

Gompf's Antibiotics Redux

**A Pocket Tool for the Medical Student, or Resident on the
Infectious Diseases Clinical Rotation**

or

Just about anyone who could use a pocket antibiotic tool

By

**Sandra G. Gompf, MD
Associate Professor, Infectious Diseases
University of South Florida
Morsani College of Medicine
Tampa, FL**

Updated JUL 2022



Gompf's Antibiotics Redux by Sandra G. Gompf, MD is licensed under a Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License.

Made in the United States of America.

ANTIBIOTIC PEARLS

1. Penicillins generally cover Gram +s, anaerobes, certain Gram –s depending on the antibiotic.
2. Cephalosporins generally cover Gram +s (EXCEPT Enterococcus!), Gram –s, few or NO anaerobes. ONLY ceftazidime/cefepime cover Pseudomonas. They do not cover SPACEK/SPICE* Gram negatives reliably; ceftriaxone/cefepime may be fine in less serious SPACEK/SPICE infections.
3. Aztreonam, a monobactam, covers ONLY Gram –s, incl. Pseudomonas. Reserve for beta lactam-allergic patients.
4. Aminoglycosides generally cover Gram +s (except tobramycin) & Gram –s, NO anaerobes, some Mycobacteria.
5. Quinolones cover Gram –s best (except moxifloxacin, best for respiratory Gram +s & anaerobes), some Mycobacteria.
6. Sulfas generally cover some Gram +s/MRSA, Nocardia, Listeria, Pneumocystis, most Gram –s except Pseudomonas.
7. Clindamycin generally covers Gram +s, incl anaerobes except Clostridia, like anaerobic/microaerophilic Strep/Peptostrep, Actinomyces (better for infections above the diaphragm).
8. Metronidazole generally covers anaerobes like Bacteroides, Prevotella, Clostridia; +/-Peptostrep (better for infections below the diaphragm).
9. Carbapenems are Big Gun Beta Lactams & Expensive. Use sparingly. Ertapenem covers most organisms except Pseudomonas. Imipenem, meropenem, & doripenem include Pseudomonas. Resistance in one carbapenem doesn't predict resistance in others.
10. A word about Enterococcus! Enterococcus in well-defined polymicrobial infections (cholecystitis, GI perforation, decubitus infections) tends to be somewhat of an opportunist & clears once source control is achieved, even if it spills into blood). Isolated Enterococcus faecalis/faecium bacteremia or hematogenous infection (septic arthritis, vertebral osteomyelitis, endocarditis, "community-acquired" bacteremia without an immediately source) should raise as much concern as Staphylococcus aureus. In that setting, even if reported as very ampicillin/penicillin-susceptible, and ampicillin is considered "drug of choice", know that these organisms have *lower*-affinity penicillin-binding proteins than other Gram + cocci, and "tolerance" may also occur with intermittent dosing of ampicillin/penicillin should be considered bacterio-STATIC. Synergistic combination therapy is preferred for these infections, such as amp + gentamicin. Amp + ceftriaxone/ceftazolin is increasingly preferred due to nephrotoxicity with gent & gent resistance. The combination of these beta lactams binds more PBPs than ampicillin alone & lowers the ampicillin MIC. In addition, know that vancomycin is bactericidal for most GPCs *except Enterococcus* (bacterio-STATIC).

Shameless plug:

Visit www.gompsidpearls.net for more regularly updated ID clinical tools & links I find useful in practice.
:}



Do's & DON'Ts

1. **Don't use an antibiotic if you don't need to.**
 - **If a bacterial infection is not high in the differential and the patient is not clinically toxic, forgo antibiotics. They are poor antipyretics.**
2. **Persistent fevers require work-up, not more antibiotics.**

If you are treating with broad antibiotics and fevers persist, **stop them**; they aren't helping.

 - Look for undrained foci of infection/pus → drain it.
 - Look for non-infectious cause → treat it.
 - True FUO in a rapidly deteriorating patient may warrant empiric anti-TB therapy → Call ID.
3. **DO use an oral antibiotic when you can; use one narrow antibiotic when you can; stop antibiotics when you can.**
 - Antibiotics are not cheap; switch to PO when reasonable.
 - Two antibiotics don't always prevent resistance better than one, and neither do broad spectrum drugs. More drugs = more resistance.
 - But NEVER give Rifampin alone! Rapid high-level resistance occurs.
 - TB/Atypical mycobacteria: NEVER use 1 drug in active TB
 - DON'T treat viral infections (or noninfectious syndromes) with antibiotics beyond the point at which you have ruled out bacterial infection.
4. **Don't combine 2 of the same class of antibiotics (e.g. 2 beta lactams, 2 quinolones) unless you have data or your friendly neighborhood ID specialist to back it up. They may unpredictably antagonize, synergize, or double the adverse effects.** E.g. Clindamycin-rifampin combo dramatically reduces clindamycin serum concentration.
[\[https://doi.org/10.1016/j.jinf.2015.03.013\]](https://doi.org/10.1016/j.jinf.2015.03.013); <http://dx.doi.org/10.1684/ejd.2013.2213>
5. **Always monitor for antibiotic adverse effects.**
 - Antibiotics are a double-edged sword. Respect them.
 - Watch for hypersensitivity/bone marrow suppression/interstitial nephritis/hepatotoxicity/drug fever with beta lactams, acute tubular necrosis/irreversible ototoxicity with aminoglycosides, & Clostridium difficile with almost all of them.
 - Watch for yeast overgrowth/Candidemia with prolonged/multiple antibiotic therapy.
 - C. diff. is easy to miss in 2 situations:
 - Colostomies – stumps/small bowel can be infected with C.diff.!
 - Spinal cord injured patients – unexplained abdominal distension & leukocytosis are a clue
 - RIFAMPIN REDUCES EFFECTIVENESS OF ORAL CONTRACEPTIVES! Tell female patients to *add barrier contraception until the next new pill pack* after finishing antibiotics.

