL. Ten microliters of these dilutions were spread on LB agar, incubated for 14–16 h at 37 °C and the colony-forming units (CFU) were counted.

To test the bacterial killing by purified myeloperoxidase, we first incubated 100 nM myeloperoxidase, 150 mM NaCl in minimum medium (70 mM phosphate buffer, 18 mM (NH4)2SO4, 100 µM CaCl2, 2 mM MgSO4 and 2% glucose, pH 7.1) in the absence or presence of 0.5 mM uric acid. The production of HOCl was initiated by addition of 100 µM H2O2, and allowed to react for 30 min at room temperature. The reaction was stopped by adding 50 µM/mL catalase for 5 min. This ensured that all remaining H2O2 was removed to avoid any unspecified effect upon bacterial killing. Alternatively to the above, 0.5 mM uric acid was added only 30 min after myeloperoxidase/Cl-/H2O2 reacting. Uric acid was allowed to react for 0, 10 or 30 min for evaluating the scavenger effect of uric acid upon HOCl rather than the competition by myeloperoxidase and inhibition of HOCl formation. After this pre-incubation, the PA14 (1 × 106 cells/mL) was added to the system and incubated for 2 h at 37 °C with constant shaking (200 rpm). Serial dilutions were immediately spread on agar plates, and the colony-forming units were determined after overnight incubation at 37 °C.

2.5. Quantification of hypochlorous acid

Measurement of hypochlorous acid production by purified myeloperoxidase and by dHL-60 was performed by 2-nitro-5-thiobenzozate (TNB) [43]. dHL-60 cells (5 × 106) were suspended in PBS glucose with or without uric acid. Cells were infected with PA14 (MOI 1:10) and incubated at 37 °C for 1 h. The myeloperoxidase/Cl-/H2O2 system was similar to the used in the bacterial killing assay: 100 nM myeloperoxidase, 150 mM NaCl, 5 mM taurine in minimum medium (70 mM phosphate buffer, 18 mM (NH4)2SO4, 100 µM CaCl2, 2 mM MgSO4 and 2% glucose, pH 7.1) in the absence or presence of 0.1, 0.2 or 0.5 mM uric acid. Reaction was initiated by adding 100 µM H2O2 and incubated for 30 min at room temperature. Reaction was stopped by incubating with 50 µg/mL catalase. The measurement of HOCl was also performed when
0.5 mM uric acid was added 30 min after myeloperoxidase/Cl₂/H₂O₂ reacting (see above) to evaluate the scavenger effect of uric acid upon HOCl rather than the inhibition of HOCl formation. A five-fold dilution of dHL-60 or myeloperoxidase/Cl₂/H₂O₂ reaction system was incubated with 80 µM TNB by 15 min in the dark. The oxidation of TNB to the colorless DTNB (5,5'-dithiobis-2-nitrobenzoic acid) by taurine chloramine was measured at 412 nm and quantified using TNB molar absorption coefficient (ε₄₁₂ = 14,200 M⁻¹ cm⁻¹). Since two TNB are consumed to form one DTNB per taurine chloramine, the molar absorption coefficient was multiplied by two, i.e. 28,400 M⁻¹ cm⁻¹ [43].

The production of HOCl by dHL-60 was also measured using a specific fluorescent probe that permeates the cell membrane [44,45]. dHL-60 (5 × 10⁶) were suspended in PBS glucose containing 10 µM DHE (10 µM) in the absence or presence of 0.5 mM uric acid or 50 µM 4-aminoxybenzoic acid hydrazide (ABAH). Cells were infected with PA14 (MOI 1:10) and incubated at 37 °C for 1 h. Alternatively, cells were activated with 100 ng/mL LPS (positive control for HOCl production). The samples were centrifuged at 180g for 10 min and the pellets were washed in PBS. Two hundred microliters 1.3% paraformaldehyde were added to the pellet and incubated at room temperature for 15 min. Cells were washed once again in PBS and stained with 4,6-diamidino-2-phenylindole (DAPI). After one final wash, cells were suspended in mounting media Fluor Mount, spread over a microscope slide and covered with a coverslip. Images were acquired with a LSM 780-NL9 confocal Carl Zeiss microscope (Göttingen, Germany) with all parameters fixed for imaging all conditions. Fluorescence intensity (λex = 515 nm, λem = 545 nm) of 10 different fields was quantified using ImageJ software.

