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

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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, & oral anaerobes--those above the diaphragm. ONLY ceftazidime/cefepime/ceftolozane cover Pseudomonas..

3. Aztreonam, a monobactam, covers <u>ONLY Gram –s</u>, incl. Pseudomonas. Reserve for beta lactam-allergic patients.

4. Aminoglycosides generally cover Gram +s (except tobramycin) & Gram –s, NO anaerobes, some Mycobacteria. Increasing resistance. Best in combination with beta lactams unless uncomplicated cystitis/pyelonephritis, but most use is declining due to nephrotoxicity & dual beta lactam options.

5. Quinolones cover Gram -s best (<u>except moxifloxacin/delafloxacin</u>, which are "respiratory quinolones" most active against Gram +s & anaerobes), some Mycobacteria.

6. Trimethoprim/sulfamethoxazole (TMP/SMX) generally covers MSSA/MRSA, Nocardia, Listeria, Pneumocystis, most Gram –s except Pseudomonas. Probably *OK* for Streptococcus pyogenes/agalactiae cellulitis as well.

7. Clindamycin generally covers Gram +s, incl anaerobes except Clostridia, like anaerobic/microaerophilic Strep/Peptostrep, Actinomyces (better for infections <u>above the diaphragm</u>).

8. Metronidazole covers *gut* anaerobes like Bacteroides, Prevotella, Clostridia; +/-Peptostrep (infections *below* the diaphragm).

9. Carbapenems in general are very broad & among our "last line" beta lactams. 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/penicillin/penicillin, which should be considered bacterioSTATIC. Synergistic combination therapy is preferred for these infections, such as amp + gen-tamicin. Amp + ceftriaxone/ceftaroline is increasingly preferred due to nephrotoxicity with gent & gent re-sistance. The combination of these beta lactams binds more PBPs that ampicillin alone & lowers the ampicillin MIC. In addition, know that vancomycin is bactericidal for most GPCs *except Enterococcus* (bacterioSTATIC).

Shameless plug:

Visit <u>www.gompfsidpearls.net</u> 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 <u>work-up</u>, 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.
 - $\succ \qquad \underline{\text{Look for non-infectious cause}} \rightarrow \text{treat it.}$
 - ➤ 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; <u>stop</u> 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.
 - o 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 <u>data</u> 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; 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 ORALCONTRACEPTIVES! Tell female patients to add barrier contraception until the next new pill pack after finishing antibiotics.

Antifungal coverage in general:

fluconazole = Cryptococcus, Coccidioides, dermatophytes, Candida EXCEPT Candida krusei/auris/some glabrata and all molds; high CSF/urine levels

itraconazole = Candida, Histoplasma, Crypto, Cocci, Aspergillus, Sporothrix, Paracocci, Talaromyces voriconazole = Candida, Histo, Crypto, Aspergillus (except a few rare species, Fusarium, NOT initially for Mucor/Rhizopus, but OK as step-down after ampho B/source control; good CSF/poor urine levels posaconazole = same as vori, + Mucor/Rhizopus; variable CSF levels

isavuconazonium 372mg (= isavuconazole 200mg) = same as posa, some Mucor, INFERIOR to caspofungin for candidemia; few drug interactions; poor CSF/urine levels

enchinocandins (caspofungin/micafungin/anidulafungin) = Candida incl C. auris, Aspergillus, SOME Cryptococcus, NOT Fusarium/Mucor/Rhizopus/Trichosporon, NOT Histo/Blasto/Coccidioides; poor levels in CSF/urine/vitreous humor. Poor urine penetration, but does penetrate renal parenchyma, so may be effective in pyelonephritis, maybe cystitis. May allow you to avoid ampho B in the case of fluconazole resistant Candida.

amphotericin B = all, +/- Fusarium, NOT Candida lusitaniae (variable)/guillermondi/auris, NOT Scedosporium (Pseudallescheria), Lomentospora (Scedosporium); Aspergillus terreus flucytosine (5-flourocystosine) - increases penetration of above drugs, rapid resistance alone; good CSF/eve/urine levels

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/isuvaconazole (active metabolite of isuvaconazonium); NOT other azoles/enchinocandins

BacteriCIDAL vs. BacterioSTATIC

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

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.

Extended Spectrum Beta Lactamase (ESBL)-Producing Gram Negative Bacilli: It's complicated & evolving as fast as these organisms!

