Strategies to combat resistance: Focus on pharmacokinetics/ pharmacodynamics with applications to β-lactams





Bangalore – 14 February 2011



Strategies to combat resistance: Focus on pharmacokinetics/pharmacodynamics with applications to β-lactams

Paul M. Tulkens Unité de pharmacologie cellulaire et moléculaire Université catholique de Louvain, Brussels, Belgium & International Society of Antiinfective Pharmacology



http://www.facm.ucl.ac.be

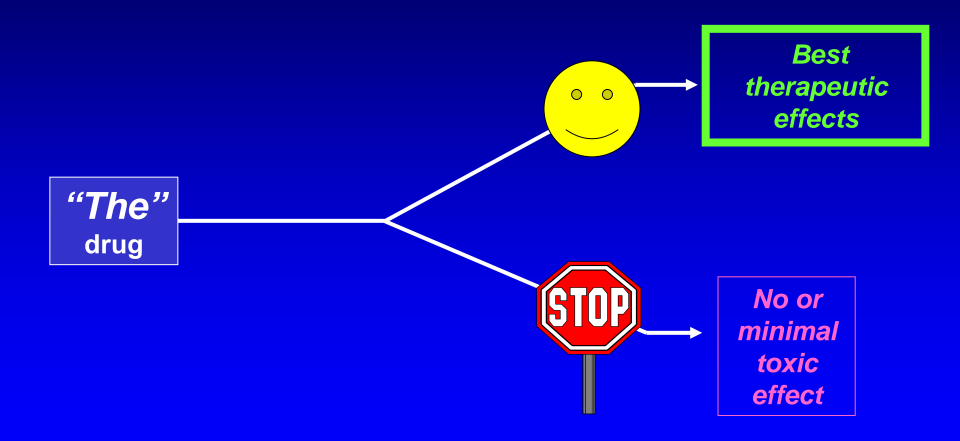




http://www.isap.org

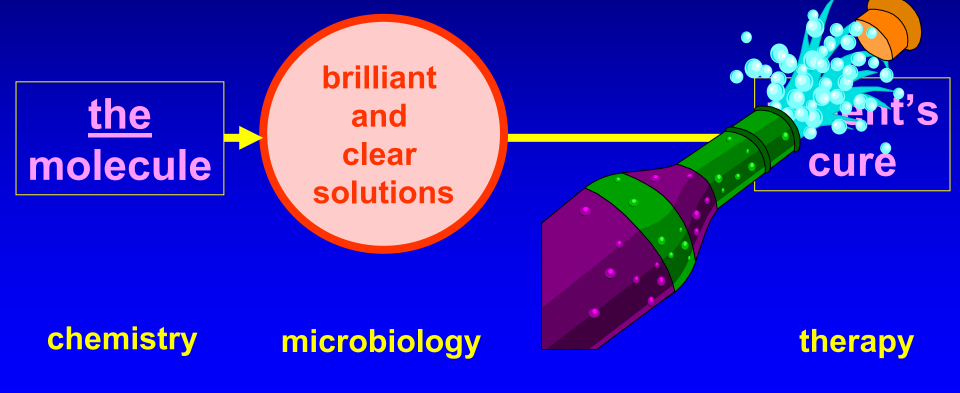


Antibiotic treatment: Wat does the clinician want?



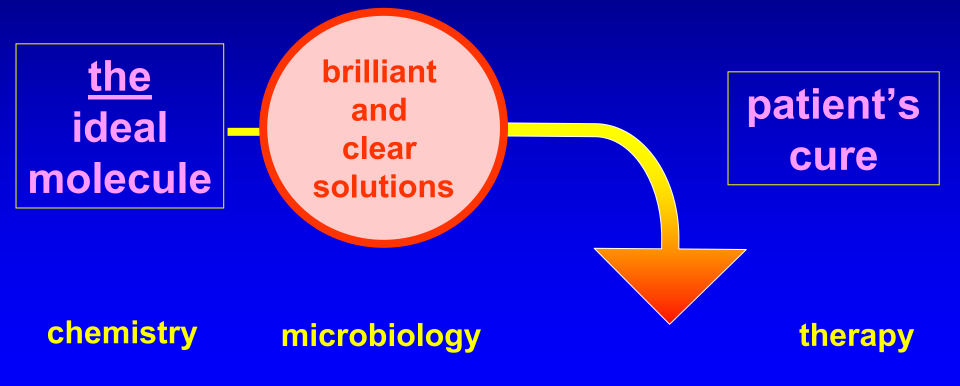


The ideal antibiotic ...





Is <u>the</u> molecule always ideal ?





Main causes of antibiotic failures...

Adapted from Pechère J.C., 1988, 1993, 1998

False failures

- erroneous diagnosis
- underlying disease uninfluenced by antibiotics
- unjustified lack of patience
- inactivation of the antibiotic

Patient related failures

- compliance failure (broadly speaking)
- inappropriate administration route (broadly speaking)
- immunodepressed hosts

- Pharmacological failures
 - insufficient amount or drug inappropriately administered
 - no attention paid to pharmacodynamic parameters
 - in situ inactivation or lack of drainage
- Micro-organism related failures
 - wrong pathogen
 - resistance acquired during treatment
 - insufficient bactericidal activity
 - inoculum effect



In a nutshell ... so far ...

Microbiology parameters: MIC !

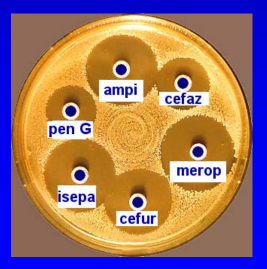
- Pharmacodynamic parameters
- PK/PD as applied to beta-lactams: Time-above MIC
- The problems if you underdose
- Take home message



Microbiology



identification



susceptibility

by static techniques drug concentration stays constant



What do I do in my country (in relation to microbiology) ?

- Survey the level of resistance of *P. aeruginosa* and *S. pneumoniae* from selected hospitals and relate it to therapy ¹
- Examine the mechanisms of resistance acquisition (with special reference to efflux pumps)²

Assess new antibiotics and novel approaches (immunotherapy)³

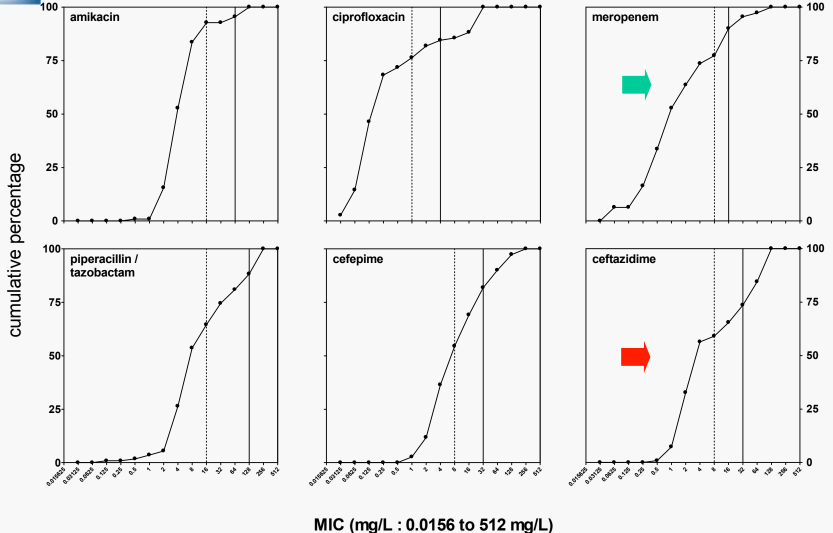
Examine the susceptibility to biocides ¹

Supported by

- 1 Regional authorities and the Fund for Industrial Research
- 2 Fund for Scientific and Medical Research
- 3 Pharlmaceutical Industry and small/medium enterprizes



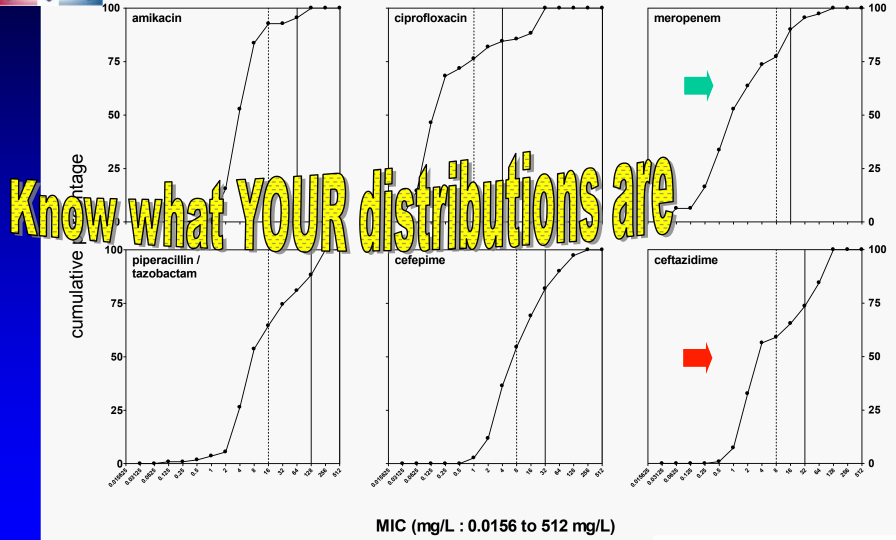
What is the situation at day 0 with P. aeruginosa in HAP ?



Riou et al. IJAA 2010; 36:513-522



What is the situation at day 0 with P. aeruginosa in HAP ?



Riou et al. IJAA 2010; 36:513-522

Moving on ...

- Does your microbiologist discuss infection cases in ICU with you ?
 - Each case
 Few cases
 - 3. Upon asking
 - 4. Never



Asking the question you always wanted to ask ...

- Does your microbiologist gives MIC of antibiotics apart from sensitivity in ICU infections ?
 - 1. Each case
 - 2. Few cases
 - 3. upon asking
 - 4. Never



Asking the question you always wanted to ask ...

