

# Activity of SPR206, a Polymyxin Derivative, Compared to Colistin Alone and in Combination Against Multidrug-Resistant Pseudomonas aeruginosa Strains

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### Introduction

Background: The emergence of multidrug-resistant (MDR) P. aeruginosa, has forced clinicians to resort to polymyxin antibiotics (polymyxin B and colistin (COL)), previously discarded due to harmful adverse effects associated with their use (1).

Motivation: Despite their resurgence in clinical treatment, the polymyxins are continually characterized by their side effect profile (2). SPR206 is a polymyxin analogue, however the side chain and tail has been extensively modified, decreasing the potential for adverse events. SPR206 has been shown to have reduced minimum inhibitory concentrations (MIC; MIC<sub>50</sub> and MIC<sub>90</sub>), values for *P. aeruginosa* when compared to COL, and other Gramnegative agents. (3).

Objective: The objective of this study was to compare the *in-vitro* activity of SPR206 to COL both alone and in combination with other Gram-negative antimicrobials against MDR P. aeruginosa strains through MIC susceptibility testing and time-kill experiments (TKE)

Significance: As MDR P. aeruginosa infections increase patient mortality and morbidity, it is important that we are equipped with both safe and efficacious novel therapeutic options.

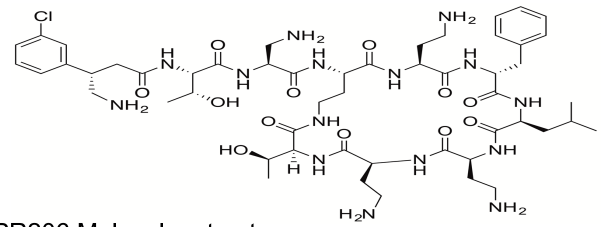


Figure 1. SPR206 Molecular structure

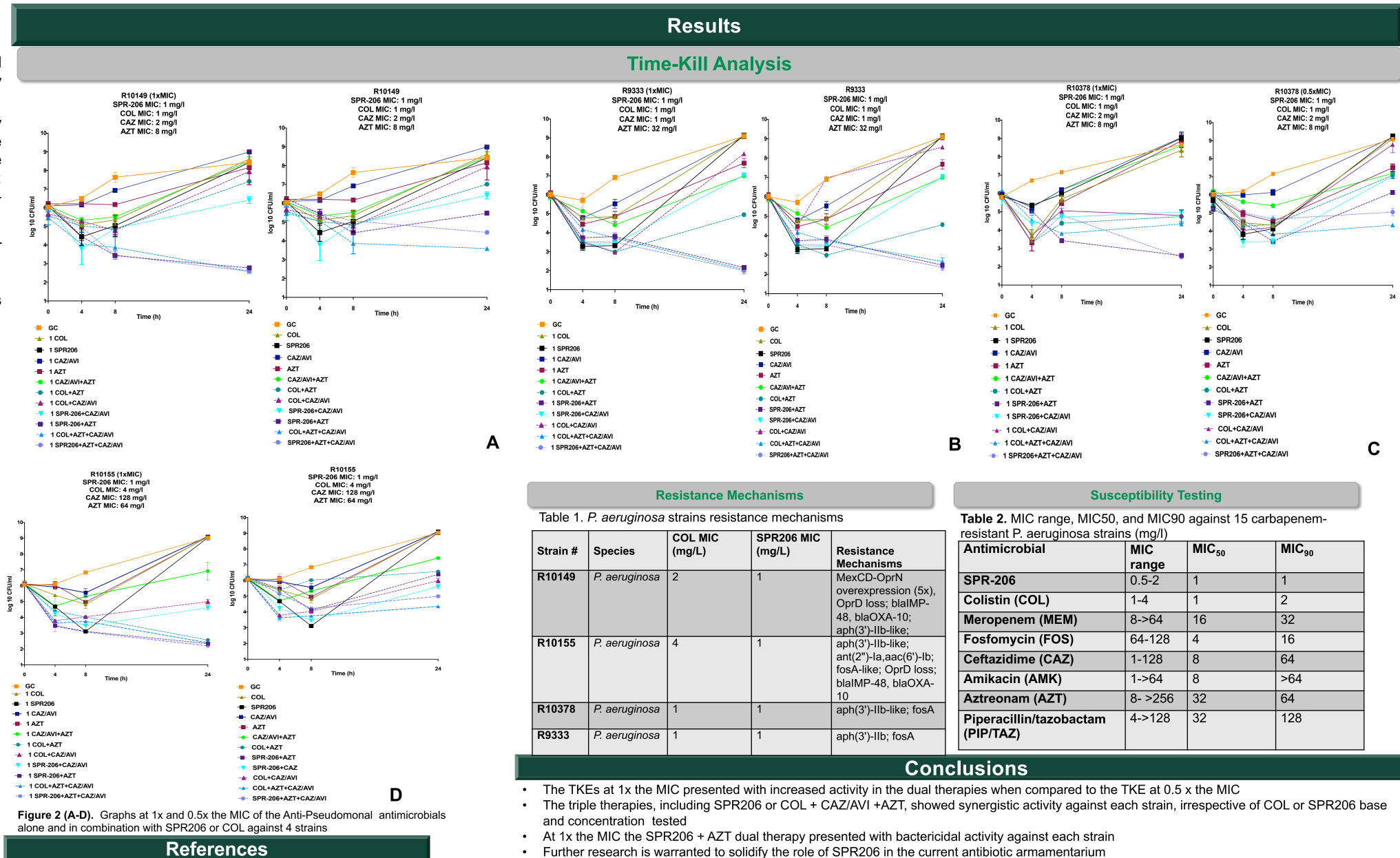
### Methods

Bacterial strains: Fifteen carbapenem (meropenem (MEM) MIC >8mg/l) P. aeruginosa strains were evaluated using MIC susceptibility testing and TKE.