SPICE/SPACE/KEC are mnemonics for bacteria that *may* either have *intrinsic* (chromosome-based, aka "constitutive") and/or *inducible* beta lactamases (chromosome- OR mobile genetic element/plasmid-mediated beta lactamases). These organisms may demonstrate variable resistance to commonly prescribed beta lactams and may require carbapenem* treatment. In addition, *inducible* beta lactamases in KEC organisms may become reversibly de-repressed (or "switched on") upon 1- 7 days' exposure to a beta lactam and result in treatment failure. Cefepime is a weak AmpC inducer and stands up to hydrolysis by AmpC β-lactamase, so is generally a good choice if the organism is susceptible.

| | Strong Inducers | Weak Inducers | |
|-----------------|--|---|--|
| Good Substrates | Aminopenicillins, first-generation cephalosporins, cefoxitin, cefotetan | Ceftazidime, ceftriaxone, cefotaxime, piperacillin, ticarcillin, aztreonam | |
| Poor Substrates | Imipenem | Cefepime, meropenem | |

TL/DR: If suspecting severe sepsis, use ceftolozane-taz, ceftaz-avibactam, meropenem, or piperacillintaz if MIC <16mg/L**; otherwise, can deescalate to cefepime if MIC \leq 1 mg/L if stabilized, off pressors, etc. Some beta lactams are weak inducers of

KEC(Y) – Grouped by Family Enterobacteriaceae with moderate-high risk of *inducible*, chromosomal beta-lactamase (AmpC, Group C, cephalosporinase)—KNOW that these may trick you by testing S to ceftriaxone/ceftazidime→ treatment with cephalosporins may *induce* R. 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.

[Aside: Klebsiella pneumoniae & some E.coli often acquire a non-inducible, plasmid-based Group A beta lactamase (TEM) with a narrow spectrum against ampicillin & 1st-gen cephalosporins. Just be aware that this pattern comes up often in susceptibility testing & does not confirm or rule out an inducible ESBL.]

Klebsiella aerogenes (formerly Enterobacter) Enterobacter cloacae Citrobacter freundii Yersinia enterocolitica

SPACE/SPICE – Treat based on susceptibility report and severity. [These species' ampC mutations are *usually not inducible*, so these DO test R to ceftriaxone/ceftazidime, so this mnemonic has fallen out of favor.] Consider: Severity of infection (bacteremia, meningitis), high inoculum/source control (UTI/pyelonephritis vs endocarditis), presence of biofilm (infected device), & MICs to cefepime (≤ 1 mg/L) & pip-tazobactam (≤ 16 mg/L)** can help guide whether to use cefepime, pip-taz, or a carbapenem. Quinolones (bactericidal), trimethoprim-sulfamethoxazole (TMP-SMX), tetracyclines, and aminoglycosides are options if susceptible.

Presence of **chromosomal ("constitutive", "intrinsic") or mobile genetic element**-acquired (**plasmid**mediated) beta lactamases. Serratia/Salmonella/Shigella Proteus (non-P. mirabilis/"Indole +")/Providencia Pseudomonas Acinetobacter baumanii complex Citrobacter species Enterobacter complex (Yersinia enterocolitica – not enough data)

E. coli and Klebsiella are the most common ESBL producers, so many labs screen those isolates if MIC for ceftazidime is >/= 2 mg/L. Remember that **Klebsiella** almost all have a constitutive chromosomebased beta lactamase (usually SHV-1) & test R to ampicillin/ticarcillin.

*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 articles gloss over this.

**<u>MERINO Trial 2018</u> suggested higher 30-day all-cause mortality with pip-taz vs meropenem, but <u>other</u> <u>trials</u> and <u>post-hoc analysis of MERINO</u> have suggested MIC > 16mg/L to pip-taz is responsible for the difference. If MIC is low, piperacillin-tazo may be acceptable as "carbapenem-sparing" in infections with source control (e.g. pyelonephritis without obstruction). See also <u>Impact of the MIC of Piperacillin-</u> Tazobactam on the Outcome of Patients with Bacteremia Due to Extended-Spectrum-Lactamase Producing Escherichia coli, 2013; Association Between Minimum Inhibitory Concentration, Beta-lactamase Genes and Mortality for Patients Treated With Piperacillin/Tazobactam or Meropenem From the MERINO Study 10/27/2020; IDSA Guidance on the Treatment of Antimicrobial-Resistant Gram-Negative Infections: Version 2.0, 3/31/2022.)

Carbapenem (CP) Resistant Gram Negative Bacilli:

(Carbapenem attached to the beta lactam ring is what preserves activity against ESBL.) Affects mainly **Klebsiella pneumoniae**, **Pseudomonas aeruginosa and Acinetobacter baumannii (CRAB)**. May possess intrinsic vs mobile genetic element carbapenemase-encoding genes that inactivate enzymes, induce efflux pumps or limit porin channels. Carbapenems are not necessarily interchangeable.

Antibiotics for Resistant Gram + Cocci

- Vancomycin, teichoplanin (Europe) cover all but vancomycin-resistant Gram +s.
- Vancomycin is bacterioSTATIC against Enterococcus.
- Dalbavancin, oritavancin, televancin 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.

Pseudomonas, Acinetobacter, Stenotrophomonas vs other GNR

- These organisms are obligate aerobes:
 - Pearl: If a GNR in blood culture shows anaerobic growth, no need to cover for Pseudomonas!
- Acinetobacter spp. are often intrinsically resistant to multiple antibiotics including ertapenem and may be variably susceptible to meropenem or imipenem (susceptibility to one carbapenem does not predict susceptibility to another!)
- Stenotrophomonas are intrinsically resistant to *carbapenems*; generally most susceptible to quinolones, sulfas, and tetracyclines.
- These species are not of the family Enterobacteriaceae & often affect patients whose microbiome has been altered by extensive antibiotic/healthcare exposure.

