





**GSK-Chair of Infectious Diseases** (Chaire GSK de Maladies Infectieuses / GSK-Leerstoel in Infectieziekten)

a joint academic activity of the Université catholique de Louvain and the Katholieke Universiteit Leuven

#### How we try to improve efficacy, decrease resistance and save money with antibiotics in hospitals ...

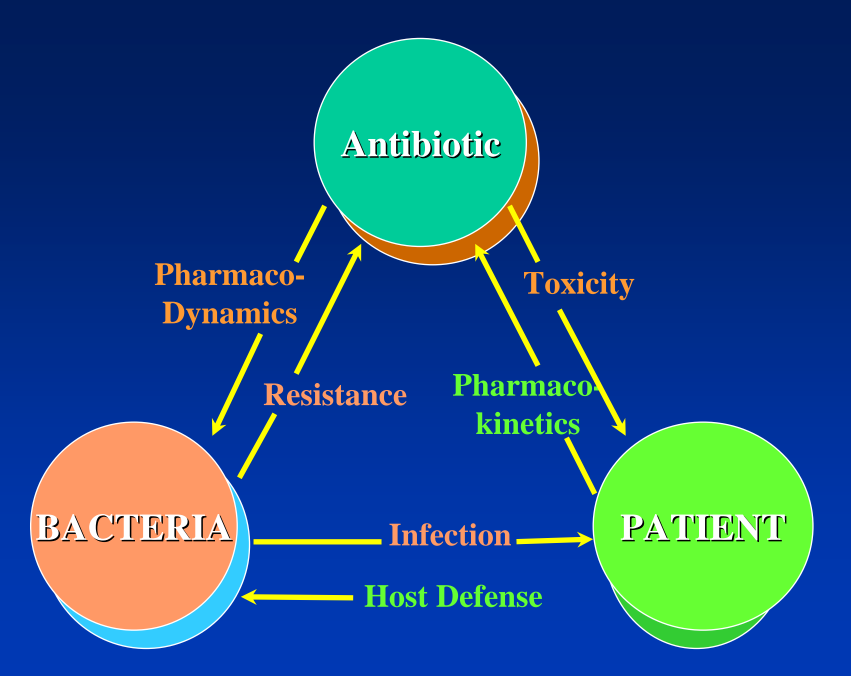
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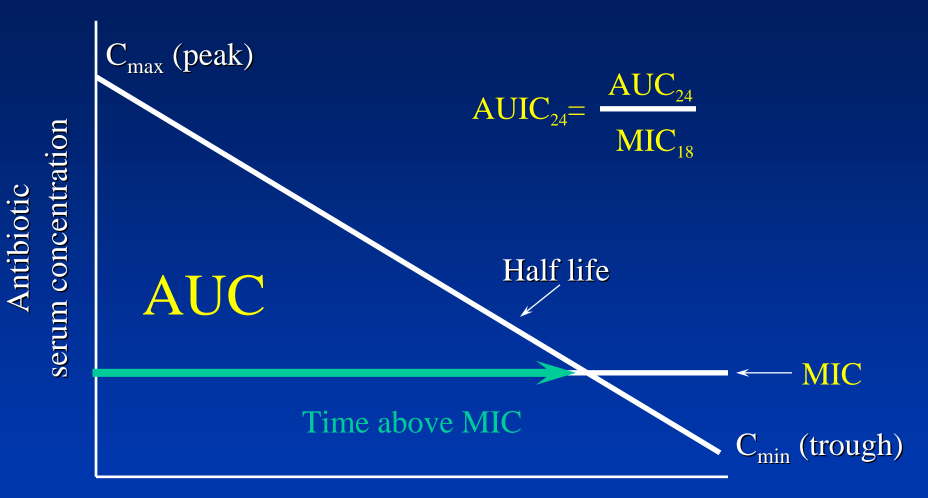
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Applied Pharmacokinetics and Pharmacodynamics: 4th Edition, 2006



