Xanthine oxidase inhibitors and sepsis

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Abstract
Xanthine oxidase activation occurs in sepsis and results in the generation of uric acid (UrAc) and reactive oxygen species (ROS). We aimed to evaluate the effect of xanthine oxidase inhibitors (XOis) in rats stimulated with lipopolysaccharide (LPS). LPS (10 mg/kg) was administered intraperitoneally (i.p.) immediately after allopurinol (Alo, 2 mg/kg) or febuxostat (Feb, 1 mg/kg) every 24 h for 3 days. To increase UrAc levels, oxonic acid (Oxo) was administered by gavage (750 mg/kg per day) for 5 days. Animals were divided into the following 10 groups (n = 6 each): (1) Control, (2) Alo, (3) Feb, (4) LPS, (5) LPSAlo, (6) LPSFeb, (7) Oxo, (8) OxoLPS, (9) OxoLPSAlo, and (10) OxoLPSFeb. Feb with or without Oxo did not aggravate sepsis. LPS administration (with or without Oxo) significantly decreased the creatinine clearance (ClCr) in LPSAlo (60%, P < 0.01) versus LPS (44%, P < 0.05) and LPSFeb (35%, P < 0.05). Furthermore, a significant increase in mortality was observed with LPSAlo (28/34, 82%) compared to LPS treatment alone (10/16, 63%) and LPSFeb (11/17, 65%, P < 0.05). In addition, increased levels of thiobarbituric acid reactive substances (TBARS), tumor necrosis factor (TNF)-α, interleukin (IL)-6, and IL-10 were observed at 72 h compared to the groups that received LPS and LPSFeb with or without Oxo. In this study, coadministration of Alo in LPS-induced experimental sepsis aggravated septic shock, leading to mortality, renal function impairment, and high ROS and proinflammatory IL levels. In contrast, administration of Feb did not potentiate sepsis, probably because it did not interfere with other metabolic events.

Keywords
allopurinol, febuxostat, inflammation, interleukin, lipid peroxidation, LPS, reactive oxygen species, sepsis, septic shock, uric acid, xanthine oxidase inhibitors

Introduction
According to the Third International Consensus 2016, sepsis should be defined as a life-threatening organ dysfunction caused by a dysregulated host response to infection.¹ In addition, septic shock should be defined as a subset of sepsis associated with a greater risk of mortality than sepsis alone,¹ especially when kidney failure is established by acute renal injury and accompanied by multiple organ failure. The Toll-like receptor 4 (TLR4) stimulation caused by lipopolysaccharide (LPS) or TLR 7 and 8 by (+) ssRNA virus induces xanthine oxidase (XO) activation with monosodium urate crystal (i.e. uric acid (UrAc)) and reactive oxygen species (ROS) formation.² XO is also activated by

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urate crystals that stimulate TLR-2 and 4 and hypoxemia due to hemodynamic changes in sepsis, which is commonly observed in ischemia/reperfusion (I/R) models.\textsuperscript{2,3} 

Prior administration of allopurinol (Alo) has a well-documented protective effect in the first 24 h in animal models of sepsis,\textsuperscript{4–11} renal I/R,\textsuperscript{3} liver,\textsuperscript{12,13} intestinal I/R,\textsuperscript{14} tourniquets on rat hind-limbs,\textsuperscript{15} brain I/R,\textsuperscript{16} cocaine-induced diastolic dysfunction,\textsuperscript{17} hyperthermia,\textsuperscript{18} fructose-fed rats,\textsuperscript{19,20} streptozotocin-induced diabetes,\textsuperscript{21} and \textit{Crotalus terrificus} snake venom.\textsuperscript{22} However, Alo did not protect against experimental sepsis,\textsuperscript{23–25} renal I/R,\textsuperscript{26} lung I/R,\textsuperscript{27} zymosan,\textsuperscript{28} or cell culture with LPS,\textsuperscript{29} or in animals with infected burns that showed improvement only on day 1,\textsuperscript{30} as the protection disappeared subsequently. Alo administration for 14 days in patients treated for burns resulted in lower mortality,\textsuperscript{31} but did not protect perioperatively against I/R injury in humans undergoing aortic aneurysm surgery.\textsuperscript{32} (Supplementary Tables 1 and 2).