Which antibiotics are bacterioSTATIC?

"In sepsis, restore Volume with a Liter of STAT NML (normal) Saline."

Vancomycin in Enterococcus – cidal for all other GPCs Linezolid/Lefamulin Sulfas/trimethoprim (*especially static against Stenotrophomonas*) Tetracyclines/Tigecycline Nitrofurantoin (cidal in cystitis if concentration > 2x MIC) "MLS antibiotic group" – clindamycin, macrolides (Note: Streptogramins are bactericidal)

Everything else is bactericidal.

Note bene: Clindamycin & linezolid are best studied as *adjuncts* for Staph or Strep toxic shock, severe streptococcal cellulitis or suspected necrotizing infection; *halt protein synthesis*—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). Other drugs whose mechanism of action is disruption

of protein synthesis: doxycycline/minocycline/tigecycline.

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 outcomes 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 pharmaco-kinetics 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 & and 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/CEK (d). And vancomycin "MIC creep" with S. aureus bacteremia—as MIC "creeps" above 1, so does treatment failure. Etc, etc.

Pharmacokinetics & Pharmacodynamics (PK/PD)

Important determinants of antibiotic dosing for maximal effect include time-dependent vs concentration-dependent activity, volume of distribution/Vd (obesity, serum albumin), and route of elimination (renal, hepatic [cytochrome P450 enzymes], GI).

Time-Dependent antibiotics depend on duration of time that free antibiotic levels remain in tissue above MIC, no post-antibiotic effect. Doses must be given at regular intervals to maintain activity. Low albumin levels may lower free antibiotic levels by allowing increased free antibiotic to be cleared before it gets into tissues (highly protein-bound antibiotics may need higher doses, esp with less susceptible organisms).

• All beta lactams, monobactam, carbapenems

Concentration-dependent antibiotics act by achieving a high initial concentration & have a persistent "post-antibiotic effect"—i.e. bacterial growth is inhibited or "stunned" for a long time even after concentrations are no longer above MIC. Allows for extended dose intervals. Low albumin levels may lower free antibiotic levels by allowing increased free antibiotic to be cleared before it gets into tissues. At higher doses, highly protein-bound antibiotics may maintain adequate serum concentration if enough free antibiotic is available; again, may need higher dose).

- Aminoglycosides
- Daptomycin
- Metronidazole
- Quinolones (mostly)

Combination of Time above MIC + Post-antibiotic Effect

- MLS group macrolides, clindamycin, tetracyclines
- Linezolid
- Vancomycin dosing is optimized by achieving certain concentration for certain period of time, or "area under the curve" (AUC) -based dosing

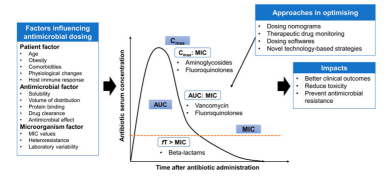


Table 1. Protein binding of antibacterials commonly used in critically ill patients and of antibacterials in development (all protein binding data have been adapted from Donnelly et al.^[26] and MIMS Australia^[27]). We have also included data on antilungal agents for the reader's reference

| Highly bound (>70%) | Moderately bound (70-30%) | Minimally bound (<30%) |
|--|---------------------------------------|------------------------------------|
| Amphotericin B (90%) | Azithromycin (7–51%) | Amikacin (0-11%) |
| Anidulafungin (>99%) | Aztreonam (60%) | Amoxicillin (17-20%) |
| Caspofungin (97%) | Cefotaxime (40%) | Ampicillin (15-25%) |
| Cefazolin (75-85%) | Cefuroxime (33-50%) | Cefepime (16-19%) |
| Cefonicid (98%) | Cephalothin (55-75%) | Ceftazidime (17%) |
| Cefoperazone (90%) | Ciprofloxacin (20-40%) | Ceftobiprole (22%) |
| Cefoxitin (80-50%) | Clarithromycin (42-50%) | Cefpirome (9%) |
| Ceftriaxone (85–95%) | Chloramphenicol (60%) | Colistin (<10%) |
| Clindamycin (90% bound to a1-acid glycoprotein) | Levofloxacin (50%) | Doripenem (8%) |
| Cloxacillin (94%) | Linezolid (31%) | Ethambutol (20-30%) |
| Dalbavancin (93%) | Moxifloxacin (30-50%) | Fluconazole (11-12%) |
| Daptomycin (90-93%, 30% to a1-acid glycoprotein) | Nitrofurantoin (40%) | Fosfomycin (0%) |
| Dicloxacillin (97%) | Benzylpenicillin [penicillin-G] (65%) | Gentamycin (<30%) |
| Doxycycline (93%) | Piperacillin (30%) | Imipenem (20%) |
| Ertapenem (85–95%) | Sulfamethoxazole (68%) | Isoniazide (0-10%) |
| Erythromycin (73–81%) | Ticarcillin (55%) | Meropenem (2%) |
| Faropenem (96–99%) | Trimethoprim (45%) | Metronidazole (<20%) |
| Flucloxacillin (95%) | Vancomycin (30-60%) | Norfloxacin (10-15%) |
| Fusidic acid (95–97%) | Voriconazole (58%) | Polymyxin B (<10%) |
| Iclaprim (93%) | | Quinupristin/dalfopristin (11-26%) |
| Itraconazole (99.8%) | | Tobramycin (<30%) |
| Lincomycin (80–90%) | | |
| Minocycline (75%) | | |
| Nafcillin (90%) | | |
| Oxacillin (93%) | | |
| Posaconazole (>97%) | | |
| Rifampicin [rifampin] (80%) | | |
| Sulfisoxazole (92%) | | |
| Teicoplanin (90–95%) | | |
| Telavancin (92–94%) | | |
| Tigecycline (71-89%) | | |