Time

# Model Antibiotics for Human PK/PD trials:

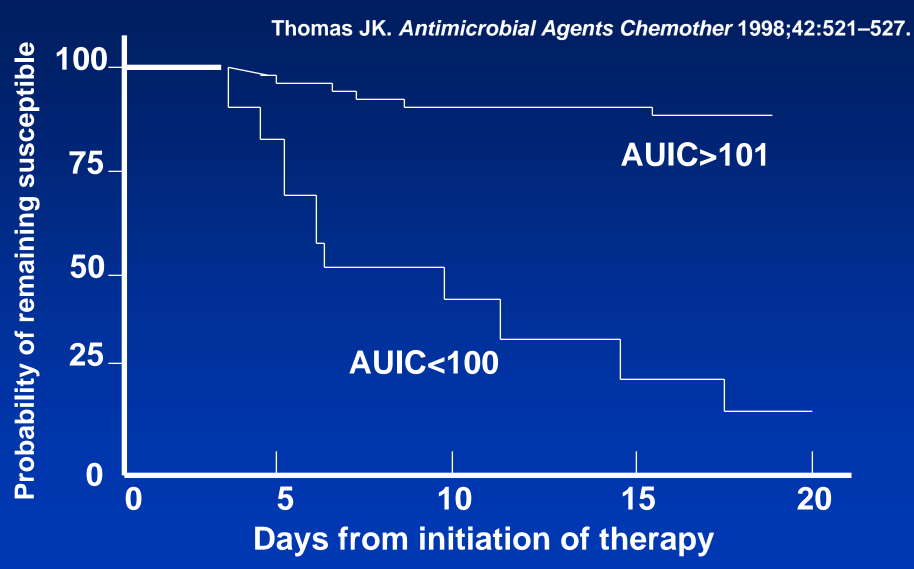
- Ciprofloxacin
- Grepafloxacin
- Tobramycin
- Piperacillin
- Ceftazidime
- Azithromycin
- Linezolid

- Cefmenoxime
- Cefepime
- Aztreonam
- Synercid
- Imipenem
- Telithromycin
- Vancomycin

## Advantages of Antibiotics

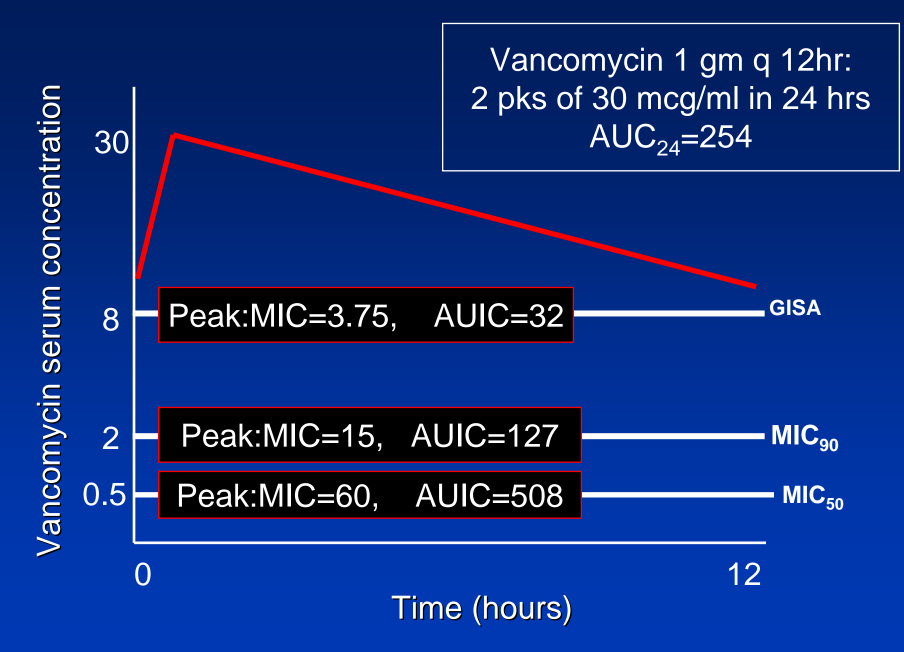
- You can readily isolate, grow and study the "receptor" for an antibiotic
  - Fortunate, because susceptibility varies tremendously between "receptors"
- Correlations between in vivo Pharmacokinetics and in vivo Pharmacodynamics are feasible; This also includes Resistance

#### **AUIC vs Resistance**



# Vancomycin – Role in Therapy

- Up until ~ 1990, it was the undisputed drug of choice for gram positives such as staphylococci and enterococci, and was always perceived as effective.
- Purpose of Serum Conc. Monitoring was to avoid toxicity; rigid range of concentration defined as peak ~ 30 mcg/ml and trough ~ 10 mcg/ml
- Problems followed increasing use
  - 1991: E. faecium became VREF
  - 1995-1998: Arrival of VISA and Declining success vs. MRSA, even when "susceptible"



