Prevalence of hypermutators among clinical Acinetobacter baumannii isolates

Patricia Komp Lindgren1*, Paul G. Higgins2, Harald Seifert2 and Otto Cars1

1Department of Medical Sciences, Section of Infectious Diseases, University of Uppsala, 75105 Uppsala, Sweden; 2Institute for Medical Microbiology, Immunology and Hygiene, University of Cologne, 50935 Cologne, Germany

*Corresponding author. Tel: +46-18-6119360; Fax: +46-18-559157; E-mail: patriciakomplindgren@gmail.com

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Objectives: The objectives of this study were to study the presence of mutators in a set of Acinetobacter baumannii isolates and to explore whether there is a correlation between mutation rates and antibiotic resistance.

Methods: The variation in mutation rate was evaluated for 237 clinical A. baumannii isolates by determining the frequency of their mutation to rifampicin resistance. For each isolate, the antibiotic resistance profile was determined by disc diffusion and/or Etest. Isolates were divided into susceptible, resistant and MDR groups according to their resistance to five groups of different antibiotics. A comparison between differences in mutation frequency (f) and strain-specific factors was performed.

Results: Of the 237 isolates 32%, 18% and 50% were classified as susceptible, resistant and MDR, respectively. The f of rifampicin resistance varied between 2.2 × 10^-21 and 1.2 × 10^-6. Of the strains under investigation, 16% had an ≥2.5- to 166-fold higher f. The presence of mutators (definition ≥2.5-fold increase in f compared with ATCC 19606) in the MDR group (22%) was significantly higher (P≤0.05) than that in the susceptible and resistant groups (11% and 7%, respectively). Furthermore, f was significantly higher in the MDR group compared with that in the susceptible and resistant groups.

Conclusions: The facts that 26 of 37 mutator isolates (70%) in the population were MDR and that there was a significantly higher general f in isolates exhibiting an MDR profile suggest that hypermutability can be of advantage for the organism in a selective environment with extensive exposure to antimicrobials.

Introduction

Acinetobacter baumannii is increasingly involved in nosocomial infections, especially in the ICU setting. This pathogen is frequently MDR and has shown a propensity to adapt to the environment through up-regulation of intrinsic resistance and acquisition of antibiotic resistance determinants.1,2 Another parameter that modulates the progression of resistance is an increase in general mutation rate (hypermutability), which can influence resistance development through mutation,3 frequency of lateral gene transfer,4 up-regulated gene expression and compensatory evolution of a bacterial population.5

Studies of mutator distribution and level of hypermutability have shown that ≤50% of clinical isolates from a variety of bacterial species exhibit a mutator phenotype/genotype. Many of these mutators are found among infection-causing bacteria, which are isolated from an environment of high selective pressure due to intense antibiotic usage.6 No previous study has evaluated the existence of mutator isolates in an A. baumannii population, to our knowledge. The aims of this study were to determine the prevalence of mutators in A. baumannii and their hypermutability and to explore whether there is any correlation between changes in the mutation frequency (f) and antibiotic resistance.

Materials and methods

Bacterial isolates, antimicrobials and culture media

Two-hundred-and-thirty-seven A. baumannii isolates originating from Europe, the USA and Sweden between 1990 and 2007 were investigated.7,8 A. baumannii ATCC 19606 was chosen as a comparator strain for f assays due to its isolation in 1948 and exposure to less antibiotic selection pressure. The WT Escherichia coli MG1655, E. coli dam (damΔ16::Kanr) and mutS (mutS::FRT) mutants were used as quality-control strains.9 All experiments were performed in freshly prepared Mueller-Hinton (MH) broth and MH agar (BD, Sweden); rifampicin was purchased from Sigma-Aldrich, Sweden.
Susceptibility testing

Resistance profiles were determined using Etest (Biodisk/bioMerieux, Sweden) and/or disc diffusion (BD/Fisher Scientific, Sweden). The EUCAST clinical breakpoints were used for sorting isolates into antibiotic-resistant profile groups (ARPGs) as follows: susceptible, susceptible to all antibiotic groups; resistant, resistant to one or two antibiotic groups; and MDR, resistant to three to five antibiotic groups. ARPGs are defined in Table 1.