Although Alo has anti-inflammatory and protective effects in several experimental models, including sepsis, there are still controversial results, especially after 24 h. Its effect has not yet been entirely elucidated following its administration immediately prior to the induction of sepsis, and few studies have compared the effects of diverse xanthine oxidase inhibitors (XOis). Therefore, we conducted experiments to investigate these effects.

Material and methods

\textit{Lipopolysaccharide}

Bacterial LPS from \textit{Escherichia coli} (EC) strain 2630: O111: B$_4$, was obtained from Sigma-Aldrich (St. Louis, MO, USA) and administered intraperitoneally (i.p.) at 10 mg/kg body weight, diluted in distilled water, to rats as one dose every 24 h for 3 days.

\textit{Xanthine oxidase inhibitors}

The following XOis were administered by gavage every 24 h for 3 days: Alo (2 mg/kg, GlaxoSmithKline do Brasil, RJ, Brazil) and febuxostat (Feb, 1 mg/kg, Takeda, IL, USA).

\textit{Oxonic acid}

Oxonic acid (Oxo, Sigma-Aldrich), which was used to increase UrAc levels, was administered by gavage (750 mg/kg per day), diluted in 0.25% methylcellulose in saline, once daily for 5 days before administration of LPS.

Animal ethics

The experimental protocol was approved by the Ethics Committee of the Universidade Federal de S\~{a}o Paulo (0220/12). Each experimental group was composed of six animals.

Experimental groups

Male Wistar rats weighing 200–250 g were divided into 10 groups: (1) Control (received water by gavage), (2) Alo, (3) Feb, (4) LPS, (5) LPSAlo, (6) LPSFeb, (7) Oxo, (8) OxoLPS, (9) OxoLPSAlo, and (10) OxoLPSFeb. At 0 and 17 h of the experimental period, blood samples were collected from the orbital sinus, and the animals were maintained in individual metabolic cages for 24 h for urine collection. Animals were euthanized 72 h after the experiment commenced under anesthesia (ketamine/xylazine, 10:1, 0.2 mL/100 g/kg, i.p.). Blood was collected, and the kidneys were perfused with saline solution, removed, and subsequently histopathologically analyzed.

Biochemical analysis

The blood creatinine (Cr) and UrAc levels were assayed spectrophotometrically according to standard procedures using commercially available diagnostics kits (Labtest Diagnostica, MG, Brazil).

Lipid peroxidation

Lipid peroxidation was measured using the thiobarbituric acid reactive substances (TBARS) assay. The reactive substances combine with thiobarbituric acid to form a red compound. Malondialdehyde (MDA) was used to construct a standard curve, and the results are expressed as MDA (mM)/mg protein. Urine samples were added to a solution of 0.375% thiobarbituric acid, 15% trichloroacetic acid, and 0.25 N HCl (Sigma-Aldrich). The samples were continually agitated while being heated to 95°C for 20 min, and then they were allowed to
cool to 22°C. The absorbance was spectrophotometrically determined at 535 nm. The results were expressed as $10^{-7}$ M/mg creatinine.

**Cytokine analysis**

Immunoassays, based on Luminex xMAP (multianalyte profiling) technology, were performed to simultaneously detect and quantify multiple cytokines (interleukin (IL)-1α, IL-4, IL-1β, IL-2, IL-6, IL-10, and tumor necrosis factor (TNF)-α) using the Cat#RECTYMAG65K-08 granulocyte-macrophage colony-stimulating factor (GM-CSF) kit. The assays were performed at the beginning (time 0), and at 17 and 72 h in all groups, but for Oxo-treated animals, the assays were also performed 5 days before.