Antifungal coverage in general:

Fluconazole = Cryptococcus, Candida EXCEPT Candida krusei/glabrata

Itraconazole = Candida, Histoplasma, Crypto, Aspergillus

Voriconazole = Candida, Histo, Crypto, Aspergillus, Fusarium, NOT Mucor/Rhizopus

Posaconazole /isavuconazole = same as voriconazole, + Mucor/Rhizopus

Caspofungin/Micafungin/Anidulafungin (echinocandins) = Candida, Aspergillus, NOT Fusarium/Mucor/Rhizopus, SOME Crypto

Amphotericin = all, +/- Fusarium, NOT Candida lusitanae/guilliermondii, NOT

Scedosporium (Pseudallescheria)

Fusarium: Vori 6mg/kg IV Q24H or 300mg PO x 1 d, then 4mg/kg/d IV or 200mg PO BID + Ampho B 1.2 mg/kg/d or ABLC 5mg/kg/d

Mucor: Ampho B 1.5mg/kg/d or liposomal ampho B or ABLC 5mg/kg/d + posaconazole/isavuconazole (active metabolite of isavuconazonium); NOT other azoles/enchinocandins

5FC increases penetration of above drugs.

BacteriCIDAL vs. BacterioSTATIC

A consideration in choosing treatments for serious infection like severe sepsis, bacteremia, meningitis, pneumonia, endocarditis, osteomyelitis, neutropenic fever. A "cidal" drug kills quickly; a "static" drug slows or stops replication and/or toxic production.

Beta lactams are CIDAL and penetrate tissues and inflamed meninges well. They are preferable in serious infection, including bacteremia, endovascular infection, CNS infection, and streptococcal cellulitis. Their microbial action is *time-dependent*, meaning that they are most effective the longer the concentration of drug in the affected site remains above the MIC of the bacteria. Thus, they can be dosed by *continuous or extended infusion*, which may also facilitate home & hemodialysis dosing. (UpToDate has a helpful chart; most drug databases like ePocrates, etc, don't offer alternative dosing recommendations.) Ceftazidime, cefepime, and ertapenem are stable enough to be given 3 times a week after hemodialysis. Ampicillin is stable at least 24 hrs for extended and continuous infusion.

Beta Lactamase Resistance: It's complicated.

SPICE/SPACEY/SPACEK are mnemonics for bacteria that either have *intrinsic* (chromosome-based, aka "constitutive") and/or *inducible* beta lactamases (chromosome- OR plasmid-based ampC).

These organisms may all demonstrate resistance to commonly prescribed beta lactams and may require carbapenem* treatment. In addition, inducible beta lactamases may become reversibly de-repressed (or "switched on") upon 1- 7 days' exposure to a beta lactam and result in treatment failure. Severity of infection (bacteremia, meningitis), source control (UTI/pyelonephritis vs endocarditis), presence of biofilm (infected device), & MICs to cefepime (≤ 1 mg/L) & pip-tazobactam (≤ 4 mg/L) ** can help guide whether to use a carbapenem.

SPACEK – Grouped by presence of **chromosomal ("constitutive", "intrinsic") or mobile genetic element-acquired beta lactamases**

Serratia

Proteus (non-P. mirabilis)/Providencia

Pseudomonas

Acinetobacter complex

Citrobacter

Enterobacter complex

Klebsiella – majority are intrinsically resistant to ampicillin/PCN (SHV-1) at this point

SPICE/Y – Grouped by Family Enterobacteriaceae with **inducible, chromosomal beta-lactamase (AmpC)**

Serratia

Providencia

Indole+ Proteus (non-P. mirabilis)

Citrobacter

Enterobacter/E.coli/Klebsiella

Yersinia enterocolitica

The SPICE organisms may produce inducible chromosome-based broad-spectrum beta lactamases as part of the Enterobacteriaceae group, and resistance/failure may be induced during beta lactam treatment, even though they initially test susceptible. **E. coli and Klebsiella** are the most common extended spectrum beta lactamase (ESBL) producers, so many labs screen those isolates if MIC for ceftazidime is ≥ 2 mg/L. Remember that **Klebsiella** almost all have a constitutive chromosome-based beta lactamase (usually SHV-1) that confers resistance to ampicillin/ticarcillin, so these drugs are never a good choice for this bacterium. Preferred treatment in serious infection is a carbapenem.

