Evaluation of Rapid Polymyxin Pseudomonas test in clinical Pseudomonas aeruginosa isolates with various degrees of multidrug resistance

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Background: Pseudomonas aeruginosa is of great concern among MDR bacteria and rapid and reliable in vitro antibiotic susceptibility testing methods are extremely necessary. Colistin is, in many cases, among the limited useful alternatives for these isolates. Unfortunately, only a few reliable in vitro methods are validated for testing susceptibility to colistin. Although EUCAST and CLSI recommend broth microdilution (BMD) as the standard method for antibiotic susceptibility testing, this method is not routinely performed in microbiology laboratories. However, some commercial products based upon BMD have tested well and offer consistent results.

Objectives: To evaluate the performance of the colorimetric Rapid Polymyxin Pseudomonas Test (RPPT) (ELITech Microbiology, France).

Methods: Eighty-seven clinical P. aeruginosa strains, prospectively collected in two microbiology laboratories exhibiting either susceptibility or various degrees of multidrug resistance, including to colistin, were used. Different susceptibility testing methods were simultaneously performed and compared with reference BMD and interpreted using 2020 EUCAST criteria.

Results: Results indicate an essential agreement (EA) of 97.7% for RPPT while the other tests did not reach 90% of EA [66.7% MicroScan, 63.2% Etest (bioMérieux, France) and 60.9% other MIC Test Strips (MTS, Liofilchem, Italy)]. The categorical agreement was 98.9% for RPPT, 87.4% for MTS, 85.1% for Etest and 64.4% for MicroScan.

Conclusions: The RPPT was able to accurately detect both colistin-susceptible and -resistant isolates within 4 h, offering a rapid alternative for a prompt decision about the inclusion of this antibiotic in a patient’s treatment.

Introduction

MDR microorganisms have increased in recent years, mainly among clinically important Gram-negative bacilli, as a result of which few therapeutic alternatives are left. Because of this, polymyxins, and more specifically colistin, regained their place as an option for treatment use.1

In vitro susceptibility testing of colistin is problematic due to certain chemical characteristics of the antibiotic, mainly its net cationic nature and its high molecular weight hindering its agar diffusion. Moreover, heteroresistance to colistin is a frequent behaviour observed in many bacterial populations.2 Taking all this into account, in vitro results and the resulting susceptibility status have a strong impact in clinical and stewardship decisions, impacting colistin’s clinical use. The reference method, according to EUCAST and CLSI is broth microdilution (BMD) following the ISO standard 20776-1. This method is laborious for daily laboratory practice and results are read after 18–20 h, thus reinforcing the need for other reliable and, if possible, rapid methods.

A comparative study was recently published in order to evaluate seven commercially available products for colistin
antimicrobial susceptibility testing (AST). In that study, five commercial BMD products and two gradient strip tests were compared with the reference method. The group concluded that commercial broth microdilution methods generally performed well, however, the performance of the two gradient tests was unacceptable. They advised laboratories not to trust colistin gradient tests or disc diffusion and recommended the use of broth microdilution-based methods. After this work, a warning from EUCAST has been issued about commercial methods to test colistin.

The Rapid Polymyxin\textsuperscript{TM} Pseudomonas test (RPPT) (ELITech Microbiology, France) is a liquid colorimetric method with freeze-dried colistin (2, 4 and 8 mg/L concentration in each well) that also contains a glucose culture medium and bromocresol purple as a pH indicator. If the isolate is able to grow in a defined colistin concentration, a colour change (green to purple) is evidenced. Results can be read less than 4 h after inoculation. A positive and a negative control are included. The objective of this study was to evaluate the performance of the colorimetric RPPT to confirm the colistin susceptibility status in a prospectively collected \textit{Pseudomonas aeruginosa} isolates exhibiting either susceptibility or various degrees of multidrug resistance, including to colistin, in two clinical microbiology laboratories in comparison with different AST methods.

**Materials and methods**

AST was performed on 75 prospectively collected \textit{P. aeruginosa} isolates that were recovered in routine daily work at the Microbiology Department, Ramón y Cajal University Hospital in Madrid (Spain) in the last quarter of 2018 and 12 isolates recovered in a contemporary period at the Microbiology Department of the Son Espases University Hospital in Palma de Mallorca (Spain). A total of 93.1\% (\(n = 81\)) isolates came from clinical samples (respiratory samples, 33; urine, 23; organic fluid, 8; abscess, 7; blood, 5; exudate, 2; and prosthesis, wound and biopsy, 1 each) and 6.9\% (\(n = 6\)) from rectal colonization. Among the group of 75 isolates, 54 exhibited various degrees of resistance: 43 MDR, 8 XDR, and 3 pandrug resistant (PDR) according to the results obtained with the routinely used automated commercial MIC gradient tests, Etest and MTS. MEs were only found with gradient tests (50%). In 12 isolates: 2 MDR, 1 XDR and 1 PDR (all from respiratory samples and 1 from organic fluid). In only one non-MDR isolate, the colistin BMD MIC value corresponded to the ATU category. The correlation of resistance MIC values obtained with the reference BMD was good for the RPPT but poor for automated microdilution and MIC gradient tests results (Figure 1).