**Histopathology**

Hematoxylin and eosin (H&E) staining of the paraffin-embedded kidney tissue slices was performed.

**Immunohistochemistry**

Paraffin-embedded tissues were cut into 4-μm thick sections using a rotary microtome (Leica Microsystems, Herlev, Denmark). The kidney slices were deparaffinized and rehydrated, and then the sections were boiled in a target retrieval solution (1 mmol/L Tris, pH 9.0, with 0.5 mM ethylene glycol tetraacetic acid (EGTA)) for 10 min for antigen retrieval. Nonspecific binding was prevented by incubating the sections in phosphate-buffered saline (PBS) containing 1% bovine serum albumin (BSA), 0.05% saponin, and 0.2% gelatin. Endogenous peroxidase activity was blocked with 5% hydrogen peroxidase (H2O2) in absolute methanol for 10 min at 22°C.

The sections were incubated with primary antibodies against proliferating cell nuclear antigen (PCNA, 1:500) or cleaved caspase-3 (1:100), overnight at 4°C. PCNA is a nuclear protein linked to DNA replication, also known as cyclin, which plays a role in initiating cell proliferation by enhancing the DNA polymerase enzyme. After the sections were washed, they were incubated with horseradish peroxidase–labeled polymer conjugated to a secondary antibody (Dako, Denmark) for 1 h at room temperature. The sites of antibody-antigen reactions were visualized using 0.5% 3,3′-diaminobenzidine tetrachloride (Dako) dissolved in distilled water with 0.1% H2O2. A total of 10 sections per animal along the kidney cortex were analyzed, and labeled cells (staining light to dark brown) were counted.

**Statistical analysis**

The results are expressed as the mean ± standard error (SE). The data were analyzed using survival Kaplan–Meier curves and a one-way analysis of variance (ANOVA) followed by Tukey tests using GraphPad Prism 5. $P < 0.05$ was considered statistically significant.

**Results**

*Coadministration of Alo but not Feb aggravated septic shock by increasing mortality*

LPS administration significantly reduced the mean arterial blood pressure (BP) of rats, measured using a tail-cuff method, in all groups. The LPS group at 72 h showed a mean arterial BP of $95.5 ± 13.2$ (\(P < 0.05\)) and the LPSAlo group at 24, 48, and 72 h showed values of $94.3 ± 12.9$, $97.7 ± 13.6$, and $96.2 ± 15.1$ mmHg, respectively, compared to the control value of $127 ± 7$ mmHg (Figure 1(a)). Alo administration immediately before LPS treatment enhanced the septic shock, which manifested as a significant increase in mortality ($P < 0.05$) in the LPSAlo group (28/34, 82%) compared to the LPS group (10/16, 63%) and LPSFeb group (11/17, 65%, Figure 1(c)).

*Oxo pre-treatment increase BP, blood UrAc levels and decreased mortality*

After 5 days from Oxo administration, a significant increase in BP occurred (from $124.7 ± 4.3$ to $149 ± 20.9$ mmHg, $P < 0.05$, Figure 1(b)) accompanied by a significant increase in blood UrAc levels (from $1.7 ± 0.2$ to $3.0 ± 0.7$ mg/dL, $P < 0.001$, Figure 2(b)). There were no deaths in the OxoLPS or OxoLPSFeb groups compared to the OxoLPSAlo group (12/18, 67%; $P < 0.02$, Figure 1(d)), and we observed a decrease in the number of deaths in the OxoLPSAlo group compared to the LPSAlo group ($P = 0.08$, Figure 1(e)). We noted significantly lower mean values of creatinine clearance (ClCr, mL/min) in all LPS-treated groups (44%, $P < 0.05$), which were the LPSAlo (60%, $P < 0.01$), LPSFeb (35%, $P < 0.05$, Figure 2(c)), OxoLPS (0.61 ± 0.14, $P < 0.01$), OxoLPSFeb (0.64 ± 0.10, $P < 0.01$), and...
OxoLPSAlo (0.57 ± 0.24, *P* < 0.01) groups, compared to the control group (1.25 ± 0.19 mL/min, Figure 2(d)).