2.6. HL-60 cell viability

Differentiated HL-60 cells (1 × 10⁶) were incubated in the absence or presence of uric acid (0.2; 0.5 and 2 mM) and infected with PA14 (MOI 1:10) for 1 or 2 h at 37 °C. After incubation, samples were centrifuged at 150g for 10 min and the pellets were incubated with 10 µg/ml propidium iodide (PI) in PBS for 15 min. Following incubation, the percentage of viable and dead cells was determined using a BD Biosciences flow cytometry (San Jose, CA, USA). Fluorescence of labeled cells was detected with λex = 535 nm, λem = 620 nm. Staurosporine (5 µM) was used as positive control [46]. To evaluate the activity of lactate dehydrogenase (LDH) in dHL-60 supernatant, the cells (1 × 10⁶) were infected with PA14 (MOI 1:10) in PBS supplemented with glucose in presence of uric acid (0.1, 0.2 or 0.5 mM) and incubated for 1 h at 37 °C with dihydrorhodamine 123 (DHR) fluorescent probe that permeates the cell membrane [44,45].

dHL-60 cells were centrifuged at 180g for 10 min and the pellets were washed in ice for 15 min. Supernatants were collected by centrifugation at 5000g at 4 °C for 10 min. Supernatants were collected and injected into LC-MS/MS system [48]. The analyses were performed in a 6600 Triple-TOF mass spectrometer (AB Sciex, CA) coupled with electrospray ionization source (ESI), operated in positive mode. The post-acquisition MRM-like data used for quantification of GSH (m/z 308.0911 → 179.0462), GSSG (m/z 613.1592 → 355.0741) and in-ternal standard NAC (m/z 164.0000 → 76.0215) were completed with collision energies for each individual transition (22 V, GSH; 32 V, GSSG; 25 V, NAC) and 80 V of declustering potential. Source temperature was 450 °C and spray voltage was set to 5500 V. The chromatographic method was developed in a Nexera UPLC system (Shimadzu; Kyoto) using a Kinetex C18 analytical column (100 mm x 2.10 mm, 2.6 µm) (Phenomenex; Torrance, CA) eluted with a mobile phase of 0.75 mM ammonium formate/0.01% formic acid (A) and methanol (B) at 0.2 mL/min. Elution gradient started with 1% B for 5 min and then increased to 80% B from 5 to 6 min. It was maintained for 4 min and returned to 1% B from 10 to 11 min. Column temperature was set at 25 °C and injection volume was 10 µL. Total peak area of GSH and GSSG was corrected by the internal standard and plotted against a standard curve to obtain absolute concentrations.

2.11. Statistical analysis

The results are presented as mean ± S.E.M of at least three independent experiments. The data were analyzed by one-way analysis of variance (ANOVA) followed by Newman-Keuls Multiple Range when appropriate. P values less than 0.05 (p < 0.05) were considered to be indicative of significance.

3. Results

3.1. Uric acid prevents bacterial clearance and inhibits IL-1β and TNF-α release by dHL-60 cell

We evaluated bacterial clearance by counting the number of colony-forming units (CFU) after incubation with dHL-60 cells at a MOI = 10. Incubation of PA14 with dHL-60 cells for one, two and three hours significantly decreased bacterial CFU number. Interestingly, treatment

\[ = 500 \text{ nm}, \ \lambda_{\text{em}} = 536 \text{ nm} \] was measured in a BioTek Synergy H1 Hybrid microplate reader (Winooski, USA).