- Does your microbiologist gives MIC of antibiotics apart from sensitivity in ICU infections ?
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 - 2. Few cases
 - 3. upon asking
 - 4. Never



No, MIC is not the acronym for "<u>M</u>inimal Interest to the <u>C</u>linician" J



What did the textbooks say about antibiotic dosages and schedules in the 70's ?

- 1. Stay above the MIC... but how much ?
- 2. Remain around for a while... but how long ?
- 3. Hope it works... against everything ?
- 4. Hope it is not toxic...

can't do much ...

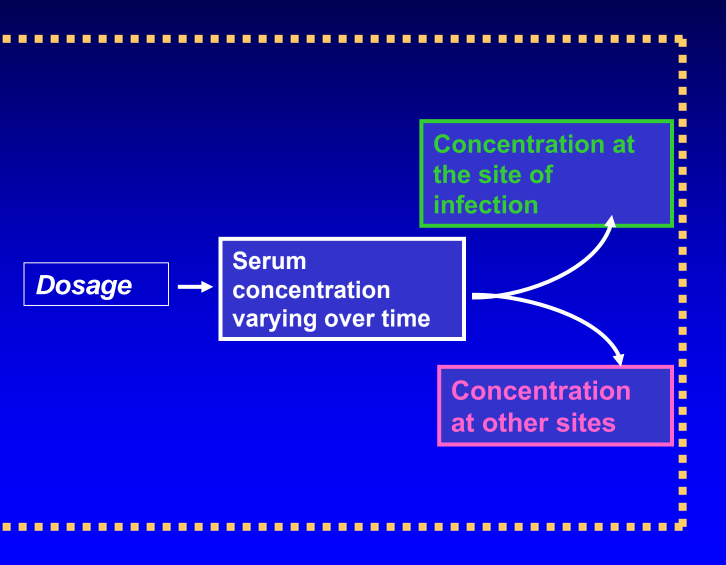


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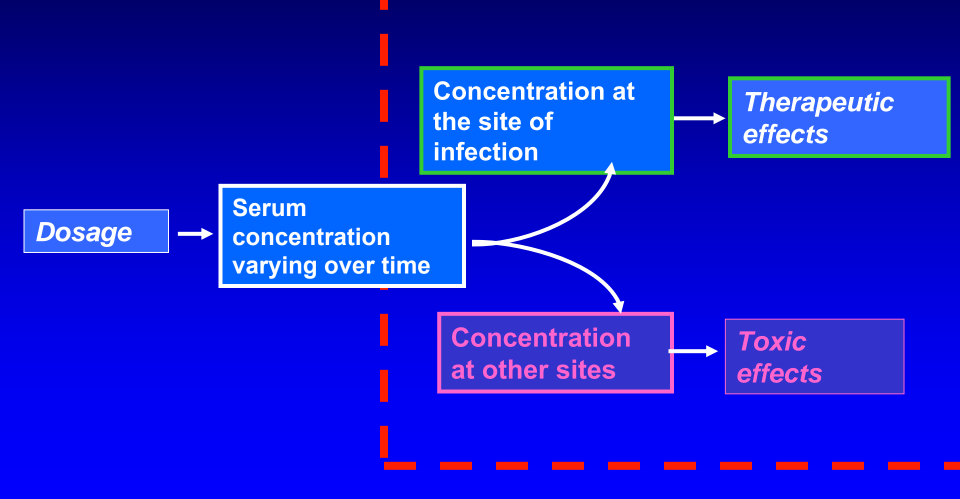


Pharmacokinetics





Pharmacodynamics





PK / PD : why does it improve the use of antibiotics ?

The basics:

- anti-infective drug usage has long been irrational or not scientifically based on a pharmacodynamic point of view
 - search for low doses for fear of toxicity
 - "errors" in drug dosages at registration
 - misunderstanding of "optimal schedules"

 pharmacokinetics was mostly used to establish "drug presence" rather than to correlate dosing with efficacy

pharmacodynamics of antiinfective drugs was largely "*terra incognita*" 20 years ago

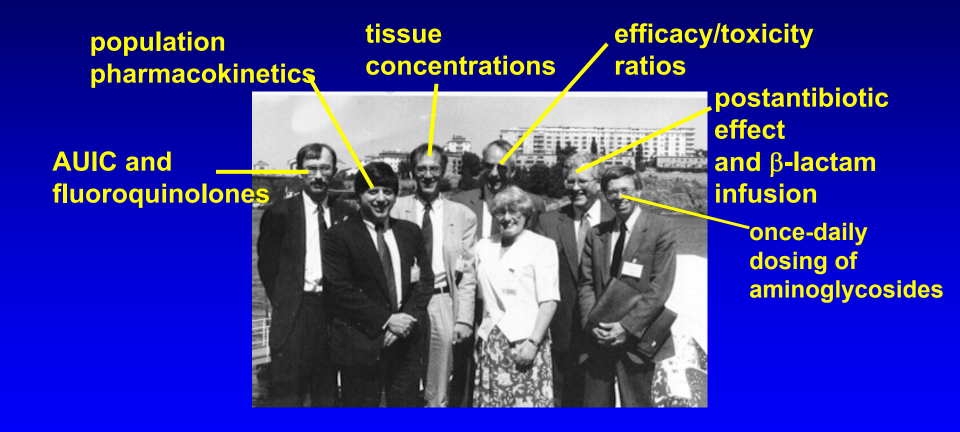


How did it start?

A bunch of goo met in Stockholm in 1989

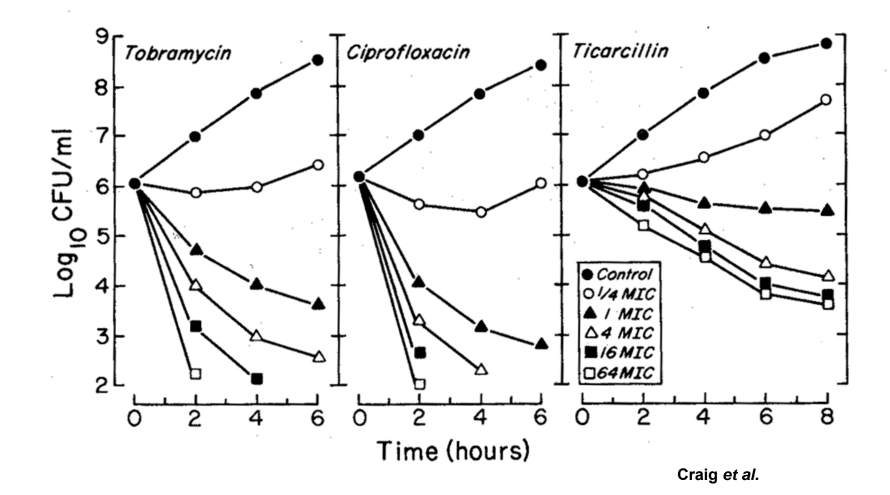


What did they think all about ?





Pharmacodynamics : influence of time and concentration ...





But then came the "Bangalore Meeting"



International Society of Anti-Infective Pharmacology Founded in 1991

7th ISAP International Symposium Advances in the Pharmacology of Antiinfective Therapy co-sponsored with Astra Research Centre, India

January 20-22 (Thursday - Saturday), 2000,

National Science Seminar complex, Indian Institute of Science, Bangalore, Karnataka, India

Craig et al



But then came the "Bangalore Meeting"

http://www.isap.org/2000/Bangalore

International Society of Anti-Infective Pharmacology

- Trends & Problems in Therapy with Antibacterial Drugs (O. Cars, Uppsala, Sweden)
- Pharmacokinetic concepts (O. Cars, Uppsala, Sweden)
- Pharmacodynamic concepts (W.A. Craig, Madison, WI)
- Intracellular pharmacodynamics (P.M. Tulkens, Brussels, Belgium)
- In vitro pharmacodynamic models in the development of antibacterials (E. Löwdin, Uppsala, Sweden)
- Animal models in early evaluation of antibacterial agents (N. Frimødt-Møller, Copenhagen, Denmark)
- Use of animal models to define pharmacokinetic and pharmacodynamic interactions and optimal doses of antibacterial drugs (w.A. Craig, Madison, WI)
- Slide Presentations on In vivo models for the determination of antibacterial efficacy: advantages and limitations (N. Frimødt-Møller, Copenhagen, Denmark & W.A. Craig, Madison, WI)
- Problems in the treatment of tuberculosis (J. Nachega, Baltimore, Md)
- Prediction of clinical efficacy of antitubercular regimens from in vitro studies (C. Paramasivan, Chennai, India)
- <u>Animal efficacy studies of antitubercular agents</u> (V. Balasubramanian, AstraZeneca R&D Indian Site,
- Bangalore, India)
- Evaluation of Clinical Efficacy and Optimal dosing from Clinical Trials (G. Drusano, Albany, NY [talk delivered by W.A. Craig, due to the illness of the speaker])
- Design of Clinical Studies of Antibacterial Agents for Efficacy and Toxicity (R. Norrby, Lund, Sweden)
- <u>Can we obtain useful susceptibility reports ?</u> (Naniwadekar, ...)
- Prediction and prevention of emergence of resistance of clinically used antibacterials (F. Baquero, Madrid, Spain)
- Regulatory efficacy requirements of new antibacterial drugs (talk delivered by W.A. Craig, Madison, WI,

on behalf of the U.S. Food and Drug Administration)

- Industrial Aspects of the development of Antibiotics (J. Edwards, Alderley Park, UK)
- Closing remarks (S. Rosell, Umeå Sweden & P.M. Tulkens, Brussels, Belgium)

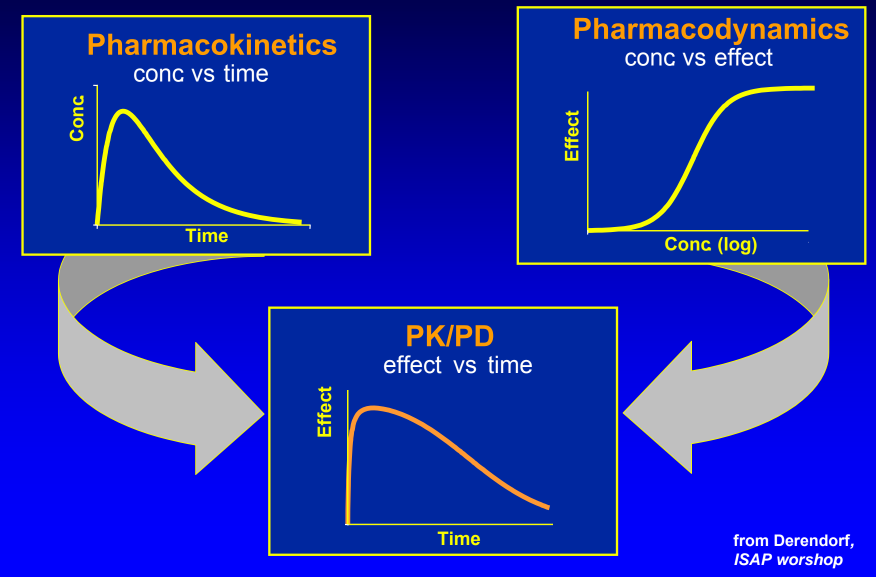
7th ISAP Interr Advances in th co-sponsored with.