*Note that carbapenems and the monobactam, aztreonam are beta lactams, as they all have a beta lactam ring. This may be confusing initially when you read about beta-lactam resistance and recommendations to use a beta lactam “versus” carbapenem; many references gloss over this.

**See Impact of the MIC of Piperacillin-Tazobactam on the Outcome of Patients with Bacteremia Due to Extended-Spectrum-Lactamase Producing Escherichia coli, 2013, <https://doi.org/10.1128%2FAAC.00135-13>.

Antibiotics for Resistant Gram + Cocci

- Vancomycin, teichoplanin (Europe) cover all but vancomycin-resistant Gram +s.
- Vancomycin is bacterioSTATIC against Enterococcus.
- Dalbavancin, oritavancin, telavancin - same mechanism as vancomycin & susceptibility generally correlates with vancomycin susceptibility; they add VRE, VISA/VRSA coverage.
- Daptomycin is bacteriCIDAL for both MRSA and Enterococcus & penetrates biofilm.
- Linezolid & tigecycline are bacterioSTATIC, NOT a choice for bacteremia unless no other options are possible, and not as monotherapy, unless you can cite legitimate supportive data.
- Tigecycline is difficult to tolerate due to 20% nausea/vomiting & associated with higher mortality than comparators for FDA-approved indications in after-market review of pooled clinical trials.
- All but vancomycin are EXPENSIVE & generally reserved for vancomycin-resistant Gram +s.

Which antibiotics are bacterioSTATIC?

“In sepsis, restore Volume with a Liter of NML (normal) Saline.”

Vancomycin in Enterococcus; cidal for all other GPCs

Linezolid

Tetracyclines/Tigecycline

Nitrofurantoin

“MLS antibiotic group” – clindamycin, macrolides (Note: streptogramins are bactericidal)

Sulfas/trimethoprim

Everything else is bactericidal & probably better for sepsis and serious infections!

Note bene: Clindamycin is used as an *adjunct* for Staph or Strep toxic shock, severe streptococcal cellulitis or suspected necrotizing infection; it *halts protein synthesis*—i.e stops production of toxins that mediate severe inflammation, necrosis, and toxic shock. Many *Staphylococcus aureus* strains carry *inducible* clindamycin resistance genes, so I suggest having susceptibilities available before relying on clindamycin alone for this pathogen (if your lab doesn't report inducible clindamycin resistance, check for erythromycin resistance-- *erm* mutation! ---as a clue). You can also use other drugs whose mechanism of action is disruption of protein synthesis: linezolid, doxycycline/minocycline/tigecycline for toxin-inhibition in severe Staph infection.

What does the MIC mean & how do we use it?

The MIC is the lowest concentration of an antimicrobial agent that completely prevents visible growth of the test strain of an organism in culture. Each antibiotic will have a different MIC for each organism. The MIC value of an antibiotic must be compared with clinical out-

comes for the disease caused by a given organism; in some cases, this is further drilled down to differentiate between clinical scenarios (e.g., meningitis vs UTI), and the pharmacokinetics of individual antimicrobials. This overview yields “*clinical breakpoints*” that define whether the strain is susceptible (S), susceptible dose dependent (SDD) or resistant (R) to the antibiotic. Clinical breakpoints are internationally reviewed & adjusted yearly according to expert consensus, epidemiology, and clinical literature by EUCAST (European Committee on Antimicrobial Susceptibility Testing), the American CLSI (Clinical and Laboratory Standards Institute), and further updated by the FDA (Food and Drug Administration). Fortunately, your friendly hospital Micro lab and their automated systems do a LOT of the interpretation work for you. If Micro says it’s susceptible, it usually is—just beware of SPICE/SPACEKs (♠). And vancomycin “MIC creep” with *S. aureus* bacteremia—as MIC “creeps” above 1, so does treatment failure. Etc, etc.

THE CLASSES (not an exhaustive list)

Penicillins – beta lactams are CIDAL, good tissue penetration

DRUG	COVERAGE	USES	TOXICITY	Cerebral Spinal Fluid (CSF)
penicillin G \$ CIDAL	Group A Strep (no resistance) Strep viridans Neisseria Capnocytophaga Actinomyces Fusobacterium Clostridia perfringens/tetani Pasteurella Treponema/ Leptospirosis NOT Staph aureus (resistant)	Skin/soft tissue (SST) or mouth infections	Hypersensitivity Stevens Johnson Interstitial nephritis Seizures (if high level) Bone marrow suppress-ion C.difficile	YES if inflamed
AminoPCN \$\$ amoxicillin* ampicillin* amox/clav amp/sulbact CIDAL	Add to the above: Listeria MSSA Most Pneumococcus Proteus Hemophilus influ. (beta lactamase negative) Salmonella/Shigella Anaerobes * <i>Klebsiella are intrinsically resistant to amp/amox</i> (clavulanate/sulbactam don't add much activity)	Otitis media Sinusitis SST Meningitis in elderly	Above	
CarboxyPCN \$\$ ticarcillin/clav (Europe) piperacillin piperacillin/tazobactam CIDAL	Adds to the above: Pseudomonas Enterobacteriaceae* Stenotrophomonas (ticar) Gut anaerobes MSSA Pip & Pip/tazo more potent for GNRs & more resistant to AmpC/ESBLs (See "SPICE" above) * <i>Klebsiella is intrinsically R to ticarcillin</i>	Adds to above: Gut/ surgical infections Nosocomial pneumonia Prostate Osteomyelitis	Above	