Overall, an EA of 97.7\% was obtained for RPPT while the other tests did not reach the minimum 90\% of EA required (66.7\% for MicroScan, 63.2\% for Etest and 60.9\% for MTS). The CA was 98.9\% for RPPT, 87.4\% for MTS, 85.1\% for Etest and 64.4\% for MicroScan. Considering error classes, a 38.7\% rate of MEs (false resistant results) was observed when the isolates were tested with MicroScan, 8.0\% with MTS, 6.7\% with Etest and 1.3\% with RPPT. A 66.7\% rate of VMEs (false susceptible results) was observed when the isolates were tested by Etest, 41.7\% when using MTS and 16.7\% with MicroScan. The acceptable inter-method error percentage of VMEs and MEs is ≤1.5\% and ≤3\%, respectively. The \(\chi^2\) coefficient for the different methods when compared with the reference BMD was: 0.953 (almost perfect) for RPPT, 0.486 (moderate) for MTS, 0.230 (fair) for Etest and 0.298 (fair) for MicroScan.

When considering different levels of colistin resistance, i.e. non-MDR, MDR, XDR and PDR, an EA of 93.9\% was obtained for RPPT with the non-MDR isolates, conversely, the worst results for this group (lowest CA) were obtained with MicroScan (24.2\%), which had a tendency to overestimate MIC values (Table 1). For MDR isolates, EA was 100\% for RPPT, 90.7\% for MicroScan, 72.1\% for MTS and 53.5\% for Etest; CA was 97.7\% for RPPT, 97.7\% for Etest and 90.7\% for both MicroScan and MTS while MEs were found with MicroScan (9.8\%), MTS (7.3\%) and both RPPT and Etest (2.4\%); VMEs were observed only with the two gradient tests (50\%). In XDR isolates, EA was 100\% for RPPT and MicroScan and 50\% for Etest and MTS. CA was 100\% for RPPT and MicroScan and 87.5\% for Etest and MTS. MEs were only found with gradient tests.
No VMEs were found in this category. Finally, in PDR isolates, EA was 100% for RPPT and 33.3% for MicroScan, Etest and MTS. CA was 100% for RPPT, Etest and MTS and 33.3% for MicroScan. A 100% rate of MEs was detected with MicroScan. No VMEs were found in this category.

The quality control strain results were within expected ranges for all test and methods except for *E. coli* ATCC 25922 with Etest, for which MICs were below the range in all studies (data not shown).

Discussion

Clinical microbiology laboratories require the implementation of a reliable and, if possible, rapid test to determine colistin susceptibility and to detect colistin-resistant isolates. At present, very few tests are validated that effectively detect colistin resistance apart from the standard ISO BMD, which cannot be performed routinely because it is a laborious and time-consuming method.

The RPPT method has previously been validated in other studies, although these were performed with collections of *P. aeruginosa* or with colistin-resistant *P. aeruginosa* generated through *in vitro* experiments, while our studied isolates had been prospectively collected from clinical samples reflecting the epidemiological overview of our institution.

According to our results, the RPPT performance for detecting the colistin-susceptibility status of *P. aeruginosa* isolates is promising. This test appears rapid and reliable for laboratory routine testing, being a particularly useful tool in the case of MDR, XDR and PDR isolates for which colistin may be one of the few therapeutic options left.

In daily clinical practice, we determine colistin MICs using MicroScan panels, however, it has been observed that, although this system shows excellent results with Enterobacterales, non-susceptible results for non-fermenters require confirmation by another validated method due to the high rate of false resistance observed. The same has been concluded for current MIC gradient strips due the inaccuracy of the results.

Our study has some limitations. The small number of colistin-resistant *P. aeruginosa* included in this study can limit definite conclusions about the true performance of the RPPT. In addition, most issues with colistin testing were found among the non-MDR isolates but this does not constitute a relevant problem as the antibiotic is not used for infections caused by these types of isolates. At present, colistin resistance is low in our hospital, even among MDR *P. aeruginosa* isolates. Nevertheless, the consistency of the results obtained with RPPT when compared with the reference method is strongly promising, thus constituting a reliable alternative method for routine work. It is also important to remark that results are obtained in less than 4 h, which is also an advantage when compared with the standard BMD method that needs an overnight wait for the results.


Table 1. Performance of four commercial methods when compared with the reference BMD method

| Method      | MDR isolates (N = 43) | XDR isolates (N = 8) | PDR isolates (N = 3) | Non-MDR isolates (N = 33) | All isolates (N = 87) |
|-------------|-----------------------|----------------------|----------------------|---------------------------|------------------------|
|             | RPPT                  | MicroScan            | Etest                | MTS                       |                        |
| Essential agreement, n (%) | 43 (100)             | 39 (90.7)            | 23 (53.5)            | 31 (72.1)                 | 85 (97.7)              |
|             | (CST R, n = 2)        | (CST R, n = 1)       | (CST R, n = 1)       | (CST R, n = 1)            | (CST R, n = 12)        |
| Categorical agreement, n (%) | 39 (90.7)             | 31 (93.3)            | 18 (54.5)            | 17 (51.5)                 | 58 (66.7)              |
| Major errors, n (%) | 23 (53.5)             | 8 (33.3)             | 1 (33.3)             | 1 (33.3)                  | 1 (33.3)               |
|             | (CST R, n = 1)        | (CST R, n = 1)       | (CST R, n = 1)       | (CST R, n = 1)            | (CST R, n = 12)        |
| Very major errors, n (%) | 23 (53.5)             | 8 (33.3)             | 1 (33.3)             | 1 (33.3)                  | 1 (1.3)                |
|             | (CST R, n = 1)        | (CST R, n = 1)       | (CST R, n = 1)       | (CST R, n = 1)            | (CST R, n = 12)        |

Total P. aeruginosa isolates (n = 87) are split by resistance according to the percentages of essential agreement and categorical agreement, and the number of major errors and very major errors.

CST R, colistin-resistant isolates.

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Transparency declarations

None to declare.

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