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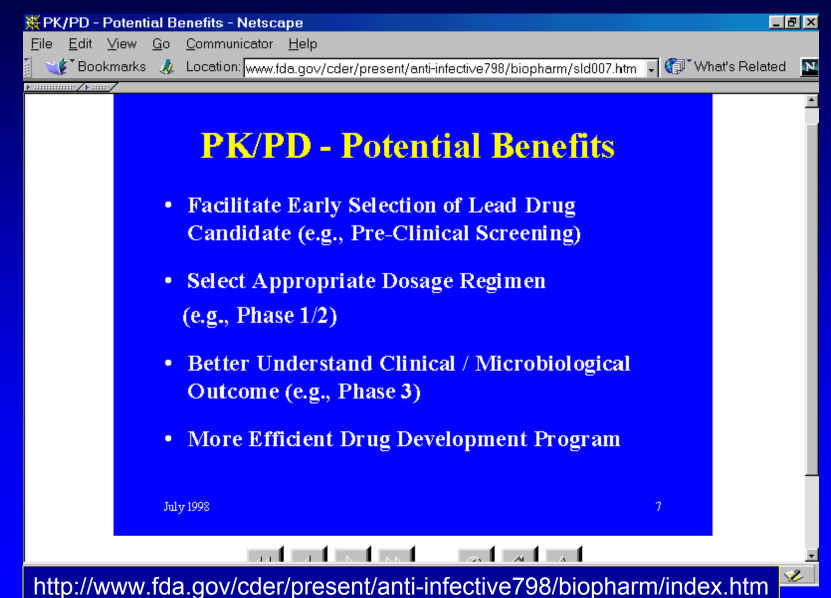
January 20-22 (Th National Science



Pharmacokinetics - Pharmacodynamics







Bangalore, 14 February 2011

More questions ...

 Do you agree with the benefit of "HIT HARD and HIT FAST" ?



1.No 2.Yes

More questions ...

• Do you agree the benefit of "HIT HARD & HIT FAST ?"

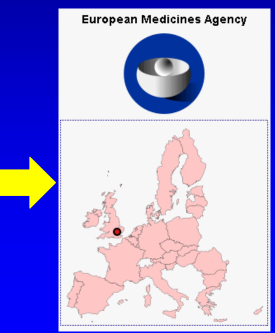


Paul Ehrlich:

,Frapper fort et frapper vite' (Hit hard and early) –

Address to the 17th International Congress of Medicine, 1913

Ehrlich P, Lancet 1913; 2:445–51.





PK /PD and resistance in Europe

" Inadequate dosing of antibiotics is probably an important reason for misuse and subsequent risk of resistance.



A recommendation on proper dosing regimens for different infections would be an important part of a comprehensive strategy.

The possibility of approving a dose recommendation based on pharmacokinetic and pharmacodynamic considerations will be further investigated in one of the CPMP* working parties... "

* Committee for Proprietary Medicinal Products – European Medicines Agency



Publications of the EMA ...





Some achievements:

- once-daily dosing of aminoglycosides registration or reregistration in several countries
 - amikacin, netilmicin (from bid to qd)
 - isepamicin (registered essentially for qd dosing)
- 24h AUC / MIC and C_{max} / MIC ratios used as guides for phase II / III trials, for treatment optimization and for registration of new antimicrobials
 - moxifloxacin
 - telithromycin
- Time above MIC as "gold standard" for β -lactams

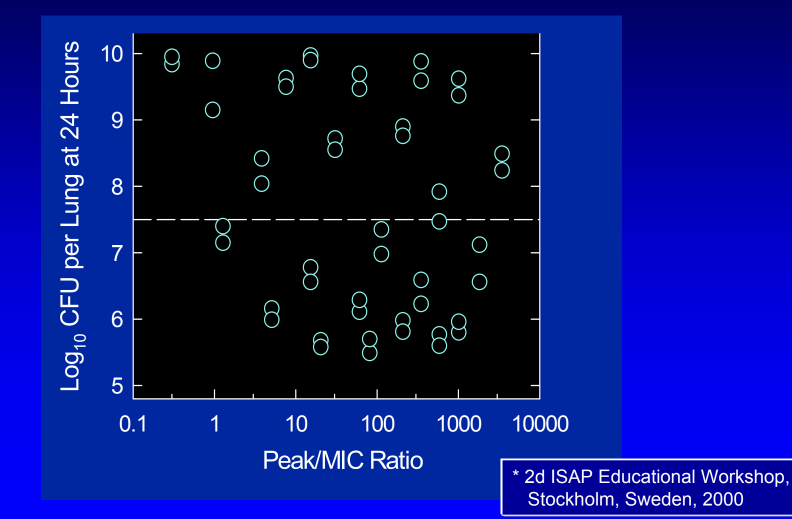


Most available antibiotics can be divided in 3 main groups with respect to PK/PD properties :

- Time-dependent (" T > MIC ")
 - $\rightarrow \beta$ -lactams (all)
- Concentration-dependent (" Cmax / MIC")
 - → aminoglycosides and, for eradication, fluroquinolones
- Total daily dose-dependent (" AUC / MIC ")
 → fluroquinolones (for global efficacy) and all others

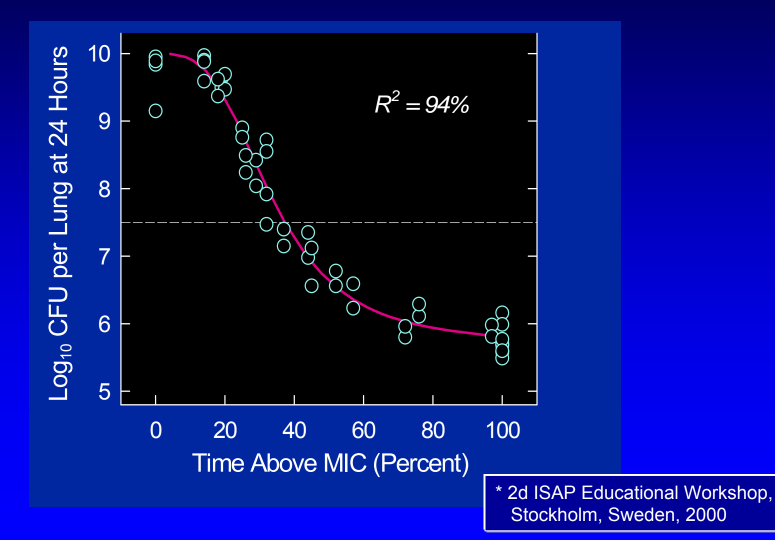


Relationship between peak/MIC and efficacy of cefotaxime towards *Klebsiella pneumoniae* in murine pneumonia (after W.A. Craig *)





Relationship between time above MIC (T>MIC) and efficacy of cefotaxime towards *Klebsiella* pneumoniae in murine pneumonia (after W.A. Craig *)





In a nutshell ... so far ...

- Microbiology parameters: MIC !
- Pharmacodynamic parameters
- PK/PD as applied to beta-lactams: Time-above MIC
- The problems if you underdose
- Take home message



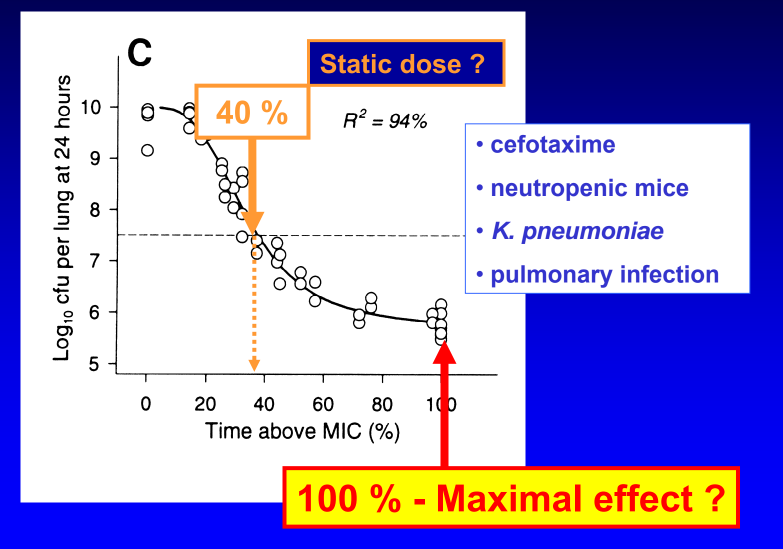
β -lactams : T > MIC ... but ...

You know it is "time above MIC", but...

- How much / How frequent ? (Static dose vs maximum effect ?)
- The same for all beta-lactams ? (Free fractions of the drug (*Fu*) ?)
- The same for all micro-organisms ?
- The same for all infections ?
- Can you apply to all patients ?

