In Vitro Activity of the Extended-Spectrum ß-Lactamase Inhibitor Enmetazobactam (formerly AAI101), in Combination with Cefepime, against 90 Molecularly Characterized Enterobacteriaceae Isolates **Expressing a Variety of Non-ß-Lactam Resistance Mechanisms**

Michael D. Huband¹, Amy A. Watters¹, Jill M. Lindley¹, Lalitagauri M. Deshpande¹, A Belley², Philipp Knechtle², Robert K. Flamm¹

¹ JMI Laboratories, North Liberty, Iowa, USA; ² Allecra Therapeutics SAS, Saint-Louis, France

Introduction

- Development of resistance due to the spread of extended-spectrum ß-lactamases (ESBLs) among Enterobacteriaceae has created the need for new therapeutic modalities
- Enmetazobactam (formerly AAI101) is a novel extended-spectrum β-lactamase inhibitor that is highly active against ESBLs and other ß-lactamases
- *In vitro*, cefepime-enmetazobactam outperforms piperacillin-tazobactam against ESBL-producing isolates of Enterobacteriaceae, and is as potent as meropenem
- Enmetazobactam, in combination with cefepime, is intended as empiric treatment of serious Gram-negative infections in settings with an elevated prevalence of ESBLproducing Enterobacteriaceae
- Cefepime (2 g) combined with enmetazobactam (0.5 g) administered q8h has recently entered Phase 3 clinical trials in patients with complicated urinary infections, including pyelonephritis
- In this study, the combination of cefepime-enmetazobactam (enmetazobactam at fixed 8 mg/L) and comparator agents were tested against 90 molecularly characterized Enterobacteriaceae isolates containing a variety of non ß-lactam resistance mechanisms
- Non-ß-lactam resistance mechanisms included: aminoglycoside resistance, fluoroquinolone resistance, colistin resistance (intrinsic and plasmid-mediated), tetracycline-resistance, and resistance due to porin alterations/deletions
- Non ß-lactam resistance isolates also expressed ß-lactam resistance mechanisms including: ESBLs (phenotype/genotype), AmpCs (plasmid encoded), and OXA-type ß-lactamases

Materials and Methods

- A challenge set of 90 molecularly characterized Enterobacteriaceae isolates exhibiting phenotypic cross-resistance to numerous antibiotic classes were recovered from patients with documented infections during 2013–2017 and consisted of 43 Escherichia coli, 34 Klebsiella pneumoniae, 3 Citrobacter freundii species complex, 3 Proteus mirabilis, 2 Enterobacter cloacae species complex, 2 Providencia stuartii, 2 Serratia marcescens, and 1 Klebsiella oxytoca
- Isolates were collected in 56 medical centres located in 19 countries and 9 US census divisions: the United States (29 medical centres; 48 isolates; 53.3% overall), Europe (14 medical centres; 25 isolates; 27.8% overall), Latin America (7 medical centres; 9 isolates; 10.0% overall), and the Asia-Pacific region (6 medical centres; 8 isolates: 8.9% overall)
- Organisms were identified by matrix-assisted laser desorption ionization-time of flight mass spectrometry and molecularly characterized via next-generation sequencing
- Reference broth microdilution susceptibility testing was conducted using frozen-form panels produced by JMI Laboratories according to CLSI M07 (2018) guidelines using cation-adjusted Mueller-Hinton broth
- Isolates were tested for susceptibility to cefepime-enmetazobactam (enmetazobactam fixed at 8 mg/L), cefepime, ceftazidime, ciprofloxacin, colistin, gentamicin, meropenem, piperacillin-tazobactam, tetracycline, and trimethoprim-sulfamethoxazole
- Isolates containing class A carbapenemases and/or class B metallo-ß-lactamases were excluded from the study due to the limited activity of cefepime-enmetazobactam against these resistance mechanisms
- For comparison purposes, CLSI susceptible-dose-dependent (SDD) breakpoint interpretive criteria for cefepime (susceptible MIC ≤ 8 mg/L) were also applied to cefepime-enmetazobactam (fixed 8 mg/L) combinations

Results

- tazobactam (Table 2)
- and Figures 1–4)

- (Table 2)



Cefepime-enmetazobactam (MIC_{50/00}, 0.12/2 mg/L) was the most active compound tested based on MIC⁹⁰ values against 90 molecularly characterized Enterobacteriaceae isolates (Table 1)