Cephalosporins – Think of progressive broadening of spectrum from Gram + to Gram - with each generation. Beta lactams are CIDAL. [⊙ = can be dosed 3 times weekly in dialysis patients]

DRUG	COVERAGE	USES	TOXICITY	CSF
1st Generation \$\$ cefalothin cefazolin cefadroxil CIDAL	GPC, E. coli, Proteus, Klebsiella (increasingly ampC+/ESBL) NOT Enterococci	SSTI Uncomplicated/Non-diabetic osteomyelitis PreOP prophylaxis	Hypersensitivity Bone marrow suppression Diarrhea C.difficile	POOR

<p>2nd Generation</p> <p>\$\$</p> <p>cefuroxime (IV/PO) cefaclor (PO)</p> <p>Cefamycins: cefoxitin (IV) cefotetan (IV)</p>	<p>GPC Pneumococcus Neisseria Some GNR except Pseudomonas Cefamycins are the only ones that reliably cover <i>anaerobes</i></p> <p>NOT Enterococci</p>	<p>Community acquired pneumonia (CAP) meningitis OM/sinusitis</p> <p>Gonorrhea</p>	<p>Hypersensitivity RASH/Stevens Johnson w/ cefaclor</p> <p>High INR/PT w/ cefoxitin/cefotetan</p> <p>Bone marrow suppression C.difficile</p>	<p>YES if inflamed</p>
<p>3rd Generation</p> <p>\$\$</p> <p>ceftriaxone (QD dosing) cefotaxime ceftazidime ☺</p> <p>CIDAL</p>	<p>Above, plus Pseudomonas for ceftazidime</p>	<p>Meningitis CAP Most community-acquired infections Gonorrhea Pyelonephritis</p>	<p>Above</p>	
<p>4th Generation</p> <p>\$\$</p> <p>cefepime ☺</p> <p>CIDAL</p>	<p>Above, plus Pseudomonas More resistant to beta lactamases/ESBLs (See "SPICE" above)</p> <p>NOT Enterococci</p>	<p>Above, plus neutropenic fever</p>	<p>Above</p>	
<p>Anti-MRSA</p> <p>\$\$\$</p> <p>ceftaroline</p> <p>CIDAL</p>	<p>Similar to 3rd generation, plus MRSA, VISA/VRSA/VRE faecalis (NOT E. faecium), pneumococcus, beta-lactamase + H. flu, Moraxella</p>	<p>Complicated SSTI, CAP (NOT MRSA-insufficient data)</p>	<p>Above</p>	

<p>Advanced-generation</p> <p>ceftolazane-tazobactam</p> <p>ceftazidime-avibactam</p> <p>CIDAL</p>	<p>NOT Enterococci or Staphylococci</p> <p>ceftolaz-taz covers GNRs incl Pseudomonas, ESBLs, NOT carbapenems</p> <p>ceftaz-avi covers KPC+ carbapenemase (1st line agent)</p> <p>ceftaz-avi covers GNRs incl Pseudomonas, adds coverage for ceftaz-R, ESBLs, some ampC-R, some carbapenemases (NOT metallobeta-lactamase)</p>	<p>Complicated UTI/pyelo</p> <p>Complicated intraabdominal infection</p> <p>ceftaz-avi adds HAP</p>	<p>Above</p> <p>Nausea, diarrhea, headache, fever, renal insufficiency (ceftolazane-t)</p>	<p>ceftazidime – YES if inflamed (NOT avibactam)</p> <p>ceftolazane – UNKNOWN</p>
---	--	---	--	---

Monobactam

DRUG	COVERAGE	USES	TOXICITY	CSF
aztreonam \$\$ CIDAL	ONLY GNRs, incl Pseudomonas	GNR infections; NOT a replacement for all aminoglyco- side uses (no syner- gy for GPC, NO Enterococcal cover- age)	Low	YES if in- flamed [Modal J et al. AAC. 1986;29:281-3.]