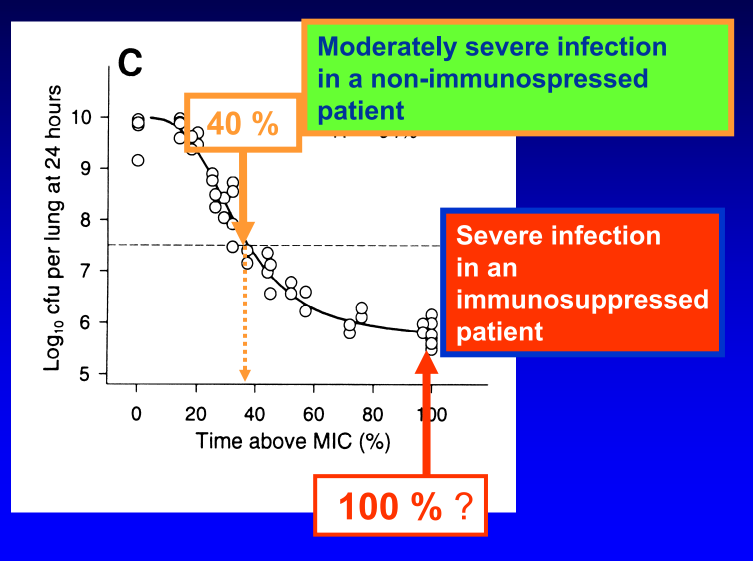


How much time above MIC ?





Here is a proposal ...



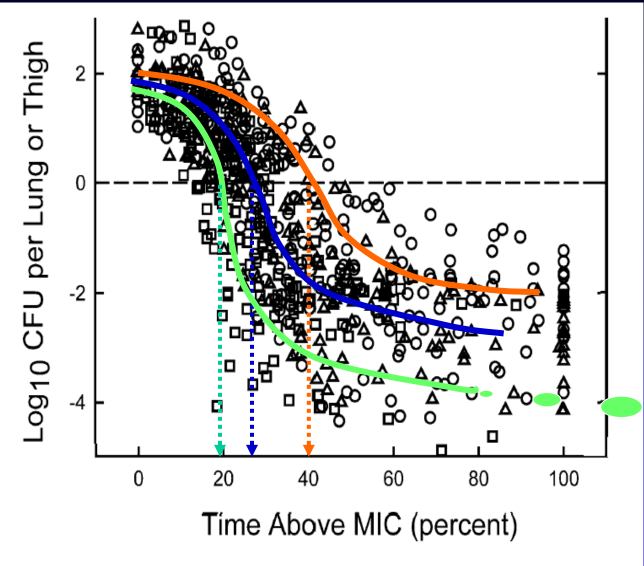


Fig. 7. Relationship between the change in \log_{10} CFU per thigh or lung for various pathogens following 24 h of therapy with different doses of penicillins (\triangle), cephalosporins (\bigcirc), and carbapenems (\Box).

The same for all β-lactams ?

Carbapenems tend to require less time above MIC

Andes & Craig Int. J. Antimicrob. Agents 2002, 19: 261-268



The same for all microorganims ?

T> MIC for static effect

| Drug | Enterobacteriaceae | S. pneumoniae |
|--------------------|--------------------|---------------|
| Ceftriaxone (free) | 38 (34-42) | 39 (37-41) |
| Cefotaxime | 38 (36-40) | 38 (36-40) |
| Ceftazidime | 36 (27-42) | 39 (35-42) |
| Cefpirome | 35 (29-40) | 37 (33-39) |
| Meropenem | 22 (18-28) | |
| Imipenem | 24 (17-28) | |



How do you adjust the dose for a given "Time > MIC" ?

- "out of the package insert" PK data
- Monte-Carlo simulations and target attainment approaches



Typical pharmacokinetics of an IV β-lactam

| time | serum co | serum concentration for | | | | |
|---------|----------|-------------------------|-----|--|--|--|
| (hours) | 0.5 g | 5g 1g | | | | |
| | | | | | | |
| 2 | 25 | 50 | 100 | | | |
| 4 | 12.5 | 25 | 50 | | | |
| 6 | 6 | 12 | 25 | | | |
| 8 | 3 | 6 | 12 | | | |
| 10 | 1.5 | 3 | 6 | | | |
| 12 | 0.75 | 1.5 | 3 | | | |

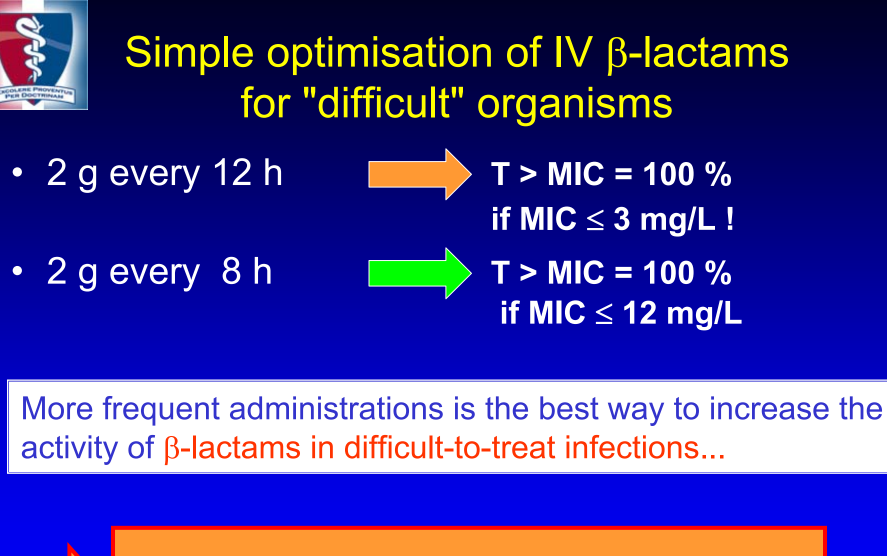
* Single administration unique; half-life 2h ; $V_d = 0.2 I/kg$



Reading the labeling (package insert)

| time | serum co | serum concentration for | | | | |
|---------|----------|-------------------------|----------|---------|--|--|
| (hours) | 0.5 g | 1 g | 2 g | | | |
| 2 | 25 Whe | re would | you like | to be ? | | |
| 4 | 12.5 | 25 | 50 | | | |
| 6 | 6 | 12 | 25 | | | |
| 8 | 3 | 6 | 12 | | | |
| 10 | 1.5 | 3 | 6 | | | |
| 12 | 0.75 | 1.5 | 3 | | | |

* Single administration unique; half-life 2h ; $V_d = 0.2 \text{ l/kg}$



PK / PD breakpoint for

IV β-lactams : MIC = 8 µg/ml

To be practical

In an environment where susceptibilities are compromised (MICs > 4 mg/L) but still "acceptable" (MIC < 16 mg/L) *

- cefepime: 2 g every 8 h
- ceftazidime: 2 g every 8 h
- meropeneme: 2 g every 8 h
- imipeneme: 1 g every 6 h

International labelling (SmPC)

Doses up to 2 g three times daily in adults ...may particularly be suited for treating nosocomial infections due to *Pseudomonas aeruginosa* or *Acinetobacter* spp.

* see discussion about breakpoints later on ...

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In an environment where susceptibilities are compromised (MICs > 4 mg/L) but still "acceptable" (MIC < 16 mg/L) *

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The label of all EU countries limit the dose of imipenem to 4 g/day !

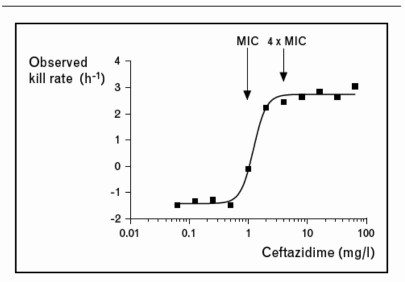
* see discussion about breakpoints later on ...



Target Concentration for β-lactams: continuous infusion

- Maximum effect time-kill at 4 x MIC
- Maximum effect in vitro model 4 x MIC
- Effect in endocarditis model 4 x MIC (Xiong et al 1994)
- Effect in pneumonia model dependent on severity of infection

Figure 2 Relationship between concentration of ceftazidime and kill rate



The relationship follows a Hill-type model with a relatively steep curve; the difference between no effect (growth, here displayed as a negative kill rate) and maximum effect is within two to threefold dilutions. The maximum kill rate is attained at around four times the minimum inhibitory concentration (MIC). Modified with permission from [16].

Mouton JW, Vinks AA. Curr Opin Crit Care. 2007 Oct;13(5):598-606.

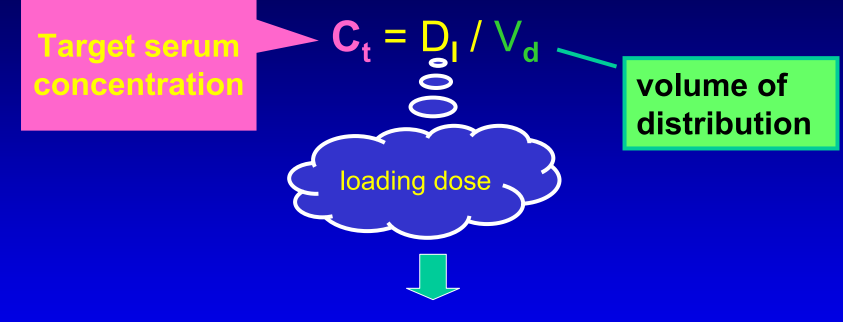


Continuous infusion of β-lactams: an overview...

- The exact role of continuous infusion of β-lactam antibiotics in the treatment of severe infections remains unclear...
- However, increasing evidence is emerging that suggests potential benefits
 - better attainment of pharmacodynamic targets for these drugs
 - More reliable pharmacokinetic parameters in seriously ill patients
 - when the MIC of the pathogen is ≥4 mg/L (empirical therapy where the susceptibility of the pathogen is unknown)
- Clinical data supporting continuous administration are less convincing, but
 - Some studies have shown improved clinical outcomes from continuous infusion
 - none have shown adverse outcomes.
 - clinical and bacteriological advantage are visible in seriously ill patients requiring at least 4 days of antibiotic therapy.
- Seriously ill patients with severe infections requiring significant antibiotic courses (≥4 days) may be the subgroup that will achieve better outcomes with continuous infusion.