Applying the CLSI SDD breakpoint of 8 mg/L to cefepime-enmetazobactam (fixed 8 mg/L) resulted in 97.8% (88/90) of the isolates testing susceptible (S) compared to 50.0%S for cefepime alone, 86.7%S for meropenem, and 58.9%S for piperacillin-

Cefepime-enmetazobactam (fixed 8 mg/L) was the most active compound tested against aminoglycoside-R (MIC_{50/90}, 0.12/2 mg/L; 100.0%S), colistin-R (MIC_{50/90}, 0.25/8 mg/L; 92.0%S), fluoroquinolone-R (MIC_{50/90}, 0.12/4 mg/L; 95.9%S), and tetracycline-R (MIC_{50/90}, 0.12/2 mg/L; 98.0%S) Enterobacteriaceae isolates (Table 2

Cefepime-enmetazobactam (fixed 8 mg/L) was the most active compound tested against molecularly characterized Enterobacteriaceae isolates containing an AmpC cephalosporinase (MIC_{50/90}, 0.25/4 mg/L; 94.1%S), ESBL phenotype/genotype (MIC_{50/90}, 0.12/2 mg/L; 96.8%S), OXA- β -lactamase (MIC_{50/90}, 0.12/4 mg/L; 94.1%S), and/or porin alterations/deletions (MIC_{50/90}, 0.5/16 mg/L; 87.5%S) (Table 2) Cefepime-enmetazobactam (fixed 8 mg/L; 97.8%S) outperformed cefepime, meropenem, and piperacillin-tazobactam against 90 molecularly characterized Enterobacteriaceae isolates

MIC₀₀ values for cefepime, meropenem, and piperacillin-tazobactam were in the resistant interpretive category with values of >64 mg/L (50.0%S), 4 mg/L (86.7%S), and >512 mg/L (58.9%S), respectively

37.8% of the Enterobacteriaceae isolates tested were aminoglycoside resistant (R). 27.8% were colistin R, 54.4% were fluoroquinolone R, and 55.6% were tetracycline R

18.9% of the 90 molecularly characterized isolates contained an AmpC cephalosporinase, 68.9% had an ESBL phenotype or genotype, 37.8% had an OXAtype β -lactamase, and 17.8% contained porin mutations (Table 2)

Figure 1 Cumulative percent inhibition results for cefepimeenmetazobactam (fixed 8 mg/L) and comparators against 34 aminoglycoside-resistant Enterobacteriaceae isolates

Figure 2 Cumulative percent inhibition results for cefepimeenmetazobactam (fixed 8 mg/L) and comparators against 25 colistin-resistant Enterobacteriaceae isolates



Table 1 Cumulative percent inhibition results for cefepime-enmetazobactam (fixed 8 mg/L) and comparators against 90 molecularly characterized Enterobacteriaceae isolates

Organism (organism group (no. of isolatos)	No. and cumulative % of isolates inhibited at MIC (mg/L) of:														МІС						
Organishi/ organishi group (no. or isolates)	≤0.004	0.008	0.015	0.03	0.06	0.12	0.25	0.5	1	2	4	8	16	32	64	128	256	512	> ^a	50	1VIIC ₉₀
All Enterobacteriaceae (90)																					
Cefepime-enmetazobactam (fixed 8 mg/L) (90)		0 0.0	2 2.2	13 16.7	26 45.6	17 64.4	9 74.4	10 85.6	1 86.7	5 92.2	3 95.6	2 97.8	1 98.9	1 100.0						0.12	2
Cefepime (90)			0 0.0	1 1.1	3 4.4	3 7.8	6 14.4	4 18.9	2 21.1	9 31.1	8 40.0	9 50.0	14 65.6	4 70.0	7 77.8				20 100.0	8	>64
Ciprofloxacin (90)	0 0.0	5 5.6	8 14.4	2 16.7	3 20.0	2 22.2	3 25.6	6 32.2	7 40.0	5 45.6	2 47.8	3 51.1	6 57.8						38 100.0	8	>16
Colistin (90)						1 1.1	49 55.6	14 71.1	0 71.1	1 72.2	3 75.6	4 80.0	4 84.4						14 100.0	0.25	>16
Gentamicin (90)						2 2.2	15 18.9	26 47.8	9 57.8	1 58.9	0 58.9	3 62.2	4 66.7	11 78.9	10 90.0				9 100.0	1	64
Meropenem (90)		0 0.0	11 12.2	35 51.1	20 73.3	6 80.0	1 81.1	2 83.3	3 86.7	2 88.9	2 91.1	4 95.6	1 96.7	2 98.9	1 100.0					0.03	4
Piperacillin-tazobactam (90)							1 1.1	3 4.4	6 11.1	17 30.0	13 44.4	7 52.2	6 58.9	5 64.4	5 70.0	4 74.4	5 80.0	5 85.6	13 100.0	8	>512
Tetracycline (90)								0 0.0	7 7.8	23 33.3	6 40.0	4 44.4	3 47.8	0 47.8	7 55.6				40 100.0	64	>64
Trimethoprim-sulfamethoxazole (90)							36 40.0	9 50.0	5 55.6	0 55.6	3 58.9	0 58.9							37 100.0	0.5	>8