Synergistic IV Antibiotic Combinations against Specific Pathogens

Rule #1 – Don't wing it! Synergy is not that common. Some combos may be antagonistic, so combine with care (i.e. Use only combos supported by good literature).

Cell wall active agents + cell wall active agents

- serious GPC/GNR infections
 - Beta lactams + aminoglycosides
 - Beta lactam increases cell wall penetration by AG→AG disrupts cell wall (superoxides) & cell wall protein production

"drugs that are strongly

protein bound (≥90%) may differ markedly from minimally protein-bound drugs in terms of tissue penetration and half-life The concentration of albumin and other protein may be altered under stress, surgery, pregnancy, and liver or kidney diseases; in such circumstances, monitoring free drug is more useful for strongly proteinbound drugs. For these patients, free drug levels correlate better with clinical picture than traditionally monitored total drug level [4]." https://www.sciencedirect.com/science/arti-cle/pii/B97801280202580

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V ceftaroline (<20%) PO cefpodoxime (18-30%; 30% urine excreted in urine; UTI indication) PO cefadroxil (20%; 90% excreted unchanged in urine; UTI indication)

• serious Enterococcus faecalis/faecium infections

• Ampicillin + cephalosporins/ertapenem

- binds more PBPs that ampicillin alone & lowers ampicillin MIC
- now preferred to avoid nephrotoxicity of amp + gentamicin combo
- MSSA/MRSA persistent bacteremia/endocarditis salvage
 - o daptomycin + cephalosporins (esp ceftaroline)/ertapenem
 - cefazolin/ceftarolin+ ertapenem (meropenem can be safely used with ceftaro if SA & GNR infection!)
 - binds more PBPs
- KPC carbapenemase-producing GNR (blakPC mutation)
 - ertapenem + meropenem
 - o erta acts as ("suicide antibiotic" preferential affinity for car-
 - bapenemase—erta resistance is often alerts Micro to test for KPC)
- Carbapenem-R Acinetobacter (CRAB)
 - **Meropenem + sulbactam** (as ampicillin-sulbactam—high dose 6-9g/d for most effective sulbactam dose) or sulbactam-durlobactam)
 - Polymixin B or collistin + carbapenem, tigecycline, or amp-sulbactam – more to come!
- Invasive Aspergillosis, when avoiding nephrotoxicity
 - Echinocandin + azole antifungal in hematologic malignancy, combination may reduce mortality vs azole alone [Ann Intern Med. 2015;162:81-89. doi:10.7326/M13-2508]

Drugs that penetrate the prostate:

- quinolones with Gram negative coverage
- trimethoprim-sulfamethoxazole
- doxycycline
- fosfomycin
- azithromycin
- beta lactams

THE CLASSES (not an exhaustive list)

| DRUG | ams are CIDAL, good tissue p COVERAGE | USES | TOXICITY | Cerebral |
|--|---|---|--|--------------------------|
| | | | | Spinal Fluid (CSF) |
| Natural Penicil- lins (PCN) \$ IV penicillin G (benzylpenicillin) PO penicillin VK CIDAL | Group A Strep (no re- sistance) Strep viridans Neisseria Capnocytophagia Actinomyces Fusobacterium Clostridia perfringens/te- tani Pasteurella Treponema/ Leptospirosis NOT MSSA (resistant) | Skin/soft tissue (SST) Oral/den- tal infec- tions | Hypersensitivity Stevens Johnson Interstitial nephritis Seizures (if high level) Bone marrow sup- press-ion C.difficile | |
| Penicillinase-re- sistant PCN \$ IV methicillin nafcillin oxacillin PO dicloxacillin CIDAL | MSSA MS S.epidermidis/other coagulase-negative Staphylococci S. lugdunensis Strep pyogenes (grp A) Strep agalactiae (grp B) Strep grp C,F,G Strep anginosus grp Strep pneumoniae (if susceptible) | SSTI Bactere- mia Endovas- cular in- fections | Fever Hypersensitivity Stevens Johnson Interstitial nephritis Seizures (if high level) Bone marrow sup- press-ion Hepatotoxicity C.difficile | YES if inflamed |
| AminoPCN \$\$ IV ampicillin amp/sulbactam PO amoxicillin amoxicillin amox/clavulanate CIDAL | Add to the above: Listeria MSSA Most Pneumococcus Proteus Hemophilus influ. (beta lactamase negative) Salmonella/Shigella Anaerobes <i>Klebsiella are intrinsi- cally resistant to</i> <i>amp/amox</i> (clavula- nate/sulbactam don't add much activity) Note: High-dose amp- sulbactam may used as a source of <i>sulbactam</i> in treating MDR Acineto- bacter | Otitis me- dia Sinusitis SST Meningitis in elderly | Above | |