2.9. Superoxide detection by dihydroethidine (DHE)

Differentiated HL-60 (1 × 10⁶ cells) were incubated with PA14 (MOI 1:10) in PBS supplemented with glucose in presence of uric acid (0.05–0.5 mM), or diphenyleneiodonium (20 µM, DPI). The reaction of superoxide with DHE (10 µM) was measured by specific production of 2-hydroxysterithidium (2-OH-E⁺). Specific fluorescence of 2-OH-E⁺ by using λex = 396 nm and λem = 579 nm allows the measurement of this product with no interference of other oxidation products like ethidium [47]. Fluorescence kinetics was measured at 37 °C in a BioTek Synergy H1 Hybrid plate reader (Winooski, USA).

2.10. Quantification of GSH and GSSG by LC/MS/MS system

GSH and GSSG were quantified by LC-MS/MS according to Carroll et al. [48] with modifications. dHL-60 cells (5 × 10⁶) were infected with PA14 (MOI 1:10) and incubated for 1 h at 37 °C with different concentrations of uric acid (0.1, 0.2 or 0.5 mM) in PBS glucose pH 7.4. After incubation, extraction buffer was added to a final concentration 0.2%TCA; 0.1 mM DTPA and 0.2 µg/mL N-acetyl cysteine (NAC, internal standard). Samples were incubated on ice for 15 min, vortexed for 45 s and re-incubated on ice for 15 min. The pH of each sample was adjusted to 2.0 by mixing with 200 µL mobile phase A (0.75 mM ammonium formate, 0.01% formic acid). Cellular debris were removed by centrifugation at 5000g at 4 °C for 10 min. Supernatants were collected and injected into LC-MS/MS system [48]. The analyses were performed in a 6600 Triple-TOF mass spectrometer (AB Sciex, CA) coupled with electrospray ionization source (ESI), operated in positive mode. The post-acquisition MRM-like data used for quantification of GSH (m/z 308.0911 → 179.0462), GSSG (m/z 613.1592 → 355.0741) and internal standard NAC (m/z 164.0000 → 76.0215) were completed with collision energies for each individual transition (22 V, GSH; 32 V, GSSG; 25 V, NAC) and 80 V of declustering potential. Source temperature was 450 °C and spray voltage was set to 5500 V. The chromatographic method was developed in a Nexera UPLC system (Shimadzu; Kyoto) using a Kinetex C18 analytical column (100 mm x 2.10 mm, 2.6 µm) (Phenomenex; Torrance, CA) eluted with a mobile phase of 0.75 mM ammonium formate/0.01% formic acid (A) and methanol (B) at 0.2 mL/min. Elution gradient started with 1% B for 5 min and then increased to 80% B from 5 to 6 min. It was maintained for 4 min and returned to 1% B from 10 to 11 min. Column temperature was set at 25 °C and injection volume was 10 µL. Total peak area of GSH and GSSG was corrected by the internal standard and plotted against a standard curve to obtain absolute concentrations.

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with uric acid significantly prevented dHL-60 bacterial clearance (Fig. 1A). Uric acid also significantly decreased the production of IL-1β (Fig. 1B) and TNF-α (Fig. 1C) induced by PA14 after 3 h. The decrease in killing activity and cytokines production was not due to a cytotoxic effect of uric acid upon dHL-60 cells because no increase in prodigium iodide (PI) staining (Fig. 2A) or LDH release (Fig. 2B) was caused by uric acid. As expected, PA14 alone had a cytotoxic effect upon dHL-60 cells after 3 h of incubation. No release of LDH was detected in absence of PA14, even at high concentrations of uric acid (Fig. 2B). Incubation of PA14 with uric acid had no effect on bacterial growth (Supplem. Fig. S1).