Roberts et al., Intern. J. Antimicrob. Agents 30 (2007):11-18

Continuous infusion in practice 1. loading dose: the correct scheme *



loading dose (in mg) = C_t (mg/L) x Vd (L)

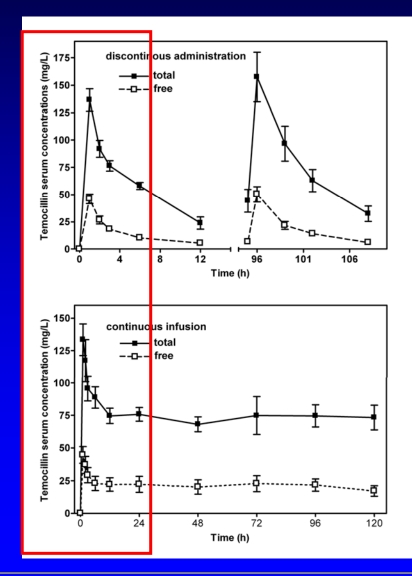
The loading dose is only dependent upon the volume of distribution and is directly influenced by the weight of the patient and his/her medical situation

Typical volumes of distribution of a β -lactam are between 0.2 L/kg (volunteers) and 0.4-0.5 L/kg (Intensive Care and burned patients)

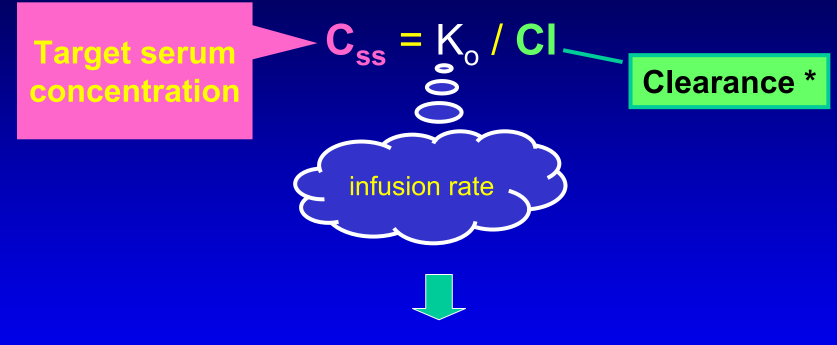
* assuming linear pharmacokinetics (almost always the case for β -lactams)

Continuous infusion in practice 1. loading dose: a simplified (useful) scheme

- Because β-lactams have a low intrinsic toxicity, transient overshooting may not be a major problem...
- Conventional treatments (discontinuous) is by means of bolus or short infusions...
- Why not giving the loading dose as a single bolus or short infusion of a classical dose (1-2 g) ?



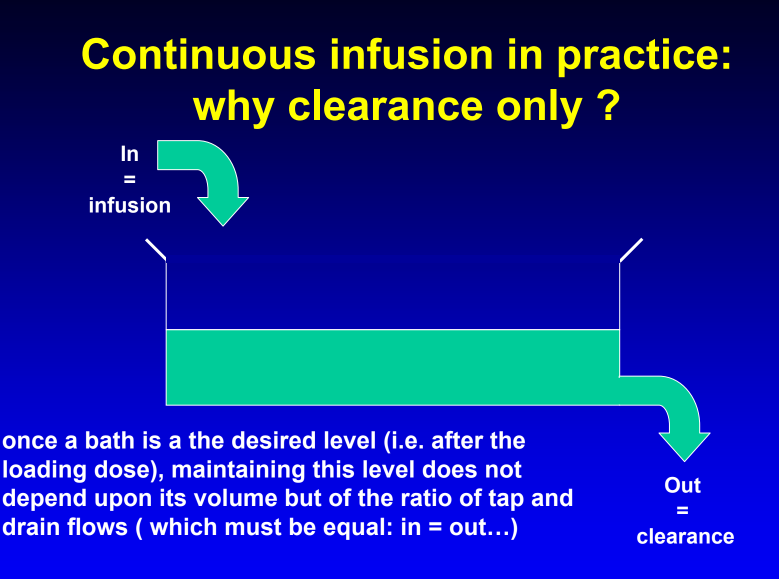
Continuous infusion in practice 2: infusion: the correct scheme *



daily dose (in mg) = 24 x clearance (L/h) x Css

* during the infusion, the necessary dose (in 24h or per min) is only dependent upon the **clearance** and <u>not</u> the weight of the patient

^{*} assuming linear pharmacokinetics (almost always the case for β -lactams)



* during the infusion, the necessary dose (in 24h or per min) is only dependent upon the **clearance** and <u>not</u> the weight of the patient

Continuous infusion of β -lactams: a simplified practical scheme for patientw with normal renal function

Journal of Antimicrobial Chemotherapy (2008) **61**, 382–388 doi:10.1093/jac/dkm467 Advance Access publication 10 December 2007

Continuous versus intermittent infusion of temocillin, a directed spectrum penicillin for intensive care patients with nosocomial pneumonia: stability, compatibility, population pharmacokinetic studies and breakpoint selection

Raf De Jongh¹, Ria Hens¹, Violetta Basma², Johan W. Mouton³, Paul M. Tulkens^{2*}

and Stéphane Carryn²

¹Dienst Voor Intensieve Zorgen, Ziekenhuis Oost-Limburg, B-3600 Genk, Belgium; ²Unité de Pharmacologie Cellulaire et Moléculaire, Université Catholique de Louvain, B-1200 Bruxelles, Belgium; ³Afdeling Medische Microbiologie en Infectieziekten, Canisius Whilhemina Ziekenhuis, NL-6500 GS Nijmegen, The Netherlands

loading dose: 2 g

the conventional unit dose

 infusion: 4 g/day (2.778 mg/min; assumed clearance: 40 ml/min) [drug diluted in 48 ml of water; infusion through motor-operated syringe at a rate of 2 ml/h; temperature 25°C or lower].

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example of

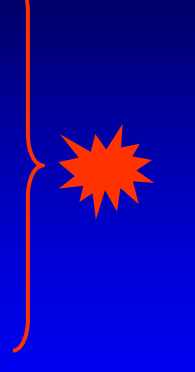
β-lactam

daily dose



Problems with continuous infusion ...

- Clearance estimates
- Variations in clearance (ICU)
- Volume of distribution (ICU, burned patients, ...)
- Non-linear clearance
- drug instability





- Clearance estimates
- Variations in clearance (ICU)
- Volume of distribution (ICU, burned patients, ...)

you may like to monitor the serum levels if MICs \geq 4 (also for discontinuous administration)

Non-linear clearance

drug instability



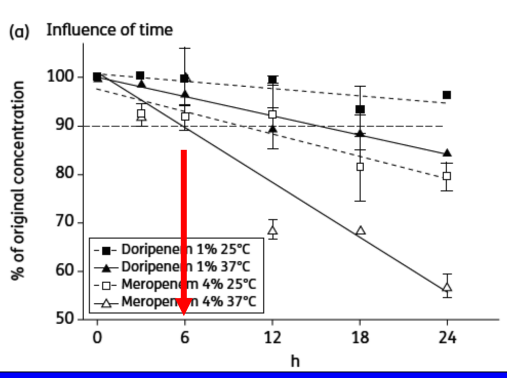


Carbapenems stability

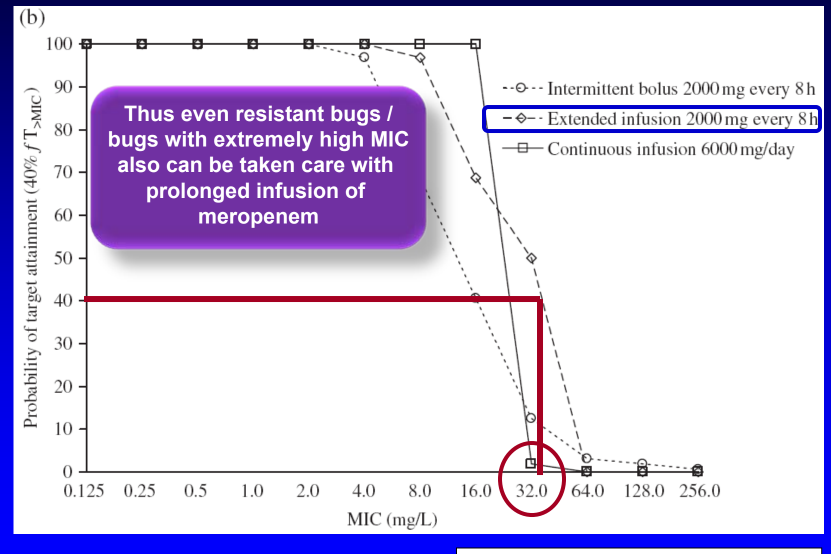
J Antimicrob Chemother (2010) 65:1073-1075 doi:10.1093/jac/dkq044 Advance publication 21 February 2010

Stability of meropenem and doripenem solutions for administration by continuous infusion

Karine Berthoin¹, Cécile S. Le Duff², Jacqueline Marchand-Brynaert², Stéphane Carryn^{1,3} and Paul M. Tulkens^{1*}



Meropenem Infusion in the Critically-III



Roberts et al. J Antimicrob Chemother 2009; 64, 142–150.