Schentag JJ. Critical Care Med 29 (4 Suppl): N100-N107, 2001

#### Vancomycin: AUICs vs Time > MIC?

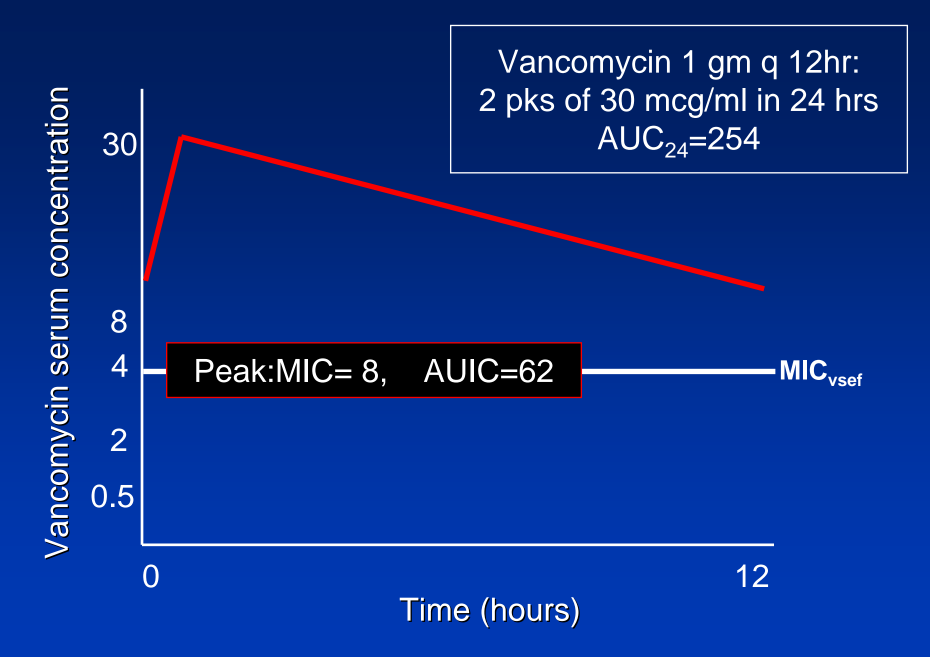
- Vancomycin is slowly cidal, and demonstrates time-dependent killing and a long half-life. With these properties, there ought to be good correlation between AUIC and time above MIC
- Dosing controls the blood levels in most patients, so any low AUICs or short time>MICs would be the result of high organism MICs
- Correlation study in 84 patients at MFH
  - This was 1993, and the goal was to understand VREF development in bacteremia patients...

## Vancomycin Outcomes vs AUICs

|                     | Outcome      |                |               |  |  |
|---------------------|--------------|----------------|---------------|--|--|
|                     | Satisfactory | Unsatisfactory | Indeterminate |  |  |
| MIC >1.0 µg/ml      | 1            | <b>4</b> a     | 0             |  |  |
| MIC <1.0 µg/ml      | 74           | 2              | 3             |  |  |
| AUIC <125           | 4            | 4 <sup>b</sup> | 0             |  |  |
| AUIC >125 (76)      | 71           | 2              | 3             |  |  |
| Total Patients (84) | 75           | 6              | 3             |  |  |

<sup>a</sup> *p* < 0.001 <sup>b</sup> *p* < 0.005

Hyatt, et al. Clin Pharmacokinet. 1995;28:143-160.



Schentag JJ. Critical Care Med 29 (4 Suppl): N100-N107, 2001

#### **Enterococcus faecium (VSEF-VREF)**

- Dangers of the inadvertent high MIC organism like *E. faecium*, with a fixed-AUC drug like vancomycin
- We lost the use of vancomycin for *E. faecium* by 1998
- Large increases in vancomycin dosing could have delayed this loss.
  - Target AUIC is 125 for VSEF (Hyatt et al. Clin PK 1995;28:143)
    - Double the dose (AUC<sub>24</sub> ~ 500) for MIC=4.0
    - Quadruple the dose (AUC<sub>24</sub> ~ 1000) for MIC=8

Peaks of ~150, troughs of 110....