3.2. Uric acid disrupts hypochlorous acid production

Because uric acid can either compete with chloride by myeloperoxidase catalysis [15] or directly react with HOCI [17], the inhibition of bacterial clearance could be related to a decrease in the HOCI availability. Thus, we quantified HOCI in dHL-60 challenged with PA14 in absence or presence of uric acid. Incubation of dHL-60 with PA14 did not induce a significant production of HOCI, as detected by taurine-chloramine TNB assay (Fig. 3A). The TNB assay is a good tool to assess absolute values of HOCI, but it is limited to HOCI or chloramines that diffuse to the extracellular space and does not assess HOCI intracellular content. As an alternative, we settled a cell-free myeloperoxidase/Cl-/H₂O₂ system to evaluate the effect of uric acid on HOCI production and bacterial clearance. In this system, the production of HOCI by was dose-dependent and significantly inhibited by uric acid (Fig. 3B).

The myeloperoxidase/Cl-/H₂O₂ system was allowed to react for 30 min and then added to PA14 cells. After 2 h of incubation, no viable bacteria were detected, showing the bactericidal effect of the system. The simultaneous incubation with uric acid significantly prevented the bactericidal effect of the myeloperoxidase/Cl-/H₂O₂ system (Table 1). This result demonstrates that the decrease in HOCI by uric acid, either by inhibiting its formation or by directly scavenging it, contributes to bacteria survival. To evaluate the solely contribution of the scavenger effect of uric acid upon HOCI in bacterial survival, we allowed the formation of HOCI by myeloperoxidase/Cl-/H₂O₂ system for 30 min, then, uric acid was incubated within the system and reacted for 0, 10, or 30 min before adding the bacteria. The total amount of HOCI was: 102.5 ± 2.5; 101.5 ± 3.7; 62.5 ± 2.5; 21.2 ± 3.7 μM for the samples in the absence of uric acid or in the presence of 0.5 mM uric acid reacting for 0 (zero), 10 and 30 min, respectively. In spite of the decrease in HOCI after 10 and 30 min of incubation with uric acid, only a mild protective effect was evidenced. The total CFU in the minimum medium (control); myeloperoxidase/Cl-/H₂O₂; myeloperoxidase/Cl-/H₂O₂ + UA 0 min; myeloperoxidase/Cl-/H₂O₂ + UA 10 min and myeloperoxidase/Cl-/H₂O₂ + UA 30 min were, respectively, 102.5 ± 2.5; 101.5 ± 3.7; 62.5 ± 2.5; 21.2 ± 3.7 μM for the samples in the absence of uric acid or in the presence of 0.5 mM uric acid reacting for 0 (zero), 10 and 30 min, respectively. In spite of the decrease in HOCI after 10 and 30 min of incubation with uric acid, only a mild protective effect was evidenced. The total CFU in the minimum medium (control); myeloperoxidase/Cl-/H₂O₂; myeloperoxidase/Cl-/H₂O₂ + UA 0 min; myeloperoxidase/Cl-/H₂O₂ + UA 10 min and myeloperoxidase/Cl-/H₂O₂ + UA 30 min were, respectively, 3.1 ± 0.4 × 10⁵/mL, 0 (zero), 0 (zero), 0 (zero) and 1.8 ± 0.9 × 10⁵/mL. This shows that the protective effect of uric acid occurs rather by a competition for myeloperoxidase and inhibition of HOCI formation than by a direct neutralization of HOCI. It is possible that the products of reaction between HOCI and uric acid still have some bactericidal effect. However, further studies are needed to address this hypothesis.