Cefepime by continuous infusion

International Journal of Antimicrobial Agents 37 (2011) 46-50



Steady-state pharmacokinetics and pharmacodynamics of cefepime administered by prolonged infusion in hospitalised patients[‡]

S. Christian Cheatham^a, Katherine M. Shea^b, Daniel P. Healy^c, Melissa L. Humphrey^d, Megan R. Fleming^a, Matthew F. Wack^e, David W. Smith^f, Kevin M. Sowinski^d, Michael B. Kays^{d,*}

^a St Francis Hospital, Department of Pharmacy, Beech Grove, IN, USA

^b Seton Family of Hospitals, University Medical Center at Brackenridge, Austin, TX, USA

^c James L. Winkle College of Pharmacy, University of Cincinnati Academic Health Center, Cincinnati, OH, USA

^d Purdue University College of Pharmacy, Department of Pharmacy Practice, W7555 Myers Building, WHS, 1001 West Tenth Street, Indianapolis, IN 46202-2879, USA

e Infectious Diseases of Indiana, Indianapolis, IN, USA

f Clarian Health Partners, Inc., Methodist Hospital, Indianapolis, IN, USA

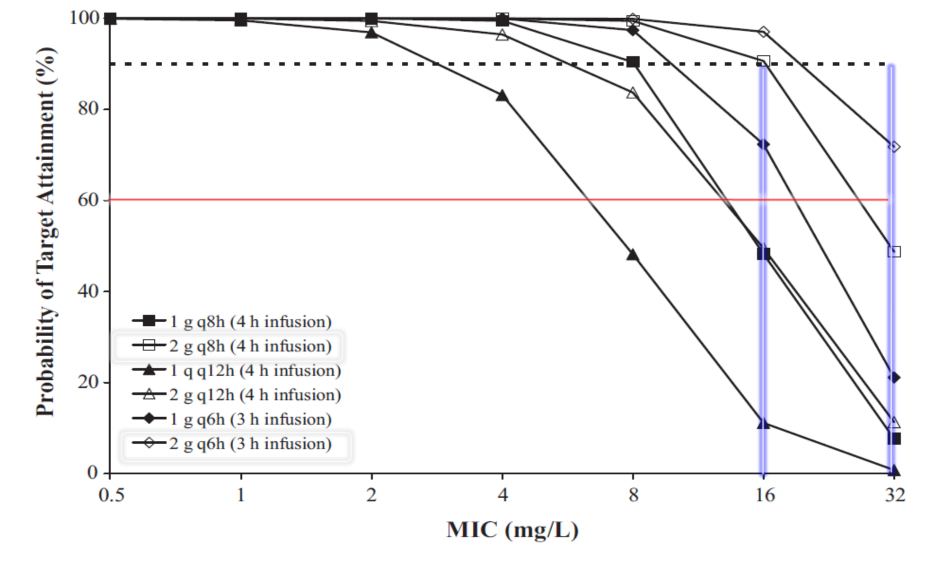


Fig. 2. Probability of target attainment (PTA) at 60% fT > MIC for six prolonged infusion regimens of cefepime at specific minimum inhibitory concentrations (MICs). The dotted line indicates a PTA \ge 90%. fT > MIC, time for which the free drug concentration remains above the MIC; q8h, every 8h; q12h, every 12h; q6h, every 6h.

To be practical : 3 h infusion for patients with normal renal function

- 1st administration: loading dose in 30 min
 2 g (cefepime / meropenem)*
- followed immediately by an 3 h infusion of
 > 2 g (cefepime / meropenem)*
- Repeat step 2 every 8 h

* piperacillin/tazobactam: loading dose: 4.5 g; infusion: 4.5 g every 6 h imipenem: loading dose max. 1 g; infusion: 1 g every 6h (max.)



Breakpoints ...

Clinicians tend to ask only (and clinical microbiologists to provide only) "S – I – R" answers based on accepted breakpoints ...

But, what is a breakpoint?





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☆ - Kr Goog

European Society of Clinical Microbiology and Infectious Diseases

Organization

Clinical breakpoints

💽 🐰

 $\frac{1}{2}$

cpoints

Expert rules

MIC - distributions and QC

Zone diameter distributions

EUCAST disk diffusion test

Frequently Asked Questions (FAQ)

Meetings

EUCAST Presentations

Documents

Information for industry

i Website changes

Links



Clinical breakpoints

| | ™ Recommend page | http://v | www.eucast. | org |
|---|---|----------|-------------|-----|
| | Procedure for harmonizing and defining breakpoints | | | |
| | definitions of clinical breakpoints and epidemiological cut off values | | | |
| | ☑ clinical breakpoints - fungi (MIC breakpoints) | | | |
| | Clinical breakpoints - bacteria (v 1.1) - Excelfile for screen (April 27, 2010) | | | |
| r | – 📙 clinical breakpoints - bacteria (v 1.1) - pdf-file for printing (April 27, 2010) | | | |
| | (technically >1 - 8) mg/L. For a breakpoint listed as S>=22 mm and R<18 mm the intermediate category is 18-21 mm | | | |
| | S-breakpoint and the R-breakpoint. For a breakpoint listed as S<=1 mg/L and R>8 mg/L the intermediate category is 2 - 8 #heterially 4 - 0 | | | |
| | therapy. In EUCAST tables, the I-category is not listed. It is implied as the values between the | | | |
| | Clinical preakpoints are for everyday use in the clinical laporatory to advise on patient | | | |

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| Clinical breakpoints Expert rules MIC - distributions and QC | | search term © Search Clinical breakpoints 💽 |
| Zone diameter distributions | | |
| | ext slides describe the EUCAST proced rmonizing European breakpoints and re | |
| EUCAST Presentatio | rational values. | |
| Documents | (tecnnically ≥1 - o) mg/L. For a breakpoint listed as S>=22 mm and R<18 mm the intermediate category is 18-21 mm. | |
| Information for industry Links | Clinical breakpoints - bacteria (v 1.1) - pdf-file for printing (April 27, 2010) Clinical breakpoints - bacteria (v 1.1) - Excelfile for screen (April 27, 2010) | |
| i Website changes | 년 clinical breakpoints - fungi (MIC breakpoints) | |
| | definitions of clinical breakpoints and epidemiological cut off values procedure for harmonizing and defining breakpoints | |
| | ™ Recommend page | www.eucast.org |

1. Data on dosing, formulations, clinical indications and target organisms are reviewed and differences which might influence breakpoints are highlighted

 Multiple MIC-distributions are collected, the wild type MIC distribution is defined and tentative epidemiological cut-off values determined (WT
 X mg/L)

4. Pharmacokinetic / Pharmacodynamic data are collected and evaluated; . Monte Carlo simulations are performed and a PK/PD breakpoint calculated based on conventional dosing regimens

 Clinical data relating outcome to MIC-values, wildtype and resistance mechanisms are assessed in relation to the tentative breakpoint

6. Pk/Pd breakpoints are checked against target species wild type MIC distributions to avoid splitting the wild type population

http://www.eucast.org

7. Tentative breakpoints by the EUCAST Steering Committee are referred to the national breakpoint committees for comments.

When steering committee and national committees agree the tentative breakpoints are subjected to the EUCAST consultation process:

- 8. Consultation process on tentative breakpoints:
 - EUCAST general committee
 - Expert committees (Neisseria, Anaerobes, others)
 - pharmaceutical industry, AST device manufacturer
 - others via EUCAST website

9. Rationale document prepared and published on website

http://www.eucast.org



European Society of Clinical Microbiology and Infectious Diseases

EUCAST and carbapenems

Enterobacteriaceae

| MIC breakpoint (mg/L) | | | Zone diameter breakpoint (mm) | |
|--------------------------|-----------------------------|--|---|---|
| S≤ | R > | (µg) | S≥ | R < |
| 1 | 4 | 10 | 24 | 18 |
| 0.5 | 1 | 10 | 25 | 22 |
| 2 | 8 | 10 | 21 | 15 |
| 2 | 8 | 10 | 22 | 16 |
| | (mg S ≤ 1 0.5 2 | (mg/L) S ≤ R > 1 4 0.5 1 2 8 | (mg/L) content (μg) S ≤ R > 1 4 0.5 1 2 8 | (mg/L) content (µg) breakpo S ≤ R > 24 1 4 10 24 0.5 1 10 25 2 8 10 21 |

- The carbapenem breakpoints for Enterobacteriaceae will detect all clinically important resistance mechanisms (including the majority of carbapenemases).
- Some strains that produce carbapenemase are categorized as susceptible with these breakpoints and should be reported as tested, i.e. the presence or absence of a carbapenemase does not in itself influence the categorization of susceptibility.
- In many areas, carbapenemase detection and characterization is recommended or mandatory for infection control purposes.



European Society of Clinical Microbiology and Infectious Diseases

EUCAST and cephalosporins

| Cephalosporins ¹ | MIC breakpoint (mg/L) | | Disk content (µg) | Zone diameter breakpoint (mm) | |
|-----------------------------|--------------------------|-----|-------------------------|----------------------------------|-----|
| | S≤ | R > | | S≥ | R < |
| Cefepime | 1 | 4 | 30 | 24 | 21 |
| Ceftazidime | 1 | 4 | 10 | 21 | 18 |
| Ceftriaxone | 1 | 2 | 30 | 23 | 20 |
| | • • | | | | |

EUCAST_breakpoints_v1.1.pdf

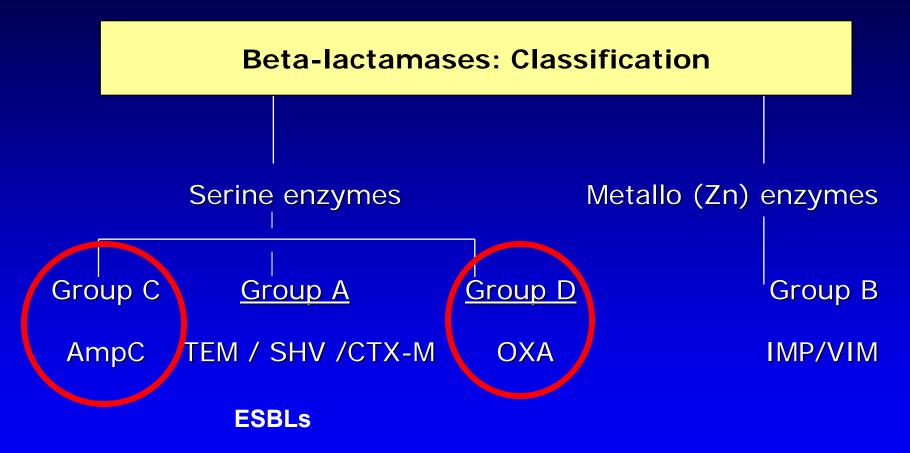
Why so low ?