Determination of f against rifampicin

To determine f, assays were performed as previously described by Baquero et al. Ten independent cultures for each isolate were grown overnight with shaking, and cells were plated onto MH plates with and without rifampicin (100 mg/L). The distribution of f values in the population was categorized as previously described (Table 1). The f values were normalized to control strain ATCC 19606 \((f = 7 \times 10^{-9})\), which was set at 1. Isolates with an \( \geq 2.5\) -to 10-fold higher f were defined as weak mutators and those with a >10-fold higher f were defined as strong mutators.

Data comparison and statistical methods

Isolates were grouped according to their ARPGs and further into hypomutators \((f < 2.5\)-fold) and mutators \((f \geq 2.5\)-fold) before analysis. The Mann–Whitney U-test and Kruskal–Wallis analysis of variance (ANOVA) statistical methods were used to compare resistance groups. For associations between ARPGs and the number of mutator isolates in each group, the Pearson’s \(x^2\) test was used, and logistic regression was used to further investigate the relationship between some of these variables. P values <0.05 were considered significant.

Results

Antimicrobial susceptibility profiles

The division of isolates and the percentage of each isolate in each ARPG are shown in Table 1. The overall occurrence of weak and strong mutators was 80% and 20%, respectively. The resistant group were spread over the whole interval, and for the MDR group, \~45% of the isolates were clustered in a sharp peak, with \( f \) ranging between \( 8 \times 10^{-9} \) and \( 2 \times 10^{-8} \). The distribution of f values in bacterial populations

Distribution of f values in bacterial populations

The frequency of rifampicin resistance was determined, and isolates were sorted according to their f, which varied between \( 2.2 \times 10^{-10} \) and \( 1.2 \times 10^{-6} \) (Figure 1). A sharp peak in frequency distribution was found at \( \sim 10^{-8} \), however, 68% of the population had a lower frequency. To the right of the peak, 15% of the isolates had a moderately higher f, whereas two isolates (<0.1%) had a frequency \( >10^{-7} \) (Figure 1a). No A. baumannii isolate displayed hypermutability as high as that of the E. coli mutS \( f = 3 \times 10^{-8} \), and only one isolate \( f = 1.1 \times 10^{-8} \) had a frequency higher than that of the E. coli dam-knockout strain \( f = 3.9 \times 10^{-7} \). When the population was sorted according to their ARPGs, the distribution of the susceptible group showed three peaks, at \( 2 \times 10^{-9} \), \( 6 \times 10^{-9} \) and \( 1 \times 10^{-8} \), respectively, while the frequencies of the resistant group were spread over the whole interval, and for the MDR group, \~45% of the isolates were clustered in a sharp peak, with \( f \) ranging between \( 8 \times 10^{-9} \) and \( 2 \times 10^{-8} \) (Figure 1b).

Hypermutability and resistance profile in mutator isolates

Compared with the A. baumannii ATCC 19606 control strain, 200 isolates had an f ranging between 0.03-fold and \~2.5-fold. Of these, 80 were classified as hypomutators (<1.0-fold), composed of 39% susceptible, 19% resistant and 43% MDR isolates, respectively. The overall occurrence of weak and strong mutators

| ARPG (n) | Susceptible (75) | Resistant (43) | MDR (119) | Total (237) |
|---------|-----------------|---------------|-----------|-------------|
| TET     | 32              | 16 (7)        | 18 (63)   | 100         |
| CEP     | 0               | 18 (54)       | 50 (92)   | 108         |
| CAR     | 0               | 0             | 0         | 0           |
| AMG     | 0               | 0             | 0         | 0           |
| FQ      | 0               | 0             | 0         | 0           |

Table 1. Allocation of 237 isolates into ARPGs and the median and geometric mean f of isolates in ARPGs

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was \( \sim 16\% \); when the population was divided into ARPGs, the prevalence of mutators in the MDR group (22%) was significantly higher \((P < 0.05)\) than the prevalence in the susceptible and resistant groups (11% and 7%, respectively). The geometric mean and median of \( f \) for the population and the percentage of mutators in each ARPG are presented in Table 1.

