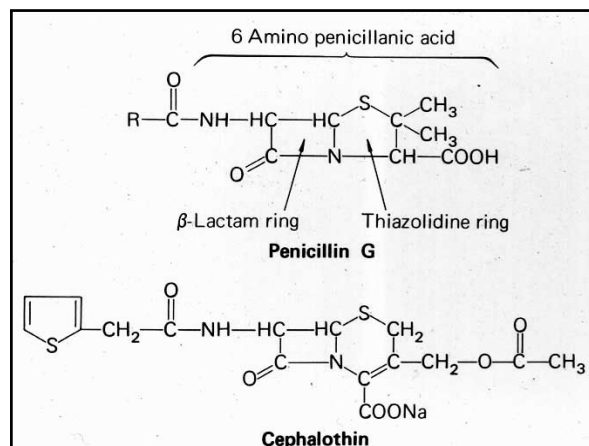


Beta-lactam antibiotics - Cephalosporins

Targets - PBP's

Activity - **Cidal** - growing organisms (like the penicillins)

Principles of action - Affinity for PBP's
Permeability properties
Stability to bacterial enzymes



Cephalosporins

Development - Giuseppe Brodtzu - Sardinian sewage

Cephalosporin C - Cephalothin

No meningeal penetration

Failed in meningococcal meningitis

Painful to give IM

Advantages

Cephalosporin nucleus - resistant to Staphylococcal penicillinase

Cephalosporin nucleus - more readily modified

Development of C'sporins

Generations - in response to clinical needs

First generation - Cephalothin (not used)

Cefazolin

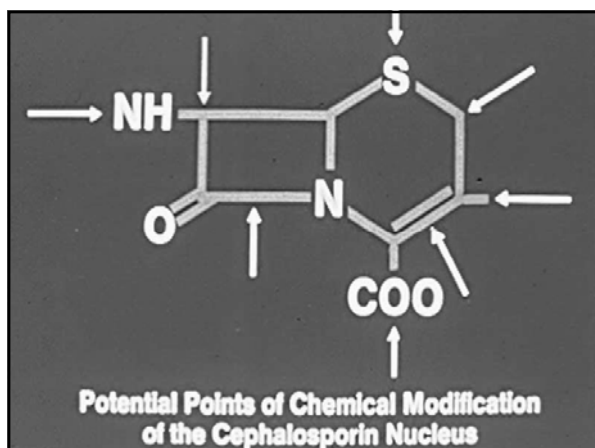
oral - Cephalexin, cefaclor

Activity - Broad spectrum:

Gram positive *Streptococci*, *S. aureus*

Gram negative - *E.coli*, *Klebsiella*

No activity against *Enterococci* - different PBP's



Second generation C'sporins

Cefuroxime

Cefoxitin

Cefotetan

70's - Beta-lactamase's recognized (*H. influenzae*)
Anaerobic infections

Cefoxitin - Methoxy group - conferred beta-lactamase stability

Induction of chromosomal beta-lactamases

Bacteroides fragilis - enteric anaerobes

Cefuroxime - Respiratory tract infections -

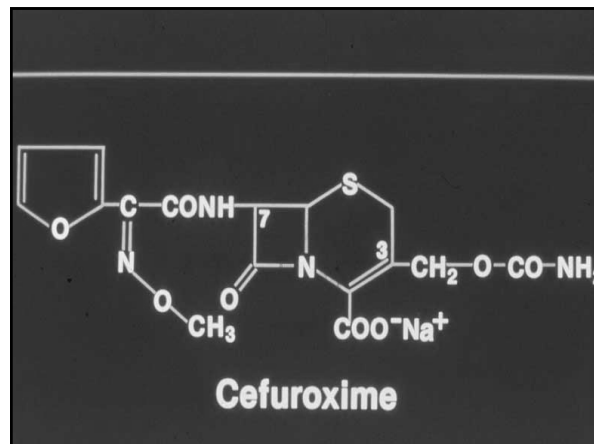
community acquired

Kinetics of c'sporin binding

Affinity for receptor - PBP

Permeability characteristics of the porin

Beta-lactamase production - within periplasmic space



Third generation C'sporins

80's - Intensive care - nosocomial infections

↓

Multi-Resistant Gram negative organisms

Chromosomal beta-lactamase - C'sporinase Inducible

Plasmid mediated enzymes - mutants with both Penicillinase and C'sporinase activity