Because TNB oxidation is limited to HOCI diffusion to the

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Fig. 1. Uric acid affects bacterial clearance (A), IL-1β (B) and TNF-α (C) release by dHL-60 cells. dHL-60 cells (2 × 10⁵) were challenged with PA14 (MOI 1:10) for 1, 2 and 3 h at 37°C. After each time point serial dilutions were spread on agar plates and the colony-forming units (CFU) were determined after overnight incubation at 37°C (A). The quantification of IL-1β (B) and TNF-α (C) in the supernatants was performed by enzyme-linked immunosorbent assay (ELISA) 3 h after challenging with PA14. Each bar represents mean ± SEM of three independent experiments. Statistical analyses were performed by one-way ANOVA followed by Newman-Keuls post hoc test; #p < 0.05 compared with no dHL-60 group in (A) or with no PA14 group in (B) and (C); *p < 0.05 compared with dHL-60 + PA14 no uric acid group. UA, uric acid.
expected. In spite of that, uric acid also greatly decreased R19-S
gulfment. Therefore, a di-
dmembrane and is not restricted to the phagosome, as in bacterial en-
Fig. S2). The oxidative burst induced by PMA initiates in the plasma
percentage relative to the positive control (dHL-60 in presence of lysis bu-
LDH activity is presented as the
lactate dehydrogenase (LDH) activity in supernatants of dHL-60 incubated or not with
Staurosporine (St) was used as a positive control. (*
performed by one-way analyses of Variance (ANOVA) followed by Newman-Keuls;
represents the mean ± SEM of three independent experiments. Statistical analyses were

extracellular milieu, we estimated the relative concentration of in-
tracellular HOCl with the HOCl-high selective membrane permeable
fluorescent probe R19-S. Incubation of dHL-60 cells with PA14 induced the
appearance of green fluorescence foci, probably originated into the
phagosome. Uric acid or the myeloperoxidase inhibitor, ABAB, de-
creased the number and the fluorescence intensity of foci (Figs. 4A and
B). Differently from dHL-60 cells challenged with PA14, activation with
PMA induced a diffuse and much more intense fluorescence (Supplem
Fig. S2). The oxidative burst induced by PMA initiates in the plasma
membrane and is not restricted to the phagosome, as in bacterial en-
gulment. Therefore, a diffuse rather than a localized fluorescence is
expected. In spite of that, uric acid also greatly decreased R19-S

Fig. 2. Uric acid does not affect dHL-60 cell viability. (A) dHL-60 cells were incubated
with PA14 (MOI 1:10) for one or two hours at 37 °C. Dead cells were stained with PI and the
fluorescence (Oλem = 535 nm, λexc = 620 nm) was measured by flow cytometry.
Staurosporine (St) was used as a positive control. (B) Cytotoxicity was measured by
lactate dehydrogenase (LDH) activity in supernatants of dHL-60 incubated or not with
PA14 (MOI 1:10) and uric acid (UA) for 3 h at 37 °C. LDH activity is presented as the
percentage relative to the positive control (dHL-60 in presence of lysis buffer). Each bar
represents the mean ± SEM of three independent experiments. Statistical analyses were
performed by one-way analyses of Variance (ANOVA) followed by Newman-Keuls;

Fig. 3. Uric acid decreases the production of hypochlorous
acid (HOCl) by the myeloperoxidase/Cl-/H2O2 system. (A) dHL-60 cells (5 x 10^5)
were incubated with PA14 (MOI 1:10) and uric acid for one hour at 37 °C. (B) Myeloperoxidase (100 nM),
NaCl (150 mM), taurine (5 mM) and H2O2 (100 µM) were in-
cubated in minimum medium in the absence or presence of uric
acid for 30 min at room temperature. Reaction was stopped by
incubating with 50 µg/mL catalase. HOCl was indirectly quanti-
ded through the oxidation of TNB (ε1ε20 = 14,200 M^-1 cm^-1) to the colorless DTNB by taurine-chloramine. Each bar
represents the mean ± SEM of three independent experiments. Statistical analyses were performed by one-way ANOVA followed
by Newman-Keuls posthoc test; *p < 0.05 compared to control

Table 1
Uric acid inhibits the bactericidal effect of the myeloperoxidase/H2O2/Cl-

system.

| Treatment                  | CFU/mL x 10^5 |
|----------------------------|---------------|
| Growth minimum medium      | 60.0 ± 12.0   |
| H2O2                      | 49.0 ± 3.0    |
| MPO/Cl-/H2O2               | 0*            |
| MPO/Cl-/H2O2/UA            | 28.5 ± 4.5*   |

Myeloperoxidase (MPO, 100 nM), NaCl (150 mM), H2O2 (100 µM) were in-
cubated or not with uric acid (UA, 0.5 mM) in minimum medium for 30 min.
Catalase (50 µg/mL) was added to remove H2O2 and stop the reaction. PA14
(1 x 10^6/mL) was incubated with this system for 2 h and colony-forming
units (CFU) were counted after overnight growth. Results represent the
mean ± SEM of three independent experiments. Statistical analyses were
performed by one-way ANOVA followed by Newman-Keuls post-hoc test;
*p < 0.05 compared with minimum media and *p < 0.05 compared with
MPO/Cl-/H2O2.