Carbapenems (Reserved for Multidrug Resistant Organisms – MDRO)

[⊙ = can be dosed 3 times weekly in dialysis patients]

DRUG	COVERAGE	USES	TOXICITY	CSF
imipenem/ cilastin meropenem meropenem- vaboractam \$\$\$	GPCs EXCEPT MRSA GNRs EXCEPT Stenotrophomonas/Burkholderia ESBL+ & "SPICE" GNRs Anaerobes (incl Cutibacterium) Listeria Pneumococcus Nocardia asteroides (NOT brasiliensis) Legionella Mycobacterium avium +/- Enterococcus mero-vaboractam adds <i>carbapenemase</i> + <i>Klebsiella pneumonia</i> (KPC), class A carbap-R Enterobacteriaceae (NOT metallo-beta-lactamase/OXA carbap-R, NOT carbap-R Pseudomonas/Acinetobacter)	Resistant GNR infections Serious gut infections Necrotizing pancreatitis	IV/IM Hypersensitivity (~10% cross-allergy with beta lactams) Seizures (if renal insufficiency or high levels used) with imipenem Candida overgrowth/infections C.difficile Encephalopathy	YES
doripenem \$\$\$ CIDAL	Above, possibly lower MICs to Pseudomonas & Acinetobacter	Above	Above	
ertapenem ⊙ \$\$\$ CIDAL	Above, without Pseudomonas coverage	Postpartum uterine infections Postsurgical Abdominal infections (not Pseudomonas)	Above	

Aminoglycosides

DRUG	COVERAGE	USES	TOXICITY/ MISC	CSF
gentamicin streptomycin spectinomycin tobramycin amikacin liposomal amikacin \$-\$\$\$ CIDAL	Gent: GPCs & GNRs incl Pseudomonas Tobra/Amik: GNRs incl Pseudomonas Amik: Mtb, NTM Strepto -Yersinia -MDR Mtb Gent/Strept -Tularemia Spectino -Gonorrhea	Synergy with beta lactams for GPC/Pseudomonas infections Usually not used alone except for UTIs	IV/Aerosol Acute tubular necrosis (reversible) Ototoxicity/ Vestibular toxicity (irreversible) When possible: -stop after 3-5 d -use once-daily dosing -avoid in elderly Liposomal amik – hypersensitivity pneumonitis, hemoptysis, bronchospasm, exacerbation of lung disease Neuromuscular blockade (may exacerbate myasthenia gravis & paralytic agents)	NO-UNKNOWN
fosfomicin \$\$ CIDAL	Enterococcus GNRs	Simple cystitis in women Off-label q3days for complicated or MDR GNRs, VRE if susceptible	PO only Above, significant diarrhea	
plazomicin \$\$ CIDAL	GNRs incl MDR/KPC/metalobetalact/CRE GNRs,variable Pseudomonas (use only if known susceptible), NOT Steno, Acinetobacter	Complicated UTI/pyelo	IV only Above Limited data	

Sulfonamides/Sulfas

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
trimethoprim-sulfamethoxazole co-trimoxazole \$ STATIC for Staph	Staph. aureus (incl MRSA) Legionella Stenotrophomonas Listeria Pneumo-cystis Nocardia Burkhold-eria cepacia Yersinia Francisella tularensis Some common coliforms	UTI MRSA SSTI Specific agents at left	IV/PO RASH/Stevens Johnson Elevated creatinine or K+ (competes with Cr for tubular secretion, blocks K+ excretion) Kernicterus in neonates C.difficile Sun sensitivity	YES

Macrolides/Lincosamides (Macrolide-Lincosamide-Streptogramin B class, or MSL—all bind 50s ribosome subunit & share resistance genes)

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
Macrolides erythromycin clarithromycin azithromycin \$\$ STATIC	Pneumo-coccus MSStaph. aureus (not MRSA) Legionella Listeria Neisseria meningitidis Hemophilus Moraxella Mycoplasma Chlamydia Actinomyces Atypical mycobacteria	LRTI/ bronchitis Sinusitis Dental/ oral infections Atypical mycobacteria (incl MAC prophy in HIV)	IV/PO Nausea/ vomiting Abdominal cramps/ diarrhea (Lowest with Azithro) C.difficile Ototoxicity with chronic use Rare association with cardiovascular mortality with QTc prolongation, low Mg ⁺⁺ /K ⁺ . Interactions: Ery/Clari induce P450! Neuromuscular blockade with Ery (may exacerbate myasthenia gravis & paralytic agents)	POOR

<p><u>Lincosamides</u></p> <p>clindamycin</p> <p>\$\$</p> <p>STATIC</p>	<p>CIDAL for Group A streptococcus, MSStaph. aureus (MRSA, but watch for inducible <i>erm</i> resistance. Clue is resistance to erythromycin)</p> <p>Pneumococcus</p> <p>Inhibits toxic proteins in severe Strep A & S. aureus/ necrotizing fasciitis.</p> <p>Oral anaerobes: Gram + such as Peptostreptococcus, Fusobacterium, Prevotella, Actinomyces, & Clostridial spp other than Clostridium difficile</p> <p>Gram – such as Bacteroides (may not cover in up to 25% of cases or strains with MIC \geq 8 mcg/mL)</p> <p>Babesiosis</p>	<p>Severe SSTI, necrotizing fasciitis, MRSA</p> <p><i>"Infections above the diaphragm"</i></p> <p>Head and neck/dental infections</p> <p>Lung abscess/ aspiration pneumonia (<i>tip: no teeth = no anaerobes</i>)</p> <p>Bacterial vaginosis</p> <p>Babesiosis</p> <p>Toxoplasma in HIV</p>	<p>IV/PO <u>C.difficile!!</u> (>30% develop it on a week of clinda)</p> <p>Watch for hepatitis/obstructive jaundice</p> <p>Neuromus-cular blockade (may exacerbate myasthenia gravis & paralytic agents)</p>	<p>POOR except for Toxo-plasmosis in HIV</p>
---	--	---	---	--