Penicillins - beta lactams are CIDAL, good tissue penetration

| CarboxyPCN \$\$ IV ticarcillin/clay (Eu- | Adds to the above: Pseudomonas Enterobacteriaceae* Stenotrophomonas (ticar) Gut anaerobes MSSA | Adds to above: Gut/ surgical infections Nosocom- ial pneu- | Above |
|---|--|--|-------|
| | (ticar) Gut anaerobes MSSA Pip & Pip/tazo more po- tent for GNRs & more resistant to AmpC/ESBLs (See "SPICE" above) *Klebsiella is intrinsically | infections | |
| | R to ticarcillin | | |

Cephalosporins – Think of progressive broadening of spectrum from Gram + to Gram - with each generation. Beta lactams are CIDAL. [\odot = can be dosed 3 times weekly in dialysis patients] ***CROSS-ALLERGY (same side chains) between aztreonam – ceftazidime – cefiderocol***

| DRUG | COVERAGE | USES | TOXICITY | CSF |
|---|--|---|--|--|
| 1st Generation \$\$ IV/IM cefazolin ☺ | GPC/MSSA/strepto- cocci, E. coli, Pro- teus, some Klebsiella (increasingly ampC+/ESBL) NOT Enterococci | SSTI Uncompli- cated/Non-diabetic osteomyelitis PreOP prophylaxis | Hypersensitivity Bone marrow suppression Diarrhea | |
| PO cephalexin cefadroxil CIDAL | | | C.difficile | POOR |
| 2 nd Generation \$\$ PO cefuroxime cefaclor IV cefuroxime cefamycins: cefoxitin cefotetan | Streptococci, uncom- plicated MSSA Pneumococcus Neisseria Some GNR except Pseudomonas Cefamycins are the only ones that relia- bly cover anaer- obes NOT Enterococci | Community ac- quired pneumonia (CAP) meningitis OM/sinusitis Gonorrhea | Hypersensitivity RASH/Stevens Johnson w/ cefaclor High INR/PT w/ cefoxitin/ cefotetan Bone marrow suppression C.difficile | YES if strongly in- flamed ceftriaxone >90% protein bound, low BBB penetration |

| 3 rd Generation \$\$ PO cefpodoxime IV ceftriaxone cefotaxime CIDAL | Above; covers viri- dans streptococci, pneumococcus, but reduced PBP binding in MSSA | Meningitis CAP Most community- acquired infections Gonorrhea Pyelonephritis Best <i>PO</i> ceph for GU is cefpodox- ime (20-30% pro- tein-bound, vs PO cefdinir, which is poorly excreted in urine & up to 70% protein-bound) | Above Ceftriaxone: <i>Pseudo-choleli- thiasis</i> (biliary sludge) | |
|---|---|--|--|--|
| 4 th Genera- tion/Anti-pseu- domonal \$\$ IV ceftazidime © cefepime © CIDAL | Above, plus Pseudo- monas More resistant to beta lac- tamases/ESBLs (See "SPICE" above) be- cause it is not porin- dependent NOT Enterococci | Above, plus neu- tropenic fever | Above Cefepime: Encephalopa- thy, non-convul- sive status epi- lepticus | |
| 5 th Genera- tion/Anti-MRSA \$\$\$ IV ceftaroline CIDAL | Similar to 3 rd genera- tion, plus MRSA, VISA/VRSA/VRE faecalis (NOT E. fae- cium), pneumococ- cus, beta-lactamase + H.flu, Moraxella Listeria NO Pseudomonas | Complicated SSTI, CAP (NOT MRSA- insufficient data) | Above | |

| Advanced-gen- eration/Anti- pseudomonal \$\$\$ IV ceftolozane-tazo- bactam ceftazidime-avi- bactam CIDAL | Viridans streptococci NOT Enterococci or Staphylococci ceftoloz-taz covers GNRs incl Pseudo- monas, ESBLs, <i>some</i> car- bapenemase-produc- ing P. aeruginosa (CRPA), NOT KPC+ ceftaz-avi covers ESBL & KPC+ car- bapenemase (1 st line agent) ceftaz-avi covers GNRs incl Pseudo- monas, adds cover- age for ceftaz-R, ESBLs, some ampC- R, <i>some</i> car- bapenemases (NOT metallobetalac- tamase) | Complicated UTI/pyelo Complicated in- traabdominal in- fection ceftaz-avi adds HAP | Above Nausea, diar- rhea, headache, fever, renal in- sufficiency (ceftolaz-taz) | celtiz(dine - CSF, Itel Inflamed (NOT avibed am) - CSF, Itel Inflamed (NOT avibed am) - CSF, Inflamed (NOT avibed am) CSF, Inflamed (avibed am 97%) 59% advint bound 30% advint bound |
|---|--|--|---|---|
| Cefiderocol IV Similar side chains of both cefep & ceftaz Trojan horse si- derophore: Che- lates Fe++, so drug is actively transported with Fe++ via sidero- phore channels CIDAL | Reliably covers XDR/carbapenem-R: Metallobetalacta- mase producers (MBL) Klebsiella pneumo- niae (KPC+) Pseudomonas aeru- ginosa (CRPA) Enterobacteriaceae (CRE) Acinetobacter bau- manii (CRAB) Stenotrophomonas Burkholderia cepa- cian NO GPC! | Complicated UTI/pyelo HAP | Hypersensitivity Bone marrow suppression Diarrhea C.difficile <u>Unclear if higher</u> <u>all-cause mor-</u> <u>tality in critically</u> <u>ill with carba-R</u> <u>GNB (CREDI- <u>BLE-CR trial)</u></u> | - CSF POOR - 58% protein bound - 60-70% renal excretion, unchanged |