3.3. Uric acid increases superoxide production in dHL-60 cells

The overall production of oxidants by dHL-60 cells was evaluated by a
non-specific rhodamine derivative fluorescent probe, DHR. Fig. 5A
shows that incubation with PA14 induced a significant increase in the
production of oxidants and that uric acid significantly prevented this
oxidation. Although DHR is less specific for HOCl than the R19-S, our in
vitro assay revealed that DHR was fifty and ten-fold more sensitive to
HOCl and to H2O2-horseradish peroxidase, respectively, than to urate
hydroperoxide (data not shown).

In one hand uric acid prevents HOCl formation and plays an anti-
oxidant role, on the other hand its oxidation by myeloperoxidase and
formation of urate free radical and urate hydroperoxide could favor a
pro-oxidation environment. This is particularly relevant in in-
flammatory cells because they express large amounts of myelopero-
xidase and NADPH oxidase. In addition, uric acid has been described to
stimulate NADPH oxidase and increase superoxide production in adi-
poocytes [34] and leucocytes [49]. In our experiments, uric acid sig-
nificantly increased the production of superoxide by dHL-60 challenged
with PA14 (Fig. 5B and C). However, a signi
ificant effect occurred only at 0.5 mM uric acid (Fig. 5C). Superoxide production was signi
ificantly prevented by the flavin adenine dinucleotide (FAD) poison DPI [50,51].
This result confirms a paradoxical effect of uric acid, whereas it exerts
an antioxidant action by decreasing HOCl production, it also acts as a
pro-oxidant by increasing superoxide production.

Therefore, the next step was to evaluate the overall redox state of
dHL-60 in presence and absence of uric acid by measuring the levels of
reduced (GSH) and oxidized (GSSG) glutathione. Uric acid by itself does
not affect baseline levels of GSH and GSSG (Fig. 6). The challenge of

Fig. 4. Uric acid does not affect the number and the
fluorescence intensity of foci. (A) dHL-60 cells were
incubated with PA14 (MOI 1:10) for one hour at 37 °C. Dead cells
were stained with PI and the fluorescence (Oλem = 620 nm, λexc = 535 nm) was
measured by flow cytometry. Staurosporine (St) was used as a positive control.
(B) Differences in the number and fluorescence intensity of foci
measured in the presence and absence of uric acid are shown
in the histograms. Each bar represents the mean ± SEM of three independent experiments. Statistical analyses were
performed by two-way analyses of Variance (ANOVA) followed by Newman-Keuls;

*p < 0.05 compared with control group.
dHL-60 with PA14 increased both GSH and GSSG levels and incubation with uric acid had no effect on them (Fig. 6). This result demonstrated that the net oxidative status is not affected by uric acid. Therefore, the antioxidant effect of uric acid by neutralizing HOCl is likely counterbalanced by the production of other oxidants: uric acid free radical, urate hydroperoxide and superoxide.