**LPS increases ROS especially with coadministration of Oxo**

In addition, we observed high levels of ROS (TBARS: nMol/mg urinary Cr, *P* < 0.001) in all of the groups compared to the control. Furthermore, the values were higher in the Alo-treated groups (LPSAlo, 293.3 ± 35.8 and OxoLPSAlo, 304.3 ± 60.4) than in the LPS (267.8 ± 44.9), LPSFeb (220.0 ± 39.1), OxoLPS (236.7 ± 33.2), and OxoLPSFeb (211.2 ± 43.6) groups (Figure 2(e) and (f)).

**Inflammatory cytokine**

In all the groups who received LPS, the levels of TNF-α increased about three times in relation to control groups in 17 h and returned to basal levels in all groups, except in the LPSAlo and OxoLPSAlo groups where this late decline was not observed (Figure 3(a) and (b); Table 1).

The levels of IL-6 in relation with controls were elevated during 17 h, by 19 times the superior values in the LPS group; by 26 times in the LPSAlo group; by 22 times in the LPSFeb group; by 25 times in the OxoLPS group; by 30 times in the OxoLPSAlo group; and by 25 times in the OxoLPSFeb group. After 72 h, lower IL-6 values were present in about 50% of cases in all groups, except in LPSAlo and OxoLPSAlo groups where these values remained higher, both about 15 times in relation to the control group (Figure 3(c) and (d)).

After 17 and 72 h, all animals treated with LPS alone or with LPS with drugs showed significantly higher IL-1α values than those of the control group, (*P* < 0.001) and LPSAlo, and the OxoLPSAlo group had even higher values than the LPS, LPSFeb, OxoLPS, and OxoLPSFeb groups (*P* < 0.001; Figure 3(e) and (f)).

Compared to the control group after 17 h, all groups showed significantly higher IL-10 levels,
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(P < 0.001) and the IL-10 increase was even higher and more significant after 72 h. After 17 and 72 h, the LPSAlO group showed significantly higher IL-10 levels (P < 0.001) than the LPS and LPSFeb groups. Oxo administration further increased the IL-10 levels in all groups. After 17 h there were no differences among the OxoLPS, OxoLPSAlO, and OxoLPSFeb groups, but after 72 h, the OxoLPSAlO group showed significantly higher IL-10 levels (P < 0.001) than the OxoLPS and OxoLPSFeb groups (Figure 3(i) and (j)).

All groups that received LPS alone or with drugs showed significantly higher IL-2 and GM-CSF levels (P < 0.001; Figure 3(k) and (l)).

Histopathological changes, proliferation (PCNA), and apoptosis rates

Minimal alterations were observed in the histopathological analysis, with rare areas that indicated circulatory disruption where vasodilatation and hemorrhage occurred. However, no significant
Figure 3. Cytokine analysis (pg/mL): TNF-α: (a) LPS, LPSAlo, and LPSFeb groups and (b) Oxo, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. All groups treated with LPS alone or with drugs showed significantly higher values than the control group ***$P < 0.001$. IL-6: (c) LPS, LPSAlo, and LPSFeb groups and (d) Oxo, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. All groups treated with LPS alone or with drugs showed significantly higher values than the control group, *$P < 0.05$, **$P < 0.01$, and ***$P < 0.001$. IL-1α: (e) LPS, LPSAlo, and LPSFeb groups and (f) Oxo, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. All groups treated with LPS alone or with drugs showed significantly higher values than the control group, **$P < 0.01$ and ***$P < 0.001$. IL-1β: (g) LPS, LPSAlo, and LPSFeb groups and (h) Oxo, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. All groups treated with LPS alone or with drugs showed significantly higher values than the control group, **$P < 0.01$ and ***$P < 0.001$. IL-10: (i) LPS, LPSAlo, and LPSFeb groups and (j) Oxo, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. All groups treated with LPS with drugs showed significantly higher values than groups treated with LPS alone after 17 h: *$P < 0.05$, **$P < 0.01$, and ***$P < 0.001$. TNF: tumor necrosis factor; IL: interleukin; LPS: lipopolysaccharide; Alo: allopurinol; Feb: febuxostat; Oxo: oxonic acid; GM-CSF, granulocyte-macrophage colony-stimulating factor.

