Intracellular penetration and activity of UB-8902 in human polymorphonuclear leukocytes

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ABSTRACT

The intracellular penetration and activity of UB-8902 in human polymorphonuclear leukocytes was evaluated. Intracellular UB-8902 concentrations were 6-fold higher than extracellular levels. Uptake was rapid, reversible, saturable, and affected by external pH. UB-8902 showed intracellular activity against Staphylococcus aureus strains presenting mutations associated with fluoroquinolone resistance in the gyrA and/or grlA genes.

Penetracion intracelular y actividad de UB-8902 en leucocitos polimorfonucleares humanos

RESUMEN

Se ha evaluado la penetración intracelular y la actividad de UB-8902 en leucocitos polimorfonucleares humanos. La concentración intracellular de UB-8902 fue más de 6 veces superior a la concentración extracellular. La penetración intracellular fue un proceso rápido, reversible, saturable y no se afectó por el pH del medio. UB-8902 mostró actividad intracelular frente a cepas de Staphylococcus aureus que presentaban mutaciones en los genes gyrA y/o grlA, asociadas a resistencia a fluoroquinolonas.

Introduction

The penetration and intracellular activity of antimicrobial agents in phagocytic cells is particularly important in conditions that enable the infecting microorganism to survive and multiply. To treat such infections, it is of clinical interest to use agents able to accumulate and remain active within the cells. A high intracellular accumulation of an antimicrobial agent does not always imply good intracellular activity. Most fluoroquinolones are able to concentrate intracellularly in human phagocytic cells and remain active against facultative and obligate intracellular pathogens such as mycobacteria, Legionella spp, Streptococcus pneumoniae, Listeria monocytogenes and Staphylococcus aureus.1–3 Nonetheless, quinolone resistance has steadily risen, and has proven to be associated with mutations, mainly in the gyrA and parC genes.4 The new quinolone UB-8902, a 7-(4-methyl)-piperazine ciprofloxacin derivative, has been designed to avoid these resistance mechanisms. UB-8902 displays good activity against multiresistant microorganisms including Acinetobacter baumannii and Stenotrophomonas maltophilia.5 The purpose of this study was to evaluate UB-8902 uptake by human polymorphonuclear leukocytes (PMNs), and investigate the mechanisms this antimicrobial agent uses to penetrate these cells. The intracellular activity of UB-8902 against isogenic Staphylococcus aureus strains presenting fluoroquinolone resistance associated with mutations in the gyrA and/or grlA genes was also evaluated.

Material and methods

Isolation of PMNs. PMNs were recovered from heparinized venous blood of healthy donors, and purified by previously
described methods. PMNs were adjusted to 5 x 10^6 cells/mL in Hank's balanced salt solution (HBSS) containing 0.1% gelatine.

**UB-8902 uptake by human PMNs.** The uptake of UB-8902 (provided by Cenavisa, S.A. Laboratories, Reus, Spain) by human PMNs was determined by a fluorometric assay. PMNs were incubated with different concentrations of UB-8902 (1–50 µg/mL). After various incubation periods at 37°C, cells were separated from the extracellular solution by centrifugation and the cell pellet was placed in 0.1 M glycine-HCl buffer to release the intracellular antimicrobial.

The antimicrobial agent was determined by fluorescence emission of the supernatants measured with a fluorescent spectrophotometer. The fluorescence excitation and emission maxima in 0.1 M glycine-HCl were 278 and 447 nm, respectively. After determination of the cell volume with radiolabeled C\(^4\) polyethylene glycol and H\(^2\) water, the accumulation ratio of antimicrobial agent in PMNs was calculated and expressed as the cellular to extracellular concentration (C/E).

**Characterization of UB-8902 uptake.** The influence of cell viability, environmental temperature (4°C vs 37°C), pH (5–8), and the metabolic inhibitors sodium fluoride and sodium cyanide 1.5 x 10\(^{-3}\) M, carbonyl cyanide m-chlorophenyl hydrzone 1.5 x 10\(^{-5}\) M, and 2,4-dinitrophenol, 1 x 10\(^{-4}\) M were evaluated.

In a series of experiments, UB-8902 uptake by human PMNs was measured after stimulating cells with 200 nM phorbol myristate acetate (PMA) and after phagocytosis of opsonized S. aureus (5% pooled human serum at a 10/1 ratio of bacteria to PMN).