Permeability limitations

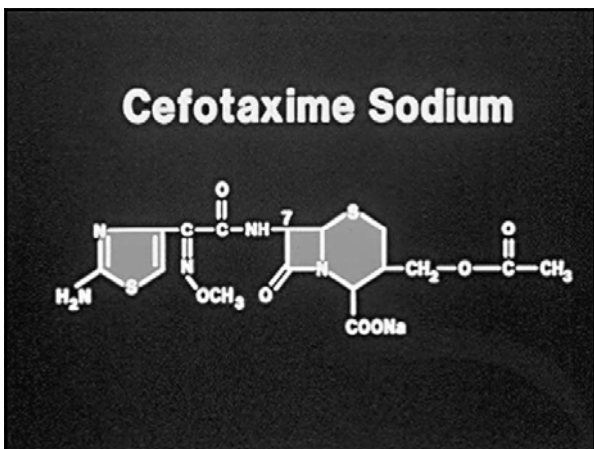
Third generation c'sporins

Cefotaxime
Ceftriaxone
Ceftazidime
Cefipime (4th?)

Highly active - Cefotaxime - *S. pneumo*
N. meningitidis, gets across BBB

Ceftriaxone - even more active - Single dose IM
get meningeal levels - Long half life !!!
N. gonorrhoeae, use in unreliable patients -
Cover *S. pneumonia* bacteremia

Use in meningitis -



IMPORTANT PHARMACOKINETIC VARIABLES OF NEW CEPHALOSPORINS

Agent	Serum protein binding (%)	Metabolism	Peak serum levels (µg/ml)		Half-Life (hours)		Urinary recovery (%)
			Ig* 0.5g †	0.5g †	Cr<90	Cr>10	
Moxalactam	50	-	60	24	2	19	75
Cefotaxime	38	+	42	12	1.1	2.5	55
Desacetyl cefotaxime	23	+	7	3	1.6	11	30
Ceftizoxime	31	-	87	14	1.4	25	85
Ceftriaxone	83-96	-	150	50	8	11-16	90*
Ceftazidime	17	-	80	18	1.8		75
Cefoperazone	90	-	125	26	1.9	2.5	12

*Based on intravenous infusion over 30 minutes.
†Intramuscularly
H.C. Neu, Bull. N.Y. Acad. Med., 60:327, 1984

Ceftazidime/Cefepime - anti- *Pseudomonas*

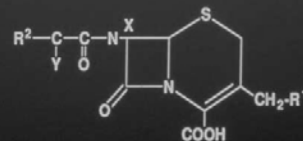
Used the side groups which have increased permeability through *Paeruginosa* porins -

? Induction (low level) of chromosomal C'sporinase

Beta-lactamase stable -

less activity against gram positive organisms

β-LACTAMASE STABILITY OF CEPHALOSPORINS



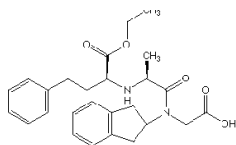
X = O-CH₃ As in Cefoxitin

Y = N-OCH₃ As in Cefuroxime
Cefotaxime

Y = N-C(CH₃)₂-COOH As in Ceftazidime

Cefepime – Fourth generation

Increased beta-lactamase stability
Also better Gram positive -



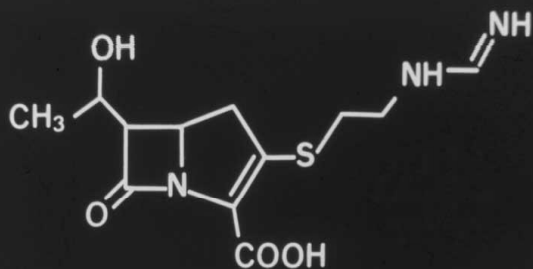
Carbapenems

Imipenem
Meropenem
Ertapenem

Beta-lactam class - PBP-2 major target
Permeability - separate porin

Huge spectrum - Aerobes, anaerobes
everything EXCEPT
Enterococci
Stenotrophomonas etc.

Concern - CNS side effects - Imipenem ??