difference ($P = 0.85$) was observed between the control and all LPS-treated groups, which is in agreement with previously reported data (Figure 4).33 After 72 h, all groups challenged with LPS alone or with drugs showed significantly ($P < 0.001$) lower values for PCNA immunohistochemical staining than the control group (1.087 ± 0.66/μm²), which was consistent with the severity of sepsis34 (Figure 5 and Supplementary Table 3). Tissues sections immunohistochemically stained for cleaved caspase-3 demonstrated a significantly higher apoptosis cell number/field in all groups that received LPS alone or with others drugs compared to the control group ($P < 0.001$; Figure 6 and Supplementary Table 4).

Discussion

Sepsis has a high mortality rate, mainly due to septic shock, aggravated by renal insufficiency and multiple organ failure.3 In this study, we would like to be clear that LPS induces endotoxemia and systemic inflammatory response syndrome (SIRS) but not sepsis as defined in the third consensus definition. LPS-treated groups had lower BP, impaired renal function, and higher ROS levels compared to the control group. Furthermore, immunohistochemistry for PCNA revealed less cell proliferation and that for cleaved caspase-3 indicated considerable apoptosis that accompanied the permanently elevated IL-10 levels related to septic shock.4,34–36 ROS are involved in the caspase-3 apoptosis cascade37 in renal tubular cells in response to systemic sepsis stress, which is a mechanism of acute kidney injury (AKI),35 where increased expression of renal IL-10 has been observed.37

Alo is readily absorbed from the gastrointestinal tract. In our experiment, LPS administration immediately after daily treatment with 2 mg/kg Alo paradoxically worsened the sepsis compared to Alo pre-treatment at doses between 10 and 100 mg/kg body weight from 4 days to 20 min before LPS treatment.5,38–40 This contrasts with the effects observed in the first 24 h, when Alo protected against experimental sepsis.5,38–40 It was previously shown that the protective effects of Alo may disappear after 24 h, with mortality rates reaching 100%.52 Matuschak et al.41 showed the potent suppressive effect of combined hypoxic stress/reperfusion (H/R) and EC stimulation on the activation and nuclear translocation of nuclear factor (NF)-Kb, in
Table 1. Cytokine analysis.