- Alternatives for MIC > 8.0 mcg/ml:
  - Quinupristin/Dalfopristin (September 20, 1999)
  - Linezolid (April 18, 2000)

#### .....What about S. aureus, esp. MRSA? 12

### MRSA: Issues With "Appropriately Dosed" Vancomycin?

- MRSA MICs are usually 0.5 to 1.0 mcg/ml
  - Slow killing of organisms in vitro and in vivo
- MRSA MBCs are increasingly 4-32 mcg/ml
  - Staphylococci that are not yet VISA or VRSA, but no longer responding to vancomycin at AUICs of 125-250
- Clinical Evidence of Problems with Vancomycin?; Failures even before VISAs with MICs ~ 2-4 mcg/ml

## Patient 2 and Patient 3

- Patient 2
  - 78-year-old male
  - Developed MRSA pneumonia day 107, treated with vancomycin
  - Initial infection  $\times$  15 days
  - 2nd infection  $\times$  15 days
  - 3rd infection  $\times$  8 days
  - 4th infection  $\times$  7 days
    - MRSA now colonized
  - Vanco MIC≤0.5

- Patient 3
  - 71-year-old female
  - Admitted from NH with MRSA pneumonia, treated with vancomycin
  - Initial infection × 10 days
  - 2nd infection  $\times$  5 days
  - Patient expired, day 20
    - MRSA not eradicated
  - Vanco MIC≤0.5

## Patient 2: Healthcare Resources Used

| Event                     | Vanco<br>AUIC <sup>a</sup> | Vanco<br>Ievels <sup>b</sup> | Total (\$) | \$/day |
|---------------------------|----------------------------|------------------------------|------------|--------|
| Initial infection         |                            |                              |            |        |
| (1/13/98-1/27/98)         | 394                        | Rdm >20                      | 29,055     | 1,937  |
| 2 <sup>nd</sup> infection |                            |                              |            |        |
| (2/2/98-2/16/98)          | 195                        | ND                           | 38,588     | 2,573  |
| 3 <sup>rd</sup> infection |                            |                              |            |        |
| (2/21/98-2/28/98)         | 266                        | ND                           | 16,385     | 2,048  |
| 4 <sup>th</sup> infection |                            |                              |            |        |
| (3/18/98-3/24/98)         | 736                        | Tr>20                        | 23,375     | 3,339  |

<sup>a</sup> Values expressed are means.
<sup>b</sup> Rdm = random; ND = not done; Tr = trough.

# Why Is Vancomycin Failing?

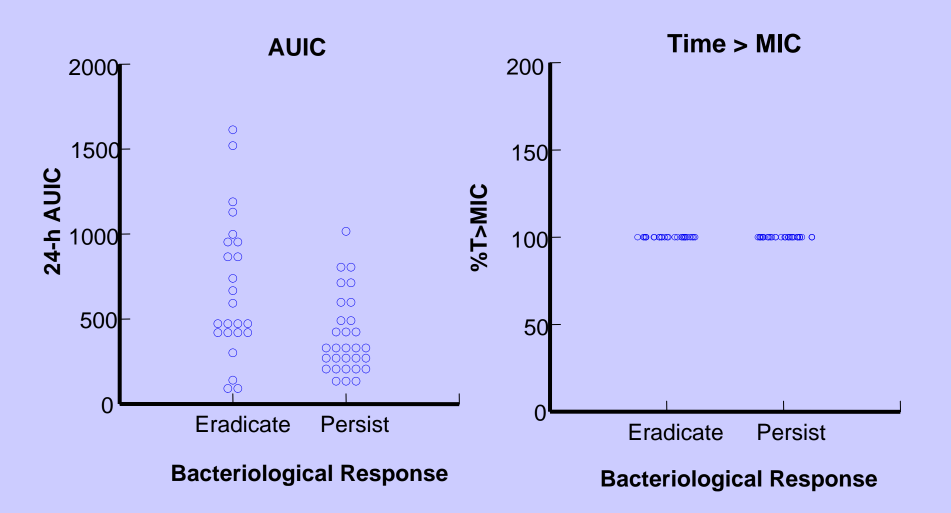
- Slowly or poorly cidal, hetero-resistance?
- MBC >> MIC for these vancomycin exposed organisms?
- Increasingly larger fractions of the organism population reach the definition of tolerance
- Vancomycin PK/PD target of 125 is too low for this drug; For MRSA, we may need AUICs of 400 or even more?