MIC value is greater than the highest concentration tester $^{\circ}$ MIC₆₀ value listed in bold.

isolates

	MIC _{50/90} (mg/L)/% susceptible (CLSI) ^a													
Organism/group (n)	FPE ^b	FEP ^c	CIP	COLd	GEN	MEM	PIP-TAZ	TMP-SMX	TET					
Enterobacteriaceae (90)	0.12 / 2	8 / >64	8 / >16	0.25 / >16	1 / 64	0.03 / 4	8 / >512	0.5 / >8	64 / >64					
	97.8% ^e	50.0%	40.0%	72.2%	58.9%	86.7%	58.9%	55.6%	40.0%					
Aminoglycoside resistant (34) ^f	0.12 / 2	16 / >64	>16 / >16	0.5 / >16	64 / >64	0.06 / 4	16 / >512	>8 / >8	64 / >64					
	100.0%	35.3%	8.8%	55.9%	0.0%	82.4%	52.9%	32.4%	35.3%					
Colistin resistant (25) ^g	0.25 / 8	16 / >64	16 / >16	>16 / >16	16 / >64	0.06 / 16	32 / >512	>8 / >8	64 / >64					
	92.0%	40.0%	20.0%	0.0%	36.0%	56.0%	48.0%	36.0%	24.0%					
Fluoroquinolone resistant (49) ^h	0.12 / 4	16 / >64	>16 / >16	0.5 / >16	32 / >64	0.06 / 8	16 / >512	>8 / >8	>64 / >64					
	95.9%	36.7%	0.0%	63.3%	38.8%	81.6%	55.1%	38.8%	34.7%					
Tetracycline resistant (50) ⁱ	0.12 / 2	16 / >64	16 / >16	0.5 / >16	1 / 64	0.03 / 2	8 / >512	>8 / >8	>64 / >64					
	98.0%	48.0%	32.0%	64.0%	56.0%	86.0%	58.0%	44.0%	0.0%					
Amp-C (17) ^j	0.25 / 4	2 / >64	1 / >16	0.25 / >16	1 / 64	0.06 / 8	128 / >512	1 / >8	16 / >64					
	94.1%	70.6%	52.9%	70.6%	70.6%	76.5%	35.3%	52.9%	47.1%					
ESBL (62) ^k	0.12 / 2	16 / >64	>16 / >16	0.25 / >16	1 / >64	0.03 / 8	16 / >512	1 / >8	64 / >64					
	96.8%	29.0%	29.0%	74.2%	54.8%	85.5%	56.5%	51.6%	41.9%					
OXA-1/30, 2, 9, 10, and/or 48 (34) ¹	0.12 / 4	16 / >64	>16 / >16	0.5 / >16	32 / >64	0.06 / 8	64 / >512	>8 / >8	16 / >64					
	94.1%	41.2%	20.6%	58.8%	32.4%	73.5%	35.3%	35.3%	38.2%					
Porin mutations (16) ^m	0.5 / 16	64 / >64	>16 / >16	16 / >16	16 / >64	0.06 / 16	256 / >512	>8 / >8	16 / >64					
	87.5%	31.2%	6.2%	31.2%	37.5%	56.2%	25.0%	25.0%	37.5%					
Multidrug-resistant (46) ⁿ	0.25 / 4	16 / >64	>16 / >16	0.5 / >16	32 / >64	0.06 / 8	32 / >512	>8 / >8	>64 / >64					
	95.7%	34.8%	10.9%	50.0%	23.9%	76.1%	41.3%	34.8	26.1%					

^a FPE, cefepime-enmetazobactam; FEP, cefepime; CIP, ciprofloxacin; COL, colistin; GEN, gentamicin; MEM, meropenem; PIP-TAZ, piperacillin-tazobactam; TMP-SMX, trimethoprim-sulfamethoxazole; TET, tetracycline; ESBL, extended-spectrum ß-lactamase ^b Cefepime susceptible-dose-dependent breakpoint for Enterobacteriaceae applied to cefepime-enmetazobactam (fixed 8 mg/L) for comparison purposes only. ^c Cefepime susceptible-dose-dependent breakpoint applied.