Monobactam

| DRUG | COVERAGE | USES | TOXICITY | CSF |
|-----------|--|---|---------------------------------------|-----------------------------|
| aztreonam | ONLY GNRs, incl Pseudomonas | <u>GNR</u> infections; NOT a replacement | Low | YES if in- flamed |
| \$\$ | · | for all aminoglyco- | Good alternative | [Modal J et al. |
| IV | Covers metal- lobetalactamase | side uses (no syn- ergy for GPC, NO | for beta lactam al- lergies EXCEPT | AAC. 1986;29:281- 3.] |
| CIDAL | carbapenemases, but not ESBL (re- sistance usually occurs together)— combination of az- treonam + ceftaz- avibactam might be used in sal- vage cases (<u>CID</u> <u>2021;72:1871</u>) | Enterococcal cover- age) | with ceftazidime, cefiderocol | 3.j |

| DRUG | imes weekly in dialysis pat | USES | ΤΟΧΙCITY | CSF |
|--|--|--|--|-----|
| \$\$\$ IV imipenem- cilastin meropenem- vaboractam imipenem-cilastin- relebactam CIDAL | GPCs EXCEPT MRSA GNRs EXCEPT Sten- otrophomo- nas/Burkholderia ESBL+ & "SPICE" GNRs Anaerobes (incl Cuti- bacterium) Listeria Pneumococcus Nocardia asteroides (NOT brasiliensis) Legionella Mycobacterium avium Enterococcus (NOT E. faecium) mero-vaboractam adds car- bapenemase+ Klebsiella pneumonia (KPC), class A car- bap-R Enterobactera- ciae (NOT metallobetalac- tamase/OXA carbap- R, NOT carbap-R Pseudomonas/Aci- netobacter) Relebactam is not ac- tive against Morganel- lacea group (Morga- nella, Proteus, Provi- | Resistant GNR in- fections Serious gut infec- tions Necrotizing pan- creatitis | IV/IM Hypersensitivity (~10% cross-al- lergy with beta lactams) Seizures with imipenem (if renal insufficiency or high levels used) Candida over- growth/ infections C.difficile | YES |
| \$\$\$ | dentia) Above, possibly lower | Above | Above | - |
| IV doripenem | MICs to Pseudomo- nas & Acinetobacter | Higher mortality than imipenem in VAP | | |
| CIDAL | | | | |
| \$\$\$ IV/IM ertapenem ☺ CIDAL | Above, without Pseu- domonas coverage | Postpartum uter- ine infections Postsurgical Ab- dominal infections (not Pseudomo- nas) | Above Encephalopathy | |

Carbapenems (Reserved for Multidrug Resistant Organisms – MDRO)

Aminoglycosides

| DRUG | COVERAGE | USES | TOXICITY/ MISC | CSF |
|---|--|--|---|---------------------------------|
| \$-\$\$\$ IV gentamicin streptomycin spectinomycin tobramycin amikacin Nebulized tobramycin liposomal amika- cin CIDAL | Gent: VSE/VRE/variable Staph, GNRs incl Pseudomonas Tobra/Amik: GNRs incl Pseu- domonas Amik: TB, non-TB Mycobacteria Strepto -Yersinia -MDR Mtb Gent/Strepto -Tularemia Spectino -Gonorrhea NO coverage for: Acinetobac- ter/Stenotropho- monas Anaerobes Pneumococcus | Synergy with beta lactams for GPC/Pseud-omo- nas infections Usually not used alone except for UTIs | IV/Aerosol Acute tubular ne- crosis (reversible) Cochlear toxicity (genetic predispo- sition) Vestibular toxicity (irreversible) When possible: -stop after 3-5 d -use once-daily dosing -avoid in elderly Liposomal amik – hypersensitivity pneumonitis, hemoptysis, bron- chospasm, exac- erbation of lung disease Neuromuscular blockade (may exacerbate myas- thenia gravis & paralytic agents) | NO or UNKNOWN (use intrathecal) |
| \$\$ PO only, in U.S. fosfomycin CIDAL | Enterococcus Staph. aureus GNRs | Simple cystitis in women Off-label q3days for complicated or MDR GNRs, VRE if susceptible <i>Prostatitis</i> | PO only Above, significant diarrhea | |
| \$\$ IV plazomicin CIDAL | GNRs incl MDR/KPC/meta- lobetalact/CRE GNRs,variable Pseudomonas (use only if known susceptible), NOT Steno, Acineto- bacter | Complicated UTI/pyelonephritis | IV only Above Limited data | |