|                | control | Alo | Feb | LPS | LPSAlo | LPSFeb | Oxo | OxoLPS | OxoLPSAlo | OxoLPSFeb |
|----------------|---------|-----|-----|-----|--------|--------|-----|--------|-----------|-----------|
| **TNF-α**      | 0       | 1.3±0.1 | 1.4±0.1 | 1.4±0.1 | 1.4±0.2 | 1.3±0.1 | 1.4±0.1 | 1.7±0.1 | 1.6±0.6 | 1.6±0.2 | 1.6±0.2 |
|                | 17 h    | 1.4±0.1 | 1.3±0.1 | 1.3±0.2 | 4.9±1.0* | 4.5±1.0* | 4.1±0.6* | 1.5±0.1 | 4.8±1.0* | 4.6±0.8* | 4.2±0.7* |
|                | 72 h    | 1.5±0.2 | 1.1±0.2 | 1.2±0.1 | 1.5±0.3 | 4.2±1.0* | 1.4±0.4 | 1.5±0.1 | 1.5±0.3 | 3.4±0.4* | 1.2±0.2 |
| **IL-6**       | 0       | 8.7±1.6 | 8.2±1.5 | 9.1±0.7 | 8.2±1.0 | 8.4±1.0 | 8.3±2.0 | 10.1±0.3 | 10.0±1.6 | 10.0±1.3 | 8.7±1.2 |
|                | 17 h    | 7.7±1.5 | 7.9±1.6 | 9.3±1.2 | 175.7±40 | 238.8±28# | 199.5±23 | 10.1±1.3 | 231.3±19 | 275.7±52## | 229.2±30 |
|                | 72 h    | 4.2±1.0* | 1.4±0.1 | 1.3±0.1 | 1.3±0.1 | 1.3±0.1 | 1.4±0.1 | 1.3±0.1 | 1.4±0.1 | 1.7±0.1 | 1.6±0.6 |
| **IL-1α**      | 0       | 1.7±0.4 | 1.8±0.5 | 1.7±0.4 | 1.7±0.4 | 1.7±0.4 | 1.7±0.4 | 1.7±0.4 | 1.7±0.4 | 1.7±0.4 | 1.7±0.4 |
|                | 17 h    | 1.8±0.5 | 1.7±0.4 | 1.8±0.5 | 16.2±4.7• | 22.2±3.9• | 18.3±5.5• | 2.0±0.0 | 20.7±4.8• | 27.7±3.6*** | 18.7±7.0* |
|                | 72 h    | 1.7±0.6 | 1.5±0.6 | 1.7±0.3 | 17.4±4.4• | 30.7±5.5*** | 15.8±1.6 | 2.0±0.3 | 18.4±3.6* | 35.4±6.2*** | 15.9±2.7* |
| **IL-1β**      | 0       | 36±6   | 31±3  | 33±5  | 33±3   | 32±3   | 34±3   | 37±6   | 38±6   | 39±6   | 38±5   |
|                | 17 h    | 38±2   | 30±4  | 30±5  | 594±64* | 655±72 | 588±66* | 38±1   | 650±86* | 625±54* | 558±42* |
|                | 72 h    | 32±4   | 29±4  | 29±4  | 629±47* | 632±95* | 490±67* | 39±3   | 662±54* | 638±55* | 565±59* |
| **IL-10**      | 0       | 32±5   | 34±8  | 35±14 | 33±3   | 32±3   | 34±3   | 37±6   | 38±4   | 39±6   | 38±5   |
|                | 17 h    | 35±3   | 33±5  | 33±12 | 475±95* | 777±60* | 497±70* | 35±6   | 659±101 | 803±59* | 765±92* |
|                | 72 h    | 31±3   | 34±10 | 34±10 | 756±60* | 1102±231* | 704±118* | 36±3   | 786±111* | 1148±153* | 825±114* |
| **IL-2**       | 0       | 5.2±1.1 | 5.5  | 5.3  | 6.6±0.3 | 5.5±0.6 | 5.7±0.5 | 5.8±1.5 | 5.8±1.3 | 5.7±1.8 | 6.0±1.2 |
|                | 17 h    | 5.9±1.8 | 5.3  | 6.9  | 152.3±25.8* | 241±18.8*** | 144.3±24.1* | 5.9±1.0 | 160.3±12.6* | 264.8±31.0*** | 162.5±16.3* |
|                | 72 h    | 6.3±2.0 | 6.5  | 6.5  | 139.0±12±* | 208.7±32.5*** | 120.5±28.1* | 6.1±1.0 | 147.8±27.6* | 250.5±33.8*** | 156.3±16.9* |
| **GM-CSF**     | 0       | 1.6±0.1 | 1.7±0.5 | 1.6±0.4 | 1.6±0.1 | 1.6±0.3 | 1.6±0.1 | 1.6±0.3 | 1.6±0.3 | 2.0±0.3 | 1.6±0.2 |
|                | 17 h    | 1.6±0.3 | 1.6±0.4 | 1.5±0.4 | 11.3±1.4* | 17.8±3.5* | 13.4±2.5** | 2.0±0.4 | 14.9±2.4* | 21.8±4.7*** | 17.0±2.5* |
|                | 72 h    | 1.6±0.1 | 1.6±0.3 | 1.6±0.3 | 8.8±0.9* | 12.3±1.6* | 10.3±1.0* | 2.1±0.7 | 11.6±1.4* | 18.0±5.0*** | 12.7±1.0* |