The efflux of PMN-associated UB-8902 was also studied. PMNs were incubated for 20 min at 37°C with UB-8902 (2 µg/mL), collected by centrifugation, and re-suspended in antimicrobial-free medium. Cell-associated UB-8902 was quantified at various time intervals after removal of the extracellular antimicrobial agent.

**Organisms and susceptibility testing.** Killing assays were performed in S. aureus 5-61 (wild-type), S. aureus 5-61 G8 (grlA mutant; Ser80Phe), and S. aureus 5-61 M14 (gyrA mutant; Ser80Phe, Glu88Lys and Ser80Phe, respectively). The minimal bactericidal concentrations (MBCs) of UB8902 and moxifloxacin (Hoechst AG) were, respectively, 0.03 and 0.06 µg/mL for S. aureus 5-61, 0.125 and 0.5 µg/mL for S. aureus 5-61 G8, and 0.5 and 4 µg/mL for S. aureus 5-61 M14.

**Intracellular activities of antimicrobial agents.** The intracellular activity of UB-8902 against mutant and wild-type *Staphylococcus aureus* in PMNs compared to that of moxifloxacin was evaluated in a 3-h assay using a previously described method. Opsonized bacteria and PMN (10:1 bacteria/PMN) were incubated for 60 min at 37°C. Extracellular bacteria were removed by differential centrifugation and cells were suspended in RPMI medium. Different concentrations of antimicrobials were then added and cells were re-incubated (3 h, 37°C). Cells were lysed in distilled water, and the colony-forming units (CFU) were counted on Mueller Hinton agar.

**Statistical analysis of data.** Data were expressed as the mean ± standard deviation. Differences between groups were compared by analysis of variance, which was used to assess statistical significance at a p-value of <.05.

**Results**

The C/E ratio for UB-8902 was 6.5 ± 0.8 after 20 min. of incubation at an extracellular concentration of 2 µg/mL. When the extracellular antimicrobial had been removed, PMN-associated UB-8902 decreased by 70% after 5 min of incubation at 37°C (Fig. 1). Cell-associated UB-8902 was saturable at extracellular concentrations higher than 10 µg/mL. The Cmax value previously described for UB-8902 is 7.9 µg/mL.

**Discussion**

High intracellular accumulation of an antimicrobial agent is required to enable eradication of an infecting microorganism.
The intracellular concentration of UB-8902 was 6.5 times higher than the extracellular level, similar to the values reported for ofloxacin, gemifloxacin, and lomefloxacin, and slightly lower than the values for other quinolones such as DX-619, trovafloxacin, and moxifloxacin. UB-8902 uptake was rapid, saturable, and reversible.

Cell viability and environmental temperature had no effect on penetration of UB-8902 into phagocytic cells. PMN uptake of UB-8902 increased significantly at an acidic pH. Of the inhibitors evaluated, only sodium cyanide significantly impaired intracellular UB-8902 penetration. It is interesting that sodium cyanide, an inhibitor of mitochondrial oxidative metabolism, reduced UB-8902 uptake by PMN, as these cells have few mitochondria, being independent of oxidative metabolism for energy. As reported for ofloxacin and its isomers, these data point to a possible passive mechanism involved in human PMN uptake of this quinolone. The effects of phagocytosis and cell membrane stimulation on UB-8902 uptake were similar to those previously observed with trovafloxacin and sparflxacin. Ingestion of opsonized S. aureus cells and cell membrane stimulation with PMA did not modify intracellular penetration of UB-8902.

At all extracellular concentrations evaluated, UB-8902 significantly reduced intracellular survival of the wild-type strain. The reduced intracellular survival of the two mutant strains was only significant at the higher extracellular concentrations evaluated (1 and 5 μg/mL). The intracellular activity of UB-8902 against the two mutant strains was similar. These results are comparable to those observed for moxifloxacin. However, moxifloxacin showed slightly higher intracellular activity than UB-8902 against the wild-type and mutant strains studied. This effect could be partly due to greater intrinsic activity and to the higher intracellular accumulation of moxifloxacin compared to UB-8902.

In summary, UB-8902 penetrates human PMNs and attains intracellular concentrations several fold higher than extracellular concentrations. A passive mechanism may be involved in UB-8902 uptake by PMN. At extracellular concentrations greater than 1 μg/mL, UB-8902 remains intracellularly active against fluoroquinolone-resistant S. aureus strains with mutations in the gyrA and/or grlA genes.

Conflict of interest

The authors have no conflicts of interest to declare.

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References