Nitrofuran

DRUG	COVERAGE	USES	TOXICITY/MISC	CNS
nitrofurantoin \$ CIDAL	GNRs EXCEPT Pseudomonas, Proteus, and Enterococcus incl susceptible VRE Multiple sites of action, inhibits syn- thesis of DNA, RNA, proteins, cell wall – higher resistance barrier than most antibiotics	Cystitis Susceptible ESBL GNRs ONLY reaches therapeutic level in URINE	PO only Nausea/ vomiting C.difficile	NONE

Quinolones (Resistance is rising due to overuse)

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
<p>"Gram negative" Quinolones</p> <p>ciprofloxacin levofloxacin norfloxacin</p> <p>\$\$</p> <p>CIDAL</p>	<p>GNRs including Pseudomonas</p> <p>Levo covers pneumococcus well</p> <p>"Atypical" pneumonia: Mycoplasma, Chlamydia, Moraxella</p> <p>Some mycobacteria/TB</p>	<p>UTI/GU infection Intraabdominal infections Endometritis</p> <p>Hospital-associated lung infections</p> <p>Levo best for acute sinusitis/ CAP</p> <p>Norflox: UTI only</p>	<p>IV/PO <i>IV=PO (bioequivalent)</i></p> <p>Dizziness/CNS Diarrhea Hypo-/hyperglycemia Sun sensitivity</p> <p>May exacerbate myasthenia gravis & paralytic agents (inhibits GABA receptors)</p> <p>May prolong QTc (watch for palpitations/syncope)</p> <p>Rare spontaneous tendon rupture (watch for pain at tendon sites)</p> <p>C.difficile</p>	<p>YES, HIGH DOSE</p>
<p>"Gram positive or Respiratory" Quinolone</p> <p>moxifloxacin</p> <p>\$\$</p> <p>CIDAL</p>	<p>Pneumococcus, Streptococci, Staphylococcus (NOT MRSA) Legionella Gut anaerobes</p> <p>Atypical mycobac/TB</p>	<p>CAP/community-associated respiratory infections Acute sinusitis</p> <p>Intraabdominal infections SSTI</p>	<p>IV/PO <i>IV=PO (bioequivalent)</i></p> <p>Above</p>	<p>UNKNOWN</p>
<p>"Gram positive or SSTI" Quinolone</p> <p>delafloxacin</p> <p>\$\$</p> <p>CIDAL</p>	<p>Streptococci, Staphylococcus (NOT MRSA) Legionella Gut anaerobes</p> <p>Atypical mycobac/TB</p>	<p>SSTI</p>	<p>IV/PO <i>IV=PO (bioequivalent)</i></p> <p>Above</p>	<p>UNKNOWN</p>

Nitroimidazole

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
<p>metronidazole</p> <p>\$\$</p> <p>CIDAL</p>	<p>Gram – anaerobes incl. Bacteroides fragilis and all Clostridia</p> <p>Entamoeba coli</p>	<p><i>"Infections below the diaphragm"</i> Intraabdominal abscess, peritonitis, diverticulitis, etc Endometritis/ Bacterial vaginosis Clostridium difficile colitis</p> <p>Entamoeba liver abscess/</p>	<p>IV/PO Disulfiram-like reaction (vomiting) if ethanol consumed within 3 days of therapy</p> <p>Aseptic meningitis/ neuropathies, rare</p>	<p>YES</p>

		dysentery NOT to be given alone for lung ab- scess/ENT infec- tions		
--	--	---	--	--

Tetracyclines/Glycylcycline

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
<p><u>Tetracyclines</u></p> <p>Minocycline Doxycycline</p> <p>\$</p> <p>STATIC</p>	<p>MRSA/MSSA Pneumococcus, +/- GAS E. coli Legionella N. meningitidis Hemophilus Moraxella Mycoplasma Chlamydia Listeria Brucella Actinomyces Rickettsia Vibrio Anaerobes: Fusobacterium, Cutibacterium, Peptostreptococcus, Clostridium, some Bacteroides fragilis/melanogenicus</p>	<p>SSTI CAP, esp under age 40 Dog/cat bite prophylaxis as alternative to amox/clav</p>	<p>IV/PO Discoloration of permanent teeth in children</p> <p>Pseudotumor cerebri, esp minocycline! (watch for headache)</p> <p>Sun sensitivity</p> <p>C.difficile</p> <p>Inhibit lipopolysaccharide-induced proinflammatory products</p>	<p>YES</p>
<p><u>Glycylcycline</u></p> <p>tigecycline >** \$\$</p> <p>eravacycline* omadacycline</p> <p>sarecycline (acne only)</p> <p>STATIC</p>	<p>Above, plus Staph. epidermidis Enterococci Corynebacterium N. gonorrhoea ESBL + E.coli/Klebs Stenotrophomonas Acinetobacter Salmonella B. fragilis/ anaerobes Clostridia incl. C.difficile</p> <p>NOT Pseudomonas or Proteus</p> <p>*eravacycline adds ESBL, carbap-R Acinetobacter</p>	<p>SSTI Intraabdominal infections CAP/HAP</p> <p>Severe C.difficile Y alveolar, soft tissue, bile/gut entry</p> <p>Poor bone/joint, CNS</p> <p>** Bacteriostatic - NOT for serious infections; > Increased mortality vs. comparators in after-market review of pooled clinical trials, incl in FDA-approved indications.</p>	<p>IV only Above</p> <p>20% tige, 6.5% erava - nausea, vomiting</p> <p>Inhibit lipopolysaccharide-induced proinflammatory products</p> <p>*/** Ampicillin/ Amoxicillin CICAL-preferred in VRE that is ampicillin-susceptible.</p>	<p>UNKNOWN</p>