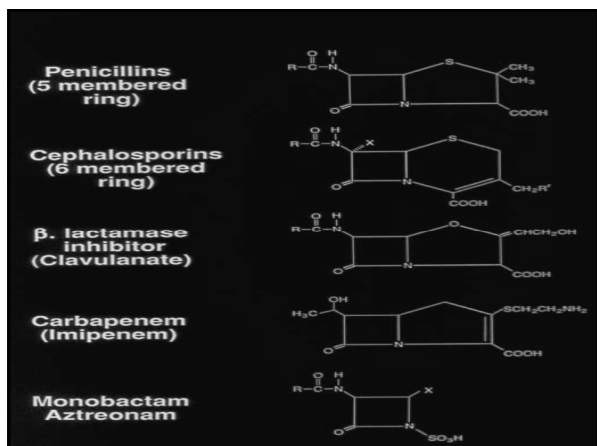
IMIPENEM

Monobactams - Aztreonam

Only binds to Gram negative PBP's

No real beta-lactam ring - therefore beta-lactamase stable

Narrow spectrum - Only aerobic gram negative rods
Use - instead of an aminoglycoside



Use of the cephalosporins:

First generation - Oral - surgical prophylaxis - skin soft tissue infections - taste good - "house cephalosporin"

Second generation - Some oral - some parenteral
Selected uses – community acquired infections

Parenteral - Third generation

Increased - due to resistant *S. pneumoniae* - susceptible to cefotaxime and ceftriaxone

Gram negative infections - hospital acquired - selection of resistant organisms

4th generation?

- Cefapime/ceftazidime
- Immunocompromised patients
- *P. aeruginosa* - known gram negs
- ? Worry about induced beta-lactamases
- ESBL + organisms - prefer carbapenem/penicillin

Resistance Rates

MYSTIC program (USA 199-2006)
>100 medical centers

Resistance to carbapenems	
Enterobacteriaceae (9,396 organisms)	0.5%
<i>Pseudomonas aeruginosa</i> (3,100 organisms)	7.2%
All (20,051)	2.8%

Pharmacology

Charged - hydrophilic - do not enter phagocytic cells

Variably protein bound (Ceftriaxone - highly bound)

Variable half-lives

Metabolism - Cefotaxime - Liver - desacetyl derivative - active

Excretion - Renal - Tubular secretion and glomerular filtration

Beta-lactams – side effects

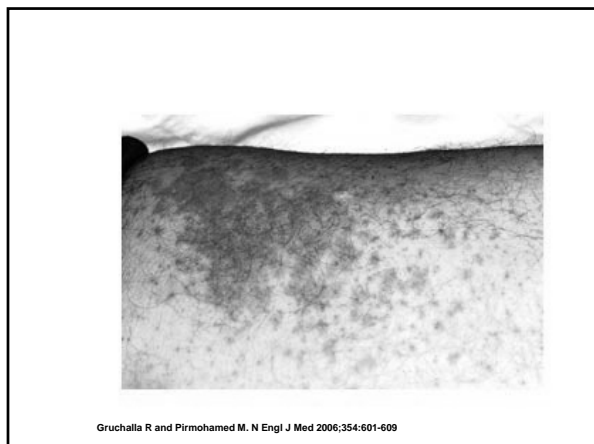
penicillin – c'sporin cross reactivity – 3-7% (depending on the drug)

Hypersensitivity – Rash
IgE-mediated allergy – Anaphylaxis

Major determinants – minor side effects
Minor determinants – MAJOR reactions

Diarrhea
Neutropenia

CNS – high doses - especially the carbapenems



Adverse Reactions to Cephalosporins

TABLE 1. ADVERSE REACTIONS TO CEPHALOSPORINS.

TYPE OF REACTION	FREQUENCY	REFERENCES
	%	
Dermatologic	1.0-2.8	Norby, ¹ Sanders et al., ² Arndt and Jick, ³ Platt ⁴
Positive direct antiglobulin test	1.0-2.0	Sanders et al., ² Platt, ⁴ Meyers ⁵
Anaphylaxis	0.0001-0.1	Gadde et al., ⁶ Sogn et al. ⁷
Fever	0.5-0.9	Sanders et al., ² Meyers ⁵
Eosinophilia	2.7-8.2	Sanders et al., ² Platt ⁴

Kelkar P and Li J. N Engl J Med 2001;345:804-809

C'sporins

Intrinsic resistance - enterococci - different targets

Acquired resistance - active change

- Acquisition of an enzyme
- Induction of an enzyme
- Selection of a mutation
- Alteration in permeability