GM-CSF: granulocyte-macrophage colony-stimulating factor. TNF-α (pg/mL): in relation to controls: significantly higher values ***P < 0.001; IL-6 (pg/mL): in relation to LPS 17 h lower values **P < 0.01 and higher values *P < 0.05 and ***P < 0.01. Higher IL-6 values compared to LPS 72 h &P < 0.05. IL-1α (pg/mL): in relation to LPS 17 h: significantly higher values *P < 0.01 and ***P < 0.001; •P < 0.001 compared to controls. IL-1β (pg/mL): in relation to control: significantly higher values *P < 0.001 and ***P < 0.001; •P < 0.001 compared to controls. L-10 (pg/mL): in relation to L 17 h: *P < 0.05; **P < 0.01; ***P < 0.001 and •P < 0.001 compared to controls. IL-2 (pg/mL): in relation to LPS 17 h: significantly higher values ***P < 0.001; •P < 0.001 compared to controls. GM-CSF (pg/mL): in relation to LPS 17 h: significantly higher values *P < 0.05; **P < 0.01; ***P < 0.001 and •P < 0.001 compared to controls.
Figure 4. Histopathological sections of kidney were obtained at 72 h (n = 3 rats/group). (a) LPS, normal and (b) LPS, normal, 100× scale bar = 50 µm. (c) LPS, sector with vasodilatation and hemorrhaging (arrow) and (d) LPS, vacuolar degeneration of proximal renal tubules (arrow). 400× scale bar = 20 µm.

Figure 5. Immunohistochemical staining of PCNA (1:500 µm²): (a) Control; (b) LPS; (c) LPSAlo; (d) LPSFeb; (e) OxoLPS; (f) OxoLPS; (g) OxoLPSAlo; and (h) OxoLPSFeb. All groups (i) treated with LPS alone or with drugs demonstrated a significantly lower proliferation rate (**p < 0.001) than the control and Oxo groups. There were no significant differences among the LPS, LPSAlo, LPSFeb, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. PCNA (arrow), LPS: lipopolysaccharide; Alo: allopurinol; Feb: febuxostat; Oxo: oxonic acid. 100× scale bar = 50 µm.
contrast with the robust hypoxic stress-induced transactivation of NF-κB without sepsis. However, administration of the anti-inflammatory compound Alo in H/R + EC increased the NF-κB transactivation, which could, in part, explain the paradoxical effect observed in the LPSAlo and OxoLPSAlo groups.

In cell cultures exposed to LPS, Alo blocked caspase-3 activity and ROS formation in the first 24 h, but this effect disappeared within 48 h, inexplicably. No deaths were observed in the OxoLPS and OxoLPSFeb groups, perhaps because the elevated UrAc level functioned as an antioxidant. UrAc is a potent extracellular antioxidant that evolutionarily confers longevity, maintains BP under conditions with low amounts of dietary sodium, and has a protective effect in multiple sclerosis and severe sepsis. However, it is also an intracellular pro-oxidant, which perhaps explains the increased rates of lipid peroxidation, the levels of proinflammatory ILs and selectively decreases the human blood monocyte production of the natural IL-1 receptor antagonist (IL-1Ra) and changes the production to highly inflammatory IL-1β. In human vaccines, aluminum hydroxide adjuvant boosts adaptive immunity by inducing UrAc and activating inflammatory dendritic cells. Oxo administration for 5 days significantly increased UrAc and BP levels, which was expected. UrAc causes endothelial dysfunction and indirectly decreases levels of nitric oxide, which regulates vascular endothelial cell tone. Furthermore, it activates the renin-angiotensin system and the proliferation of smooth muscle cells by stimulation of inflammatory responses. In the kidney, it triggers afferent arteriopathy, induction of the epithelial-mesenchymal transition, and interstitial tubular fibrosis. Hyperuricemia may be an early marker of sepsis severity.