Glycopeptides, lipoglycopeptides

DRUG	COVERAGE	USES	TOXICITY/ MISC	CSF
vancomycin \$ CIDAL ex- cept <i>STATIC</i> for <i>Entero-</i> <i>cocci</i>	Gram + cocci except VRE/VRSA <i>Most</i> Gram + rods (but see below) Corynebacterium Listeria C.diff (only PO) Increasing vanco- mycin MICs > 1 assoc with treat- ment failures ("MIC creep") Intrinsic re- sistance in: Leuconostoc Lactobacillus Propionobacterium Pediococcus Erysipelothrix Clostridia(non-diff.)	SSTI due to MRSA HAP/CAP due to MRSA Infections due to VRE	Vanc IV≠PO – PO not ab- sorbed from gut Vanc requires a central IV line , due to phlebitis (which may cause fevers, unneces- sary antibiot- ics/cultures/increased lengths of stay...) "Red man syndrome" with vanc (histamine release) if infused too rapidly—infuse over 1-2 hours Leukopenia Thrombocytopenia Rare interstitial necrosis Ototoxicity (abrupt, irreversi- ble, usually elderly) Rarely, DRESS	YES
dalbavancin \$\$\$	MSSA, MRSA, Group A, B strep- tococci, Strep anginosus group	SSTI, osteomyeli- tis/prosthetic joint infection, endocarditis, bloodstream infection 1500mg IV x1 OR 1000mg IV then 500mg in 7 days	Nausea, headache, diarrhea "Red man syndrome" with vanc (histamine release) if infused too rapidly—infuse over 1-2 hours	
televancin \$\$\$	MSSA, MRSA/VISA/VRSA, Group A, B strep- tococci, Strep anginosus group, VSEnterococcus	SSTI HAPneumonia due to MRSA/VISA	N/V, foamy urine QTc prolongation Mortality > with mod/sev renal impairment compared with vanco Possibly teratogenic—avoid in pregnancy unless mater- nal benefit exceeds fetal risk "Red man syndrome" with vanc (histamine release) if infused too rapidly—infuse over 1-2 hours Interferes with coag tests but not coagulation	
oritavancin	MSSA, MRSA, Group A, B, C streptococcus,	SSTI 1200mg IV x1, over 3	Headache, N/V "Red man syndrome" with	

<p>\$\$\$</p> <p>CIDAL including <i>Enterococci</i></p>	<p>Streptococcus anginosus group, VSEnterococcus</p>	<p>hr</p>	<p>vanc (histamine release) if infused too rapidly—infuse over 1-2 hours</p> <p>Artificially prolong PT/INR for up to 12 hr (5.1); aPTT for up to 120 hours, and may prolong PT and INR for up to 12 hr and ACT for up to 24 hr—Use Factor Xa assay for coagulation testing</p> <p>Coadministration with warfarin may result in higher exposure of warfarin and increase risk for bleeding; monitor frequently for signs of bleeding</p>	
---	--	-----------	--	--

Cyclic Lipopeptides

DRUG	COVER-AGE	USES	TOXICITY/ MISC	CSF
daptomycin \$\$\$\$ CIDAL	All Gram + cocci incl. Vanc-/Amp- resistant* Entero- coccus MRSA/VRSA	SSTI Bacteremia Osteomyelitis, Joint infections May be active in biofilms (which usually inactivate antibiotics)	IV only Nausea/vomiting Rhabdomyolysis & associated renal insufficiency (weekly creatinine, CPK) Rare asthmatic pul- monary eosinophilia NOT for primary pneumonia because it is inactivated in alveolar fluid, BUT seems effective in embolic lung infec- tion/septic emboli due to Gram +s, since the infection is more parenchymal. *ampicillin/ amoxicillin (CIDAL) preferred in VRE that is amp-susceptible.	UN- KNOWN