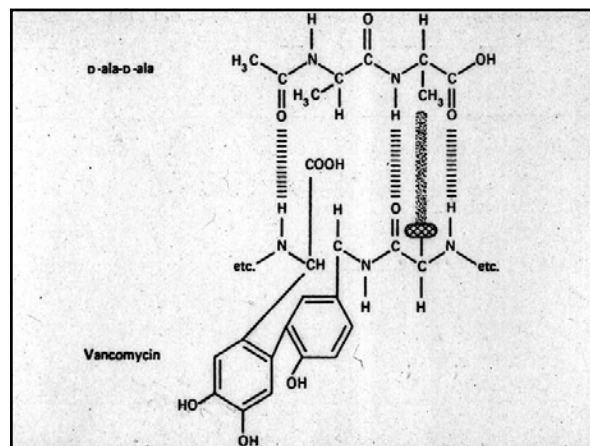
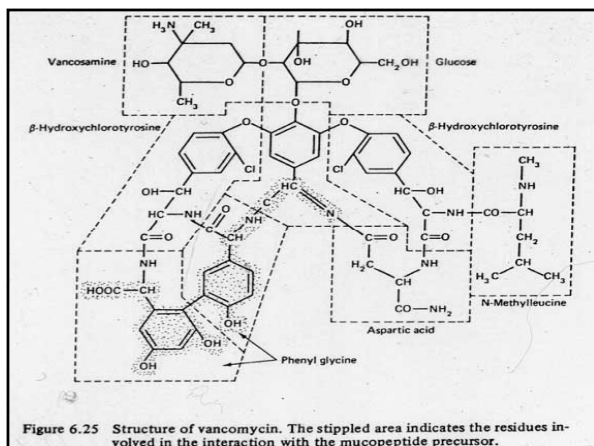
Vancomycin

History - Developed in the 50's - anti-Staph drug

Re-"discovered" - MRSA - and MRSE - *Staphylococci* with altered PBP-2A **mecA** gene - no longer binds penicillin (C'sporins don't bind either)

Target - **D-ala-D-ala** - pentapeptide blocks two steps in cell wall synthesis

Cidal - Only gram positives - Highly resistant *S. pneumo*



Methicillin resistant *Staphylococci*

- *mecA* mutations - altered PBP's
-
- often linked to overexpression of beta-lactamase
-
- Use different class of antimicrobial agent

MRSA - methicillin resistant *S.aureus*

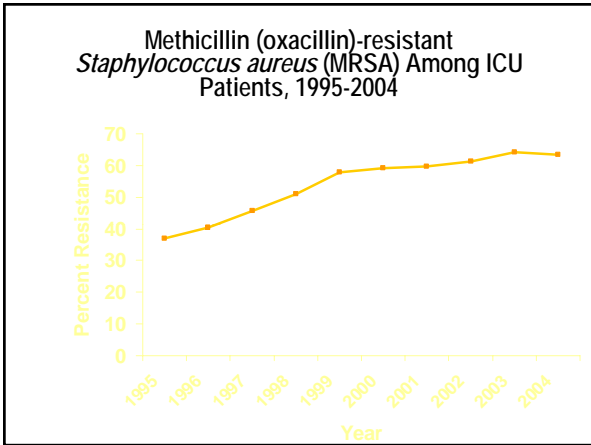
- *mec A* mutations - Altered PBPs
- Overexpression of beta-lactamases as well in some strains
- Use different class of antibiotic

Use of Vancomycin

Staphylococci – resistant to penicillin – “methicillin resistant - Altered PBP's

Coagulase- negative staphylococci – Catheter infection

S. aureus – MRSA – Methicillin Resistant *Staphylococcus aureus*



MRSA - types

Nosocomial – Multi-resistant – large chromosomal insertions – *mecA* cassette

Community – SCC's (small covalent circles)
Integrated elements along with the recombinases

Very common – moving back into the hospital

Community Acquired MRSA

Increasingly common – smaller mobile genetic unit
USA300 - epidemic clone in the USA

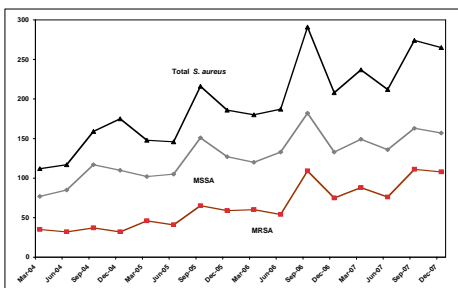
Still susceptible to other antibiotics (unlike hospital –acquired)
(TMP/SMX, fluoroquinolones, Clindamycin)