1. The cephalosporin breakpoints for Enterobacteriaceae will detect all clinically important resistance mechanisms (including ESBL, plasmid mediated AmpC). Some strains that produce beta-lactamases are susceptible or intermediate to 3rd or 4th generation cephalosporins with these breakpoints and should be reported as found, i.e. the presence or absence of an ESBL does not in itself influence the categorization of susceptibility. In many areas, ESBL detection and characterization is recommended or mandatory for infection control purposes.





What about ESBL ?





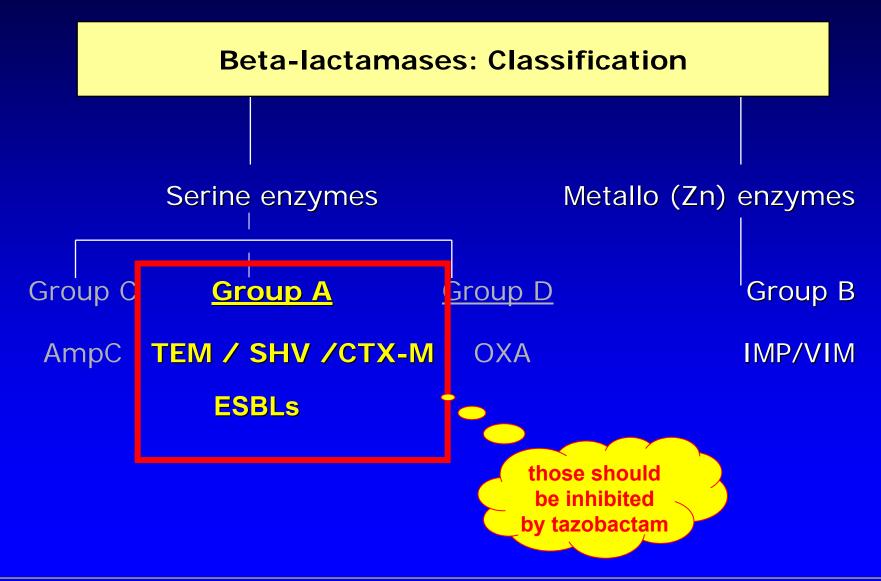
Class A and D of β-lactamases are poorly active on 3d generation cephalosporins

| Table 130.1 Functional classification of β-lactamases | | | | | | | |
|--|-----------------|---|---|--|--|--|--|
| Group | Molecular class | Preferred substrates | Preferred substrates Active β-lactams | | | | |
| Group 1: serine cephalosporinases not inhibited by clavulanic acid | с | Cephalosporins I and II (>> cephalosporins III, monobactams, penicillins) | Carbapenems Temocillin cephalosporins III and IV; variable upon level of expression) | AmpC from gram-negatives; variable upon the species | | | |
| 2d: cloxacillin-hydrolyzing β-lactamases generally inhibited by clavulanic acid | D | Penicillins Cloxacillin Cephalosporins I and II | Carbapenems Cephalosporin III Monobactams* Piperacillin + tazobactam | OXA-1 to -30, PSE-2 from Enterobacteriaceae and <i>P. aeruginosa</i> OXA-11 to -19, 28, 32, 45 are ESBLs in <i>P. aeruginosa</i> (R to Ceph 3) OXA-23, -24, -58 are carbapenemases in <i>Acinetobacter baumannii</i> | | | |



Van Bambeke F, Glupczynski, Y, Mingeot-Leclercq, MP, Tulkens PM Mechanisms of Action. In: Infectious Diseases (3d edition; J. Cohen, W. Powderly & S. Opal, eds), chapter 130, pp 1288-1307, Elsevier/Mosby, 2010

So, now you are left with the ESBL...





An innovative approach for ESBL...

- take a 4th generation cephalosporin (cefepime [PM])
 - should cover (partly AmpC) and resist to OXA
- add a β -lactamase inhibitor (tazobactam [TZ])
 - → will take care of many ESBL

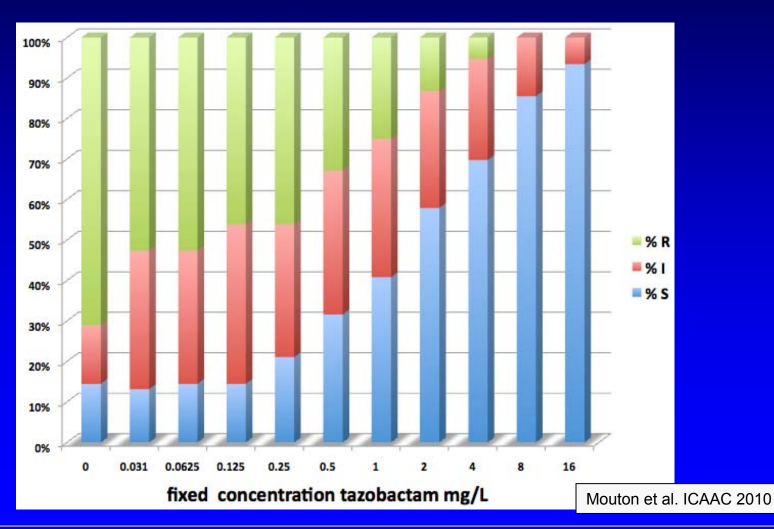
Mouton et al. ICAAC 2010 76 ESBL producing *Enterobacteriaceae* were selected from a variety of clinical specimens.

| | %S | %I | %R | MIC50 (mg/L) | MIC90 (mg/L) |
|--------------|----|----|----|-----------------|-----------------|
| PM | 15 | 14 | 71 | >32 | >32 |
| PM/TZ 1mg/L | 41 | 34 | 25 | 2 | 32 |
| PM/TZ 4mg/L | 70 | 25 | 5 | 0.5 | 4 |
| PM/TZ 16mg/L | 93 | 7 | 0 | 0,25 | 1 |



An innovative approach for ESBL...

Percentage sensitive(S), intermediate(I) and resistant(R) cefepime (breakpoints EUCAST: ≤1 S – R >8)





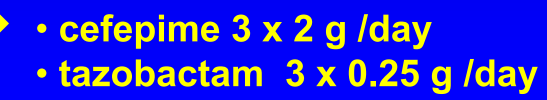
An innovative approach for ESBL...

Conclusions:

- The combination of cefepime and tazobactam may offer an alternative treatment option for ESBL harboring strains.
- If the same amount of tazobactam is used as current piperacillin/tazobactam regimens and breakpoint determinations, most strains would be categorized as susceptible.

Mouton et al. ICAAC 2010

In India, due to high ESBL: consider cefepime + tazobactam







In a nutshell ... so far ...

- Microbiology parameters: MIC !
- Pharmacodynamic parameters
- PK/PD as applied to beta-lactams: Time-above MIC
- The problems if you underdose
- Take home message



A simple experiment ...

Exposure of E. aerogenes to anrti-Gram (-) penicillin (temocillin) to 0.25 MIC for 14 days with daily readjustment of the concentration based on MIC détermination

| | Initial MIC (mg/L) ^a | | | TEM-exposed MIC (mg/L) | | | Revertant MIC (mg/L) | | |
|----------------------|------------------------------------|-----|-------|---------------------------|-------|-------|-------------------------|-----|-------|
| strains | | | | | | | | | |
| | ТЕМ | FEP | MEM | TEM | FEP | МЕМ | TEM | FEP | MEM |
| 2114/2 ° | 8 | 2 | 0.25 | 2048 | > 128 | 16 | 32 | 4 | 0.5 |
| 2502/4 ^c | 8 | 2 | 0.125 | 8192 | 4 | 0.25 | 4096 | 1 | 0.125 |
| 3511/1 ° | 32 | 2 | 0.125 | 4096 | 32 | 0.125 | 4096 | 8 | 0.5 |
| 7102/10 ^d | 512 | 32 | 1 | 16384 | > 128 | 4 e | 8192 | 64 | 1 |

^a figures in bold indicate values > the R breakpoint for Enterobacteriaceae (EUCAST for MEM [8] and FEP [4]; BSAC and Belgium for TEM [16])

^b dotblot applied with antiOmp36 antibody; signal quantified for grey value after subtraction of the signal of a porin-negative strain (ImageJ software); negative values indicate a signal lower than the background

^c ESBL TEM 24 (+) ; ^d ESBL (-) and AmpC (+) [high level] ; ^e Intermediate (I) according to EUCAST



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| strains | | | | | | | MIC (mg/L) | | |
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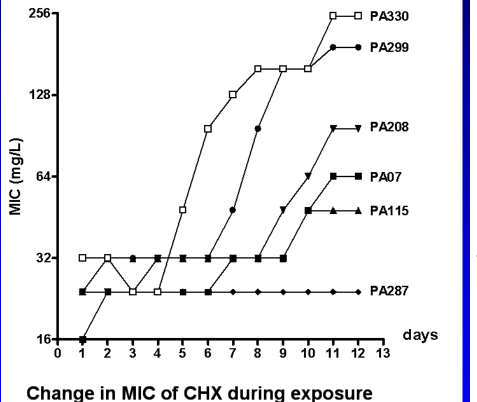
° ESBL TEM 24 (+) ; d ESBL (-) and AmpC (+) [high level] ; e Intermediate (I) according to EUCAST





And this happens also with biocides

Exposure of *P. aeruginosa* to sub-MIC concentrations of chlorhexidine



to 0.5 MIC with daily concentration readjustment

 initial
 after 13 days exposure to CHX

 Search
 Image: Search

 Image: Sea

Typical change in colony size and swarming abilities after 13 days of exposure to 0.5 MIC

Tan et al. ECCMID 2011, in press



And in the clinics ?