Alo is the drug of choice for reducing UrAc levels, although it is contraindicated in the initial phase of gout treatment. Recent meta-analyses have shown that continuous use of Alo in patients with hyperuricemia preserves renal function, and

Figure 6. Staining for cleaved caspase-3 (1:100): apoptotic cell number/field. (a) Control; (b) LPS; (c) LPSAlo; (d) LPSFeb; (e) Oxo; (f) OxoLPS; (g) OxoLPSAlo; and (h) OxoLPSFeb. All groups (i) the number of apoptotic cells (arrow) per field was significantly higher in groups treated with LPS alone or with drugs compared to the control group (P < 0.001). There were no significant differences among the LPS, LPSAlo, LPSFeb, OxoLPS, OxoLPSAlo, and OxoLPSFeb groups. LPS: lipopolysaccharide; Alo: allopurinol; Feb: febuxostat; Oxo: oxonic acid. 100× scale bar = 50 µm.
the subsequent slower induction of proteinuria protects against arterial hypertension and slows the progression to chronic kidney disease and mortality. However, some adverse effects such as hepatitis, nephropathies, and hypersensitivity reactions limit the clinical use of Alo. Although occurring at a frequency of <0.1% in humans, hypersensitivity to Alo increases the risk of death by 27%, with the main risk factors as follows: male sex with arterial hypertension and/or renal insufficiency and overdosage, recent initiation of therapy, use of concomitant diuretics or nonhormonal anti-inflammatory drugs, and the presence of HLA-B*5801.

In the LPSAlO group, the shock was possibly triggered by the hemodynamic changes of sepsis evolving into AKI that was aggravated by Alo or the animals developed interstitial nephritis induced by Alo. However, no substrate such as eosinophilia was found in histopathology. There may be another unknown mechanism underlying the paradoxical beneficial therapeutic effect of Alo and negative effect of renal function, as the use of Feb did not worsen the sepsis. Feb does not have structural similarity to purines such as Alo, which may interfere with other stages of purine metabolism.

Feb is an XOi with a favorable toxicological profile and high bioavailability, and it reduces UrAc levels in a more sustained and potent manner than Alo with adverse effects in less than 2% of patients; such effects include diarrhea and increased hepatic transaminases, without serious cutaneous reactions. Feb is primarily metabolized by hepatic oxidation and glucuronidation and eliminated by both the liver and kidney. In contrast to the renal elimination of the active metabolite of Alo (oxypurinol), renal elimination plays a minor role in the elimination of Feb, facilitating safer prescription including for the treatment of tumor lysis syndrome (TLS). Although TLS is classically treated with Alo, studies on Feb in clinical trials have shown a more promising profile. In this experiment, baseline TNF-α levels were approximately 1 pg/mL and increased significantly (P < 0.001) by least four-fold in the first 17 h of LPS administration. The values returned to baseline levels at 72 h, which represents a successful inflammatory response according to the literature as well as the determined IL-2 levels.
ROS and proinflammatory ILs. Clinically, the known risk factors for Alo adverse reactions include recent initiation of therapy and renal failure. In contrast, when we administered Feb, no damage to the sepsis-regulating system was observed, probably because it did not interfere with other metabolic events, which probably occurs with Alo. The prescription of Alo requires physician expertise for use in patients with sepsis who at any time might progress to shock and renal failure. In contrast, Feb, a XOi with hepatic metabolism and renal and hepatic excretion, did not aggravate or improve septic shock with renal failure. Further studies are needed to elucidate the role of XOs in septic shock with renal failure.

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