Sulfonamides/Sulfas (INTRACELLULAR ACTIVITY)

| DRUG | COVERAGE | USES | TOXICITY/MISC | CSF |
|--|---|---|--|-----|
| \$ IV/PO trimethoprim-sulfa- methoxazole (co-trimoxazole) STATIC for Staph, may be CIDAL for others | Staph. aureus (incl MRSA) (NOT Enteroco- ccus, Pseudo- monas, Aci- netobacter) Legionella Stenotropho- monas Listeria Pneumocystis Nocardia Burkholderia cepacia Moraxella Yersinia Francisella tula- rensis Toxoplasma Atypical myco- bacteria (M. marinum) Some common coliforms | Cystitis/pyelone- phritis <i>Prostatitis</i> MRSA SSTI Specific agents at left <i>Dosing de- pends on what</i> you're treating | IV/PO RASH/Stevens Johnson Nausea Fever Bone marrow sup- pression Hemolysis (if G6PD deficient) Hepatotoxicity <i>Elevated creatinine</i> <i>despite normal</i> <i>GFR</i> (competes with Cr for tubular secretion <i>Hyperkalemia</i> (blocks Na+ chan- nels & thus K+ ex- cretion) Kernicterus in neo- nates C.difficile Sun sensitivity | YES |

Pleuromutilins (INTRACELLULAR ACTIVITY)

| DRUG | COVERAGE | USES | TOXICITY/MISC | CSF |
|---|------------------------------|--------------------------------------|----------------------------------|--------------------------|
| \$\$\$ | S. pneumoniae MSSA, ?MRSA | CAP/community- acquired pneumo- | Nausea, hepato- toxicity, CNS | UNKNOWN 70% fecal ex- |
| IV/PO | Hemophilus in- fluenzae | nia | Hypokalemia | cretion |
| Lefamulin** | Mycoplasma | ** Bacteriostatic - | | |
| Protein synthesis in- hibitor – multiple | pneumoniae Chlamydia | be aware of this when empirically | Prolonged QTc | |
| binding sites to ribo- some, higher re- | pneumoniae Legionella | treating serious in- fections. | Teratogenicity | |
| sistance barrier | pneumophila | | C. difficile | |
| STATIC | | | | |
| | | | | |

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

| DRUG | COVERAGE | USES | TOXICITY/MISC | CSF |
|----------------|---|-------------------|---------------------------------|---|
| Macrolides | Pneumococcus IF | LRTI/ | IV/PO | |
| \$\$ | local resistance is | bronchitis | | |
| | <25% | Sinusitis | Nausea/ | |
| 11/100 | MSStaph. aureus | Dental/ | vomiting | |
| IV/PO | (not MRSA) | oral infections | Abdominal | |
| Erythromycin | Legionella | | cramps/ | |
| | Listeria | Prostatitis | diarrhea (Lowest | |
| PO | Hemophilus | | with Azithro) | |
| clarithromycin | Moraxella | Atypical mycobac- | 0 -1145 - 11- | D. |
| azithromycin | Mycoplasma | teria | C.difficile | ydi |
| , | Pertussis | (incl MAC prophy | Ot - t | am |
| | | in HIV) | Ototoxicity with chronic use | Ч. |
| STATIC | Neisseria menin- | | chionic use | a |
| | gitis | | Rare association | hel |
| | - | | with cardiovascu- | gio |
| | | | lar mortality with | Ļ |
| | Chlamydia/He- | | QTc prolongation, | ma |
| | mophylus ducreyi | | low Mg++/K+. | las |
| | | | 0 | do |
| | Campylobacter | | Interactions: | Ψ, |
| | Helicobacter pylori | | Ery/Clari induce | for |
| | Salmonella/Shi- | | P450! | fer |
| | gella | | | bet |
| | | | Neuromuscular | 1 |
| | Deuteurelle | | blockade with Ery | late |
| | Bartonella Barrolia burgdor | | (may exacerbate | Adequate – better for Mycoplasma, Legionella, Chlamydia |
| | Borrelia burgdor- pheri (<i>Lyme</i> dis- | | myasthenia gravis | PA |
| | ease | | & paralytic | |
| | Babesia microti | | agents) | |
| | Babesia miorod | | | |
| | | | | |
| | Actinomyces | | | |
| | Atypical mycobac- | | | |
| | teria | | | |
| | | | | |
| 1 | 1 | 1 | 1 | 1 |