# **PK/PD study in S. aureus LRTI**

- 108 patients in 1998 that qualified for PK/PD and LRTI out of a total of 160 pts at MFH that year (Mean Age=74, 67% on Ventilator at baseline); Main reason for exclusion was insufficient proof of LRTI
- All patients had PK/PD as AUIC<sub>24</sub>; for endpoints we could often derive time to bacterial eradication (via daily cultures) and time to clinical cure (via daily scoring). We also collected the usual cure-failure micro and clinical data typical of registration trials.
- Clinical success was 59% overall; 54% for MRSA, 71% for MSSA
  - Oxacillin vs MSSA was 100% effective
  - Failure overall was associated (LR analysis) with MRSA, low albumin, low CCr, multi-lobe involvement and AUIC <400</li>

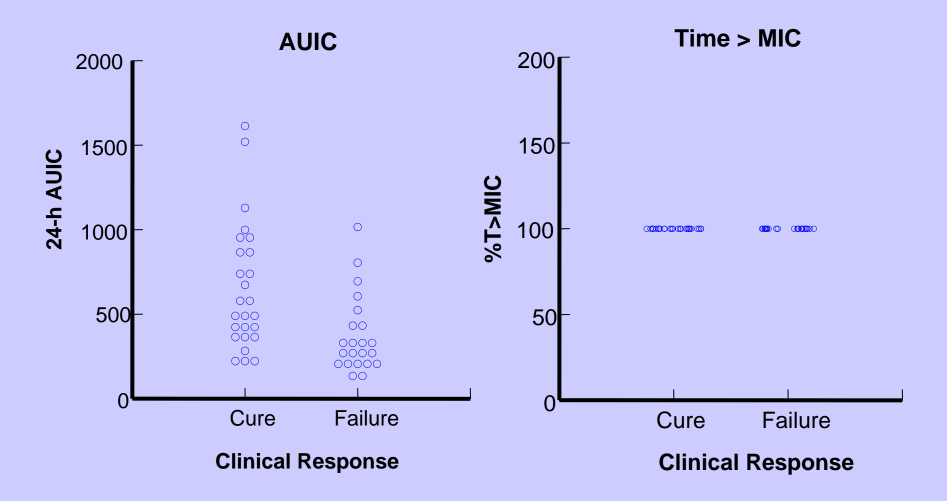
### AUIC vs T>MIC and Microbiological Response

Moise, Forrest, Schentag et al. Clinical Pharmacokinetics 2004; 43: 925-942

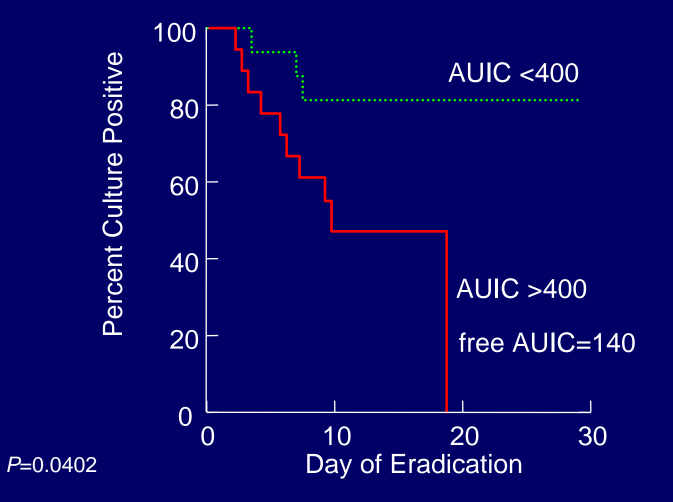


#### AUIC vs T > MIC and Clinical Response

Moise, Forrest, Schentag et al. Clinical Pharmacokinetics 2004; 43: 925-942



#### Comparison of Vancomycin days to eradication for MRSA Infections



Moise & Schentag. Clinical Pharmacokinetics 2004; 43: 925-942

#### **Computerized Estimation of AUIC**

- Selected patients who are now undertreated will benefit from the addition of a second antibiotic, or higher doses
  - Less resistance, fewer failures, shortened therapy
- Most cephalosporin doses will be lowered (elderly patients, low MIC organisms)
  - Cost Savings in the antibiotic budget

## Use of AUIC in Patient Care

- 77 yoM, 70 in, 155 lb, with COPD, Lung Ca, and Diabetes, 7 days post-op LLL resection.
- Now with new S&S of LRTI, on a Ventilator
- Cefazolin for prophylaxis day 1, currently receiving no ABX. Serum creatinine is 1.2 mg/dl
- Cx taken, Ceftazidime 1.0 gm Q12hr is ordered.
- You were consulted for antibiotic management