^d Percentage of wild type based on the epidemiological cutoff value for colistin. Susceptibility >90.0% listed in hold

Composed of Citrobacter freundii species complex (1), Enterobacter cloacae species complex (1), Escherichia coli (12), Klebsiella pneumoniae (16), Proteus mirabilis (2), and Providencia stuartii (2). Composed of E. coli (7), K. pneumoniae (11), P. mirabilis (3), P. stuartii (2), and Serratia marcescens (2 Composed of C. freundii species complex (1), Escherichia coli (25), K. pneumoniae (19), P. mirabilis (2), P. stuartii (2). Composed of C. freundii species complex (2), E. cloacae species complex (1), Escherichia coli (25), K. pneumoniae (16), P. mirabilis (3), P. stuartii (2), and S. marcescens (1) Composed of E. coli (10), Klebsiella oxytoca (1), and K. pneumoniae (6).

Composed of *C. freundii* species complex (1), *E. cloacae* species complex (2), *Escherichia* coli (29), *K. oxytoca* (1), *K. pneumoniae* (28), and *P. mirabilis* (1). Composed of E. cloacae species complex (2), Escherichia coli (7), K. oxytoca (1), K. pneumoniae (22), P. mirabilis (1), and S. marcescens (1). Composed of C. freundii species complex (1), E. coli (4), K. pneumoniae (10), and P. stuartii (1).

Figure 3 Cumulative percent inhibition results for cefepimeenmetazobactam (fixed 8 mg/L) and comparators against 49 fluoroquinolone-resistant Enterobacteriaceae isolates



^o Composed of 43 Escherichia coli, 34 Klebsiella pneumoniae, 3 Citrobacter freundii species complex, 3 Proteus mirabilis, 2 Enterobacter cloacae species complex, 2 Providencia stuartii, 2 Serratia marcescens, and 1 Klebsiella oxytoca.

Table 2 Activity of cefepime-enmetazobactam (fixed 8 mg/L) and comparators against 90 recent molecularly characterized Enterobacteriaceae

Composed of C. freundii species complex (1), E. cloacae species complex (1), Escherichia coli (16), K. pneumoniae (22), P. mirabilis (3), P. stuartii (2), and S. marcescens (1).

Figure 4 Cumulative percent inhibition results for cefepimeenmetazobactam (fixed 8 mg/L) and comparators against 50 tetracycline-resistant Enterobacteriaceae isolates



Conclusions

- The novel extended-spectrum ß-lactamase inhibitor enmetazobactam, combined with cefepime, was highly active against 90 recent molecularly characterized Enterobacteriaceae isolates expressing a variety of ß-lactam and non-ß-lactam resistance mechanisms
- Cefepime-enmetazobactam (enmetazobactam fixed at 8 mg/L) activity was unaffected by resistance mechanisms targeting aminoglycoside, fluoroquinolone, colistin, and tetracycline classes of antibacterial agents
- Cefepime-enmetazobactam (fixed 8 mg/L) outperformed cefepime, meropenem, and piperacillin-tazobactam against this collection of molecularly characterized Enterobacteriaceae isolates, including isolates containing AmpC, ESBL, and/or OXA-ß-lactamases
- Cefepime-enmetazobactam may represent a novel empiric option for treatment of serious Gram-negative infections in settings that have high rates of ESBL-producing Enterobacteriaceae and that restrict carbapenem use as part of antimicrobial stewardship measures

Acknowledgements

This study and poster presentation were funded by a grant from Allecra Therapeutics SAS (Saint-Louis, France)

References

Clinical and Laboratory Standards Institute (2018). M100-S28. Performance standards for antimicrobial susceptibility testing: 28th informational supplement. Wayne, PA: CLSI. Clinical and Laboratory Standards Institute (2018). M07-A11. Methods for dilution antimicrobial susceptibility tests for bacteria that grow aerobically; approved standard - eleventh edition. Wayne, PA: CLSI.

Contact

Michael D. Huband JMI Laboratories 345 Beaver Kreek Centre, Suite A North Liberty, IA 52317 Phone: (319) 665-3370 Fax: (319) 665-3371 Email: michael-huband@jmilabs.com



To obtain a PDF of this poster: Scan the QR code or visit https://www .jmilabs.com/data/posters/ECCMID19 -enmetazobactam-cefepime.pdf Charges may apply. No personal information is stored.