International Journal of Antimicrobial Agents 36 (2010) 513-522



Contents lists available at ScienceDirect

International Journal of Antimicrobial Agents

journal homepage: http://www.elsevier.com/locate/ijantimicag

In vivo development of antimicrobial resistance in *Pseudomonas aeruginosa* strains isolated from the lower respiratory tract of Intensive Care Unit patients with nosocomial pneumonia and receiving antipseudomonal therapy

Mickaël Riou^{a,1}, Sylviane Carbonnelle^{a,2}, Laëtitia Avrain^{a,b}, Narcisa Mesaros^{a,3}, Jean-Paul Pirnay^c, Florence Bilocq^c, Daniel De Vos^{c,d}, Anne Simon^e, Denis Piérard^f, Frédérique Jacobs^g, Anne Dediste^h, Paul M. Tulkens^{a,*}, Françoise Van Bambeke^a, Youri Glupczynskiⁱ

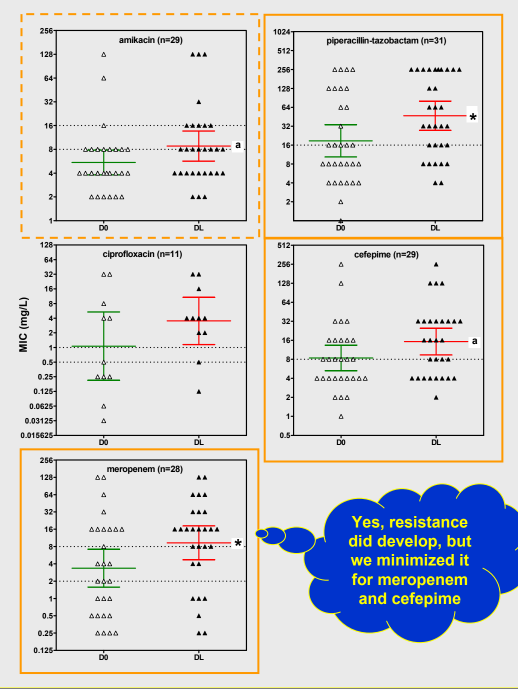
Antimicrobial



What happens during treatment ?

- D0: initial isolate
 DL: last isolate obtained
- individual values with geometric mean (95 % CI)
- S (lowest line) and R (highest line) EUCAST breakpoints
- p < 0.05 by paired t-test (twotailed) and Wilcoxon nonparametric test
- a p < 0.05 by Wilcoxon nonparametric test only

Note: stratification by time between D0 and DL gave no clue (too low numbers)

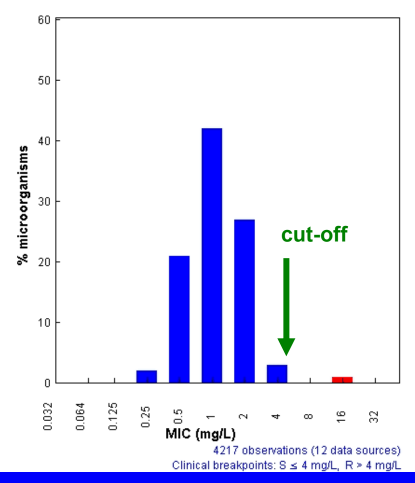


Bangalore, 14 February 2011

And what about colistin ?

You first need to consider the MIC distribution.

Here are the data of EUCAST for *Pseudomonas* Colistin / Pseudomonas aeruginosa EUCAST MIC Distribution - Reference Database 2011-02-12



And what about colistin ?

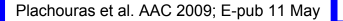
Dosage (colistine methane sulfonate [CMS]): 240 mg every 8h (= 3 x 10⁶ UI)

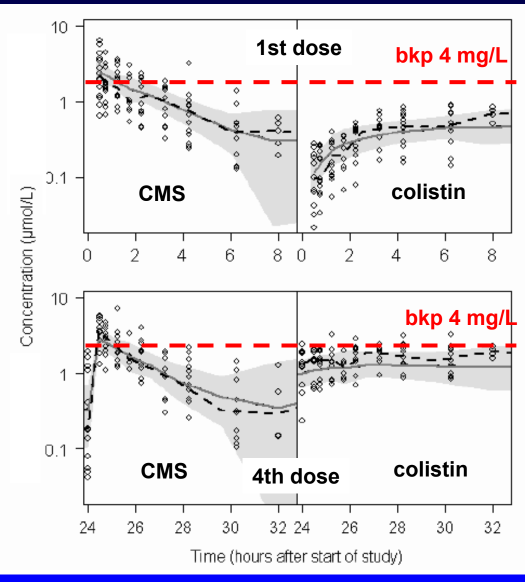
CMS • t_{1/2} ~ 2.3 h,

Colistin:

- t_{1/2} ~ 14.4 h.
- Cmax (pred.)
 - •1st dose: 0.60 mg/L
 - s.s.: 2.3 mg/L.

Problem #1: Low initial blood levels suggest the necessity of a loading dose

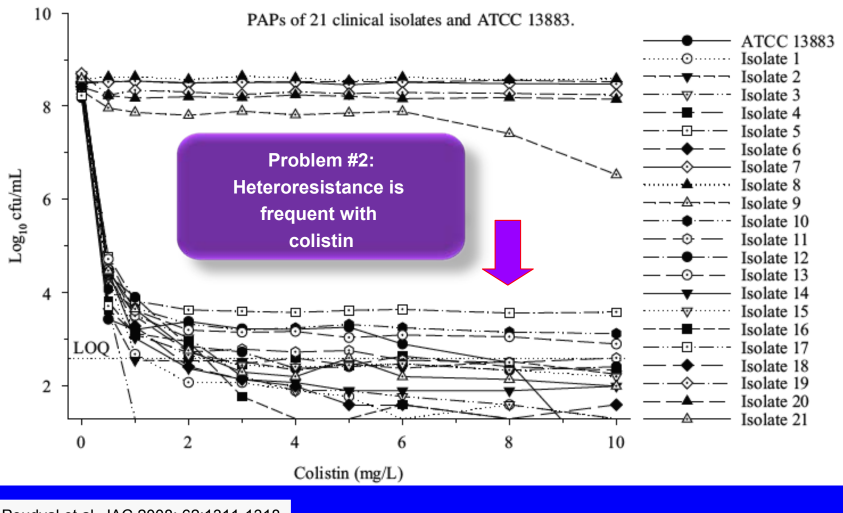




Bangalore, 14 February 2011

And what about colistin ?

Population analysis profiles of K. pneumoniae isolates



Poudyal et al. JAC 2008; 62:1311-1318



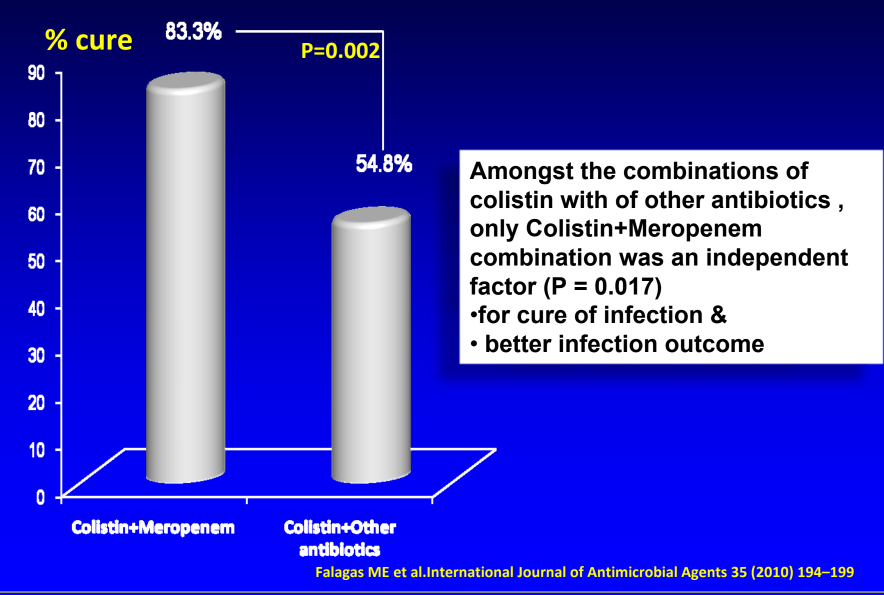
Short communication

Colistin therapy for microbiologically documented multidrug-resistant Gram-negative bacterial infections: a retrospective cohort study of 258 patients

Matthew E. Falagas^{a,b,c,*}, Petros I. Rafailidis^{a,b}, Elda Ioannidou^a, Vangelis G. Alexiou^a, Dimitrios K. Matthaiou^a, Drosos E. Karageorgopoulos^a, Anastasios Kapaskelis^{a,b}, Dimitra Nikita^d, Argyris Michalopoulos^{a,e}

- Retrospective cohort <u>clinical study of 258 patients</u>
- <u>52.3% isolates were polymyxin–only-susceptible</u>
- Remainder were susceptible to colistin & at least 1 other antibiotic

Patients with polymyxin-only-susceptible infections



 In how many patients you are implementing "once-daily dosing" of aminoglycosides?

0%
 25%
 50%
 75%
 100%



Journal of Antimicrobial Chemotherapy (1991) 27, Suppl. C, 49-61

Pharmacokinetic and toxicological evaluation of a once-daily regimen versus conventional schedules of netilmicin and amikacin

Paul M. Tulkens

Laboratoire de Chimie Physiologique, Université Catholique de Louvain, and International Institute of Cellular and Molecular Pathology, Brussels, Belgium

In conclusion, these very sensitive tests of nephro- and oto-toxicity suggest that od dosing of amikacin or netilmicin is, if anything, safer than bd or tid dosing.



"Take home" message

- dosage is key to success and protection against resistance...
- dosage should match bacterial susceptibility... and knowledge of MIC is essential
- for β-lactams, get TIME > MIC to reach maximal efficacy ... and dose appropriately...
 → 3h infusion of meropenem and cefepime may help
- Use of correct breakpoints will also help in avoiding the use of "weak antibiotics" ... or to decide dosage escalation to avoid emergence of resistance ...
- New combinations tailored to local needs (viz. cefepime + tazobactam) with 3h infusion) are useful ...



WHO statement 2000

The most effective strategy against antibiotic resistance is:

- "to unequivocally destroy microbes"
- "thereby defeating resistance before it starts"

WHO Overcoming Antimicrobial Resistance, 2000



And a few sights from Belgium...







Brussels







I hope the future will be fine with you...



http://www.isap.org





http://www.facm.ucl.ac.be

All slides are available from here