4. Discussion

Uric acid is a facile electron donor and is rapidly oxidized by myeloperoxidase and lactoperoxidase to form the single and two electron-oxidants urate free radical and urate hydroperoxide [15, 29]. Because both enzymes are important mediators of the innate immune response, we hypothesized that oxidation of uric acid could contribute against bacterial infection. In addition, soluble uric acid, at the concentration found in human plasma, can activate formation of neutrophil extracellular traps [38], increase production of superoxide [34, 49], activate release of cytokines and recruit inflammatory cells to the site of inflammation [32, 33, 35, 52]. All these events could contribute to a bactericidal adjuvant effect of uric acid. Unexpectedly, incubation of dHL-60 with uric acid significantly inhibited the bactericidal properties of these cells when challenged with PA14 (Fig. 1A). This inhibition was not due to a decrease in dHL-60 viability by uric acid. Differentiated HL-60 cells underwent significant lysis only three hours after incubation with PA14 and uric acid did not potentiate it (Fig. 2). In accordance, previous results show that PA14 did not induce cytotoxicity to murine macrophages at the first and second hour of treatment [53, 54].

The decrease in dHL-60 bactericidal activity by uric acid was likely related to the decrease in HOCl availability. HOCl is the strongest microbicidal produced by neutrophils [2, 7, 9]. It is much more reactive and less diffusion than hydrogen peroxide and, therefore, a proper killing agent inside the phagosome [55]. In agreement to this, inhibition of chloride transport and HOCl production decreased the bactericidal property of neutrophils against P. aeruginosa (PA01) [8].

Uric acid has also been described by disrupting the formation of
hypothiocyanite by lactoperoxidase and the killing activity of the lactoperoxidase/H2O2/thiocyanate system against Pseudomonas aeruginosa [29]. As for lactoperoxidase, uric acid is a substrate for myeloperoxidase-Compound I and Compound II and competes with chloride for the enzyme (Fig. 7) [15]. Comparing the rate constants of the reaction of myeloperoxidase-Compound I with uric acid, \( k = 4.6 \times 10^5 \text{M}^{-1} \text{s}^{-1} \), or chloride \( k = 2.5 \times 10^4 \text{M}^{-1} \text{s}^{-1} \) [11,15] it is likely that, at physiological concentrations of both compounds: uric acid 0.2–0.5 mM and chloride 140 mM, chloride will be the main substrate for the enzyme. Nevertheless, when we incubated myeloperoxidase with physiological concentrations of both competing substrates, we still found a significant decrease in HOCl production despite the large excess of chloride over uric acid (Table 1). In fact, the large Michaelis-Menten constant for myeloperoxidase and chloride reaction, \( K_M = 175 \text{mM} \) [12], might limit the oxidation of this substrate in neutral pH 7.4. A much lower \( K_M \) has been estimated for uric acid reacting with myeloperoxidase (\( K_M = 74 \text{μM} \)) (estimation based on data from [15]) or lactoperoxidase (\( K_M = 100 \text{μM} \)) [29] in the same conditions.

Besides competing for the catalytic activity of myeloperoxidase, uric acid can also directly scavenge HOCl. The estimate rate constant for this reaction at pH 7.0 is \( 3 \times 10^5 \text{M}^{-1} \text{s}^{-1} \) [17,18,56]. Therefore, uric acid can act as a pro-oxidant by propagating free-radical chain reactions [21,23]. One-electron oxidation of uric acid by heme-peroxidases produces urate free radical and urate hydroperoxide [15,29], both intermediates can...
oxidize glutathione, methionine and thiol in proteins, promoting a pro-oxidative redox imbalance [25,26]. In addition, soluble uric acid can increase oxidative status by inducing NAPDH oxidase membrane assemblage and, consequently, superoxide production [34,35,49] and also by increasing reactive oxygen species in mitochondria [52]. In our hands, uric acid increased even further the superoxide production triggered by bacteria engulfment (Fig. 5B and C). The NADPH oxidase is likely the main source of superoxide during phagocytosis. However, further studies are necessary to address what is the main source of superoxide production affected by uric acid.