Often relatively virulent – Panton-Valentine toxin
Skin- soft tissue infections
Fulminant pneumonias

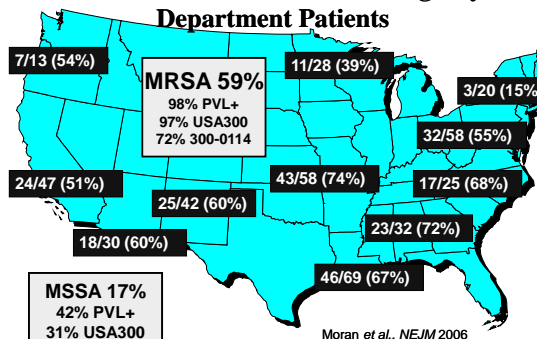
Adolescents

Up to 70% of outpatient isolates !!!! At some centers

S. aureus Outpatient Skin/Soft Tissue Clinical Isolates at Columbia 2004-2007



MRSA Prevalence as a Cause of Skin/Soft Tissue Infections in Adult Emergency Department Patients



Vancomycin - properties

Small glycoprotein (MWt @ 1,450) derived from *Nocardia orientalis*

Activity - most G(+) bacteria including Streptococci, Corynebacteria, Clostridia, Listeria, and Bacillus species.

Bactericidal at levels 0.5 - 3 mg/L

Staphylococci including β -lactamase producing and methicillin resistant species are killed at levels <10 mg/L

Vancomycin - Pharmacokinetic properties

Vd @ 0.7 L/kg
 Protein binding @ 55%
 Elimination: > 90% renal

Half-life @ 7 hrs (with normal CLcr)

Vancomycin is not removed by standard HD or PD, but it is removed by CVVH

Side effects of vancomycin:

Red man syndrome - histamine-mediated erythematous flushing of the face, neck and trunk, a reaction which occurs during the infusion, and may be associated with hypotension.

Nephrotoxicity and ototoxicity ?? < 1% of pts especially those receiving other 'toxic' drugs like aminoglycosides.

A relationship between vancomycin level and nephrotoxicity or ototoxicity has not been established.

It is now widely believed that the earlier reports of nephrotoxicity may have been related to impurities in the product.

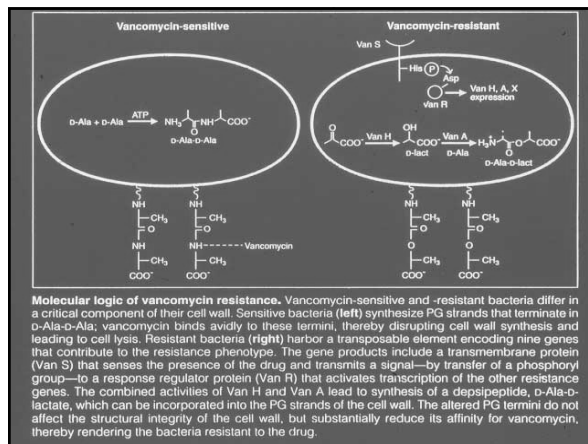
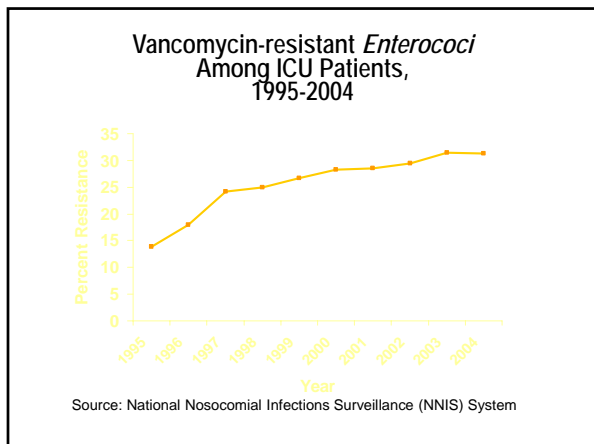
Vancomycin resistance

VRE = Vancomycin resistant enterococci
 ? From oral use of vancomycin

Selection of enterococci – altered cell wall structure

Several mechanisms:
 D-ala-D-ala changed to a lactate
 No metabolic cost

Several Vanco resistance cassettes



Vanco resistant *S. aureus*

- VISA – Vanco intermediate –
MIC's 4-16 micrograms/ml

Multiple point mutations
Thickened peptidoglycan layer
? Sponge effect
(GISA = glycopeptide-intermediate strains)