Streptogramins

DRUG	COVER-AGE	USES	TOXICITY/ MISC	CSF
quinupristin- dalfopristin \$\$\$\$ CIDAL	Vanc-/Amp- resistant* Enteroco-ccus faecium MSSA Group A Strep NOT Entero- co-ccus fae- calis or MRSA	SSTI/non-MRSA Bacteremia Endocarditis due to VRE faecium Extremely limited use.	IV only Needs central IV line due to frequent pain, phlebitis, fever >30+% Myalgias/ Arthralgias Nausea/ Vomiting/ Diarrhea *ampicillin/ amoxicillin (CIDAL) preferred in VRE that is amp-susceptible.	UNKNOWN

Oxazolidinone

DRUG	COVER-AGE	USES	TOXICITY/ MISC	CSF
linezolid \$\$\$\$ STATIC except CIDAL for streptococci	All Gram + cocci incl. ** vanc-/amp- resistant* Entero- coccus MRSA/VRSA TB/Atypical myco- bacteria Binds 23S rRNA- blocks formation of 50s/70s ribosomal initiation complex	SSTI MRSA HAP/CAP due to MRSA Osteomyelitis/ Joint infections (very Y bone pene- tration) **NOT for bacteremia without a well- defined and remov- al or draining focus, NOT for endovas- cular infections	IV=PO (bioequivalent) Nausea/ vomiting/ diarrhea Temporary tooth staining Headache Thrombocytopenia/ Neutropenia after 7 days Peripheral/ Optic neuropathies with extended use Lactic acidosis (nausea, fatigue) Serotonin syndrome: Avoid high tyramine food/drink (> 100mg tyramine per meal). E.g. aged cheeses, dried/processed meats, ethanol, sauerkraut, soy sauce, or yeast ex- tract/supplements, fer- ments */**ampicillin/amoxicillin (CIDAL) preferred in VRE that is amp-susceptible. **Associated with treat- ment failure in bacteremia , incl line & endovascul- ar infections.	GOOD Myrianthefs et al. Serum and CSF concentrations of linezolid in neurosurgery patients. AAC 2016; 50(12): 3971-6.
tedizolid \$\$\$\$ STATIC	All Gram + cocci incl. ** VRE, Amp- resistant* Entero- coccus, MRSA/VRSA Binds 50s riboso- mal subunit	SSTI	IV=PO (bioequivalent) 6 days tedizolid Qdaily = 10 days linezolid BID = higher lipid solubility/higher tissue levels Nausea/headache/diarrhea Lower thrombocytopenia than linezolid; similar neuropathic events; no longer term data Serotonin syndrome: Avoid high tyramine food/drink (> 100mg tyramine per meal). E.g. aged cheeses, dried/processed meats, ethanol, sauerkraut, soy sauce, or yeast ex- tract/supplements, fer- ments	NO DATA – suspect similar to linezolid

Colistin/Polymixin B (Reserved for multi-drug resistant organisms - MDRO)

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
colistin polymixin B colistimethate \$\$\$ CIDAL	Gram - including Pseudomonas, Acinetobacter membrane disruption, binds lipopolysaccharide (LPS)/ Gram - endotoxin	Intraabdominal infections UTI/GU infections Pneumonia/ Hospital-associated respiratory infections Potent anti-LPS binding/ neutralizing activity	IV/Aerosol 30% Nephrotoxicity! Peripheral/ Optic neuropathies Neuromuscular blockade (may exacerbate myasthenia gravis & paralytic agents)	YES

Rifamycins

DRUG	COVERAGE	USES	TOXICITY/MISC	CSF
rifampin \$-\$\$ Only rifampin is discussed here, in context of use outside of mycobacterial infections CIDAL	Very broad, incl GPC/GNR, mycobacteria; use is <i>condition-specific</i> RAPID RESISTANCE if given alone – <i>Use in combinations</i> Inhibits DNA-dependent RNA polymerase	Only used alone as prophylaxis against Neisseria meningitidis (2 days), Hemophilus influenza b (4 days) in contacts/nasal carriage Combination treatment in serious S. aureus, Streptococcal infections Combination treatment of Legionella, Anthrax, Brucella, Bartonella, Anaplasma, Ehrlichia Combination treatment of tuberculous and non-tuberculous Mycobacteria	IV/PO Red urine, sweat, tears, saliva – hold soft contact use Nausea, abd pain Hepatotoxicity (avoid ethanol & hepatotoxins), hyper-bilirubinemia Type I & Flu-like hypersensitivity Autoimmune reactions Many drug interactions – always check an updated reference	YES

References:

<http://webedition.sanfordguide.com/>

www.drugs.com

www.emedicine.medscape.com

www.epocrates.com

www.micromedix.com

Acknowledgment:

Gratitude for support of my teaching efforts is owed to my alma mater, the University of South Florida Morsani College of Medicine Division of Infectious Diseases and International Medicine, and my distinguished professors there. I remain honored and very humbled to call them my colleagues these several years. I strive always to measure up to their high standards.

The opinions and information presented in any of my teaching materials, in print or electronically, remain my own intellectual property, and do not reflect the opinions or representations of any employer(s) or professional affiliates of which I am a part, past or present.

Thank YOU, dear Colleague, for your dedication to the Art and Science of Medicine. I hope that you find this tool of help in your care of the VIP at the center of our efforts:

The Patient.

Dr. G