Thirty-seven isolates had an \( \geq 2.5\)- to 166-fold increase in \( f \) (73% with \( \geq 2.5\)- to 5-fold, 19% with \( > 5\)- to 10-fold and 8%...
with >10-fold (Table S1, available as Supplementary data at JAC Online). Of the mutators, 70% were MDR isolates with a median/mean increase of 3.6/11.1-fold (Figure 2a). There was no significant difference in f between the mutators in the susceptible, resistant and MDR groups (P=0.51–0.74) (Figure 2a). However, there was a significant correlation between MDR and increased f (P<0.05) (Figure 2b).

Of the 34 weak mutators, 7, 3 and 24 of the isolates had a susceptible, resistant and MDR phenotype, respectively. All the weak resistant mutators (f=3.19–4.29-fold) were susceptible to cephalosporins and carbapenems, with a varied range of susceptibility to the other ARPGs. Four weak MDR mutators (f=2.51–3.25-fold), which displayed antibiotic resistance to three of five antibiotic groups, were all susceptible to the carbapenem group in combination with susceptibility to either the tetracycline-like, aminoglycoside or fluoroquinolone group of antibiotics. Of the remaining 20 weak MDR isolates, nine were resistant to all antibiotic groups (f=3.14–8.29-fold), 5 were resistant to all except tetracycline-like antibiotics (f=2.57–5.89-fold) and 6 were resistant to all except carbapenem-group antibiotics (f=2.71–10.0-fold) (Table S1).

The remaining three strong mutators, one susceptible isolate and two MDR isolates, displayed 11.1-fold (AB95), 19.5-fold (AB279) and 166-fold (AB190) increased f (Table S1). In comparison with the control mutator E. coli strains (dam and mutS mutants, 55.7- and 429-fold increased f), the AB190 isolate can be regarded as a strong mutator.

### Discussion

This study is the first one, to our knowledge, on the occurrence of mutators in an A. baumannii population, and we have shown that 16% of isolates from our collection displayed an ≥2.5- to 166-fold increase in f. The prevalence of mutators and their f values were significantly higher in the MDR group compared with those of the susceptible and resistant groups. We also found 0.42% and 15.2% of isolates to be strong and weak mutators, respectively.11,13

Variation in mutation rates and its effect on resistance development and bacterial adaptability have been studied extensively over recent years. It was shown that even a small (2–4-fold) increase in mutation rate can drive the evolution of fluoroquinolone resistance, and, furthermore, a slight increase in f favours the evolution of MDR.9,14 In E. coli, selection with rifampicin and ciprofloxacin showed that mutator strains generated both higher resistance levels and resistance mutations with <1000-fold higher f.15

Studies of clinical isolates have revealed that weak mutators can be present at earlier stages of infection, and a modestly elevated mutation rate can give them an adaptive advantage.16

Another aspect of mutators is the notion that some genotypes can exhibit increased frequency of recombination, interspecies recombination and transformation.15,16 These events might explain in part the ability of A. baumannii to adapt, whereby a mutator can integrate DNA that carries resistance markers and/or increase the chance of gaining mutations that promote the survival of the organism in clinical settings.

Carbapenems are still the drugs of choice to treat A. baumannii infections, even though resistance rates are increasing.2 To understand the progression of carbapenem resistance, Zander et al.19 investigated three isolates recovered from patients during an outbreak in a hospital in Krakow, Poland. Sequencing of blaOXA-51-like genes showed that carbapenem resistance was caused by a conversion of OXA-66 into OXA-82 and that blaOXA-82 was also associated with the IS element ISAba1. A second study looked at related isolates possessing similar plasmids that encode the carbapenemase OXA-58 and exhibit varying levels of carbapenem resistance. Sequencing revealed genetic variability composed of multiple copies of the blaOXA-58 gene and that extra copies were due to IS-element transposition or recombination events.20 These adaptive modifications could be elevated in a population composed of mutators, where hypermutability can drive the progression of survival and evolution of genetic elements such as β-lactamase genes. However, whether any of the genetic changes seen in these studies was due to an altered mutation rate is not known.

In conclusion, we have shown that 16% of the A. baumannii strains were weak-to-strong mutators and that there was a strong correlation with an increased f with an MDR phenotype. The fact that a high percentage of MDR A. baumannii isolates show an increased mutability in clinical settings calls for further studies that could form the basis of novel treatment strategies.

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### Transparency declarations

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### Supplementary data

Table S1 is available as Supplementary data at JAC Online (http://jac.oxfordjournals.org/).

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