Media/ Antibiotics: COL, MEM, Fosfomycin (FOS), Amikacin (AMK), piperacillin//tazobactam (PIP/TAZ), aztreonam (AZT), ceftazidime (CAZ) were purchased commercially from Sigma Chemical Co. (St. Louis, MO, USA), Avibactam (AVI) was purchased from Fisher scientific SPR206 was obtained from Spero Therapeutics Cambridge, Massachusetts

Susceptibility Testing: MIC values were determined by broth micro-dilution in duplicate, per the current Clinical Laboratory and Standards Institute (CLSI) Guidelines for all strains. MIC testing via broth microdilution was performed for SPR-206, COL, MEM, FOS, AMK, AZT, CAZ/AVI, and PIP/TAZ. Avi was supplemented at a 4:1 ratio to CAZ.

Time-Kill Experiments:. Dual therapy and triple therapy combinations, either COL or SPR206-based, were tested against four representative strains in 24h time-kill experiments (TKE). Each antibiotic was tested at 1x the MIC, or the peak concentration, whichever was lower. A >2 log10 CFU/ml was defined as synergistic activity, and a >3log10 CFU/ml was defined as bactericidal activity.



Sievert, DM et al., Infection Control Hosp Epidemiology.2013. 34:1-14.

- Evans, BA., et al., Curr Pharm Des. 2013. 19:223-38.
- Grosser, L. et al. ASM Microbe 2019. 4. Melvin P., et al., *Performance Standards for Antimicrobial Susceptibility Testing*, 28th
- Edition. 2018.

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## Conclusions

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robial	MIC range	MIC <sub>50</sub>	MIC <sub>90</sub>
5	0.5-2	1	1
(COL)	1-4	1	2
nem (MEM)	8->64	16	32
/cin (FOS)	64-128	4	16
ime (CAZ)	1-128	8	64
n (AMK)	1->64	8	>64
am (AZT)	8- >256	32	64
llin/tazobactam <u>(</u> )	4->128	32	128