The increase in superoxide levels could affect myeloperoxidase catalytic activity because superoxide can directly react with the enzyme or can provide the hydrogen peroxide substrate. Superoxide reacts with different myeloperoxidase intermediates [60] and the most relevant reactions in the phagosome have been tested experimentally [61] and by simulation [6]. In the phagosome, the reaction of superoxide with native myeloperoxidase to form myeloperoxidase-compound III accounts for 93% of total myeloperoxidase. The 7% remaining of the native myeloperoxidase is responsible to keep up the rate of HOCl production at nearly maximum [6]. This same simulation proved that a two-fold increase in superoxide slightly decreases HOCl production [6]. Therefore, the 1.5-fold increase in superoxide caused by uric acid in our experiments (Fig. 5C) probably did not significantly affect myeloperoxidase catalytic activity in the phagosome. This reinforces that the main decrease in the levels HOCl is likely due to the competition between chloride and uric acid by myeloperoxidase-compound I (Fig. 7).

Uric acid did not change the GSH/GSSG ratio (Fig. 6C), showing that any antioxidant effect that could result from HOCl disruption might be counterbalanced by the production of oxidants like uric acid free radical, urate hydroperoxide and superoxide. It is noteworthy to mention that the levels of both GSH and GSSG were greatly increased in dHL-60 challenged with PA14. It might reflect de novo synthesis of glutathione since toxins produced by PA14 can up-regulate genes related to oxidative stress [62]. A similar increase in total glutathione was found in macrophages and polymorphonuclear incubated with lipopolysaccharide or Burkholderia pseudomallei [63,64].

A shift in the redox balance can modulate cytokines synthesis, release and their intracellular response [64–68]. In our study, uric acid caused a slight decrease in IL-1β but largely decreased TNF-α level (Fig. 1B and C). An alteration in the redox state could modulate the transcriptional activity of NF-κB, altering the synthesis of IL-1β and TNF-α [69,70]. Whereas some studies have shown induction of NF-κB by HOCl and inhibition by antioxidants [71–74], other studies revealed that a reductive environment is crucial to NF-κB activation [68,75]. Certainly, the modulation of NF-κB is not identical among different cells [70]. In addition, besides phosphorylation, the full transcriptional activity of NF-κB depends on thioredoxin reduction of disulfide bonds in NF-κB p50 subunit that migrates to the nucleus [68,75–77]. Therefore, an increase in the oxidative state caused by uric acid could modulate the transcriptional activity of NF-κB and synthesis of TNF-α. However, alterations on redox balance may not be the solely mechanism for modulation of cytokines synthesis and release by uric acid.

Oxidation of uric acid was positively correlated with bronchiectasis and with myeloperoxidase, neutrophil elastase, methionine sulfioxide, glutathione sulfonamide, chlorotyrosine and inflammatory cytokines levels in bronchoalveolar lavage of children suffering of pulmonary infection by P. aeruginosa. This clinical study strengthened the link between uric acid oxidation and worst outcome in patients with P. aeruginosa infection and suggests the investigation of oxidized glutathione and uric acid as biomarkers of early cystic fibrosis lung disease [39]. A correlation between plasma uric acid levels and worst outcomes has also been found in sepsis [78–81]. However, in some cases it was not possible to identify the independent contribution of uric acid because some patients also presented kidney injury [81].

In conclusion, this study shows that uric acid decreased the levels of HOCl without providing a reductive environment in dHL-60. The decrease in HOCl was likely the main cause of inhibition of the microbicidal activity in these cells, showing the crucial role of HOCl in bacterial killing in an isolated cell system. Uric acid also prevented the release of cytokines and this may contribute to inhibition of autocrine effect that could result from HOCl disruption. The decrease in HOCl could modulate the transcriptional activity of NF-κB and synthesis of TNF-α. However, alterations on redox balance may not be the solely mechanism for modulation of cytokines synthesis and release by uric acid.

**Acknowledgments**

The authors thank Adriano Britto for technical support on quantification of glutathione by MS/MS and Lucia R. Lopes for discussion on superoxide experiments.

**Funding sources**

This study was supported by Fundação de Amparo a Pesquisa do Estado de São Paulo (FAPESP): CEPID Redoxoma 2013/07937-8; Young Investigator 2011/18106-4 and Conselho Nacional de Pesquisa.
Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.redox.2018.02.020.

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