| | | 0 00Ti | 11//20 | |
|--------------|---------------------|----------------------|--------------------------|--------------------------------------|
| Lincosamides | CIDAL for Group | Severe SSTI, ne- | IV/PO | |
| | A streptococcus, | crotizing fasciitis, | C.difficile!! (>30% | |
| \$\$ | MSStaph. aureus | MRSA | develop it on a | |
| | (MRSA, but | <i></i> | week of clinda) | |
| IV/PO | watch for induci- | "Infections above | | |
| | ble erm re- | the diaphragm" | Dysgeusia | |
| clindamycin | sistance. Clue is | | | |
| | resistance to | Head and | Deeb favor ee | |
| | erythromycin) | neck/dental infec- | Rash, fever, eo- | |
| STATIC | | tions | sinophilia | |
| | Pneumococcus | Lung abscess/ | Erythema multi- forme | |
| | | aspiration pneu- | torme | |
| | Inhibits toxic pro- | monia | | ≥ |
| | teins in severe | (tip: no teeth = no | Reversible neutro- | POOR except for Toxoplasmosis in HIV |
| | Strep A & S. au- | anaerobes) | penia/thrombocy- | . <u>L</u> |
| | reus/ | | topenia | sis. |
| | necrotizing | Bacterial vagi- | | or |
| | fasciitis. | nosis | Watch for hepati- | asr |
| | | | tis/obstructive | bld |
| | Oral anaerobes: | Babesiosis | jaundice, severe | DX(|
| | Gram + such as | | liver injury | Ĕ |
| | Peptostrepto-coc- | Toxoplasma in | | for |
| | cus, | HIV | Neuromuscular | pt |
| | Fusobacterium, | | blockade (may ex- | ce |
| | Prevotella, | | acerbate myas- | ex |
| | Actinomyces, & | | thenia gravis & | R |
| | Clostridial spp | | paralytic agents) | õ |
| | other than Clos- | | | A |
| | tridium difficile | | | |
| | | | | |
| | | | | |
| | Gram – such as | | | |
| | Bacteroides | | | |
| | (may not cover in | | | |
| | up to 25% of | | | |
| | cases or strains | | | |
| | with MIC $>/= 8$ | | | |
| | mcg/mL) | | | |
| | | | | |
| | Babesiosis | | | |
| | | | | |

Nitrofuran

| DRUG | COVERAGE | USES | TOXICITY/MISC | CNS |
|----------------------------|--|---|--------------------------------|-----|
| \$ PO nitrofurantoin | GNRs EXCEPT Pseudomonas, Proteus, and Enterococcus incl susceptible VRE | Cystitis Susceptible ESBL GNRs | PO only Nausea/ vomiting | |
| CIDAL | Multiple sites of ac- tion, inhibits syn- thesis of DNA, RNA, proteins, cell wall – higher re- sistance barrier than most antibiot- ics | ONLY reaches therapeutic level in URINE | C.difficile | NON |

Quinolones (Resistance is rising due to overuse; single-step mutation→ resistance may arise while on therapy; INTRACELLULAR ACTIVITY; high concentration in bone, prostate, CSF)

| DRUG | COVERAGE | USES | TOXICITY/MISC | CSF |
|-----------------------------------|---------------------------------|------------------------|--|----------------|
| "Gram negative" | GNRs including | Cystitis/pyelone- | IV/PO | |
| Quinolones | Pseudomonas | phritis Bractatitia | IV=PO (bioequiva- | |
| \$-\$\$ | Levo covers | Prostatitis | lent) | |
| Ψ-ΨΨ | pneumococcus | Intraabdominal in- | Dizziness/CNS | |
| IV/PO | & Stenotropho- | fections | Diarrhea | |
| ciprofloxacin | monas well | En dans stuitis | Hypo-/hyperglyce- | |
| levofloxacin | "Atypical" pneu- | Endometritis | mia Sun sensitivity | YES, HIGH DOSE |
| | monia: | Hospital-associ- | ourischistivity | ă |
| PO | Mycoplasma, | ated lung infec- | May exacerbate | ц Б |
| norfloxacin | Chlamydia, | tions | myasthenia gravis | Ξ |
| | Moraxella | Levo best for | & paralytic agents (inhibits GABA re- | ĔS |
| CIDAL | Some mycobac- | acute sinusitis/ | ceptors) | ~ |
| | teria/TB | CAP | | |
| | | Norfleyy UTL enty | May prolong QTc | |
| | | Norflox: UTI only | (watch for palpita- tions/syncope; | |
| | | | avoid if QTc > | |
| | | | 500msec, cardio- | |
| | | | myopathy) | |
| | | | Rare spontaneous | |
| | | | tendon rupture | |
| | | | (watch for pain at | |
| | | | tendon sites) | |
| | | | *Aortic dissection | |
| | | | association* | |
| | | | 0.1111111 | |
| "Gram positive | Pneumococcus, | CAP/community- | C.difficile IV/PO | |
| or Respiratory" | Streptococci, | associated respir- | IV=PO (bioequiva- | UNKNOWN |
| Quinolone | Staphylococcus | atory infections | lent) | |
| | (NOT MRSA) | Acute sinusitis | | |
| \$\$ | Legionella Gut anaerobes | Intraabdom-inal | Above >60% <i>liver</i> excre- | |
| PO | Gut anaciobes | infections | tion | |
| moxifloxacin | Atypical myco- | SSTI | | |
| monitoraciti | bac/TB | | | |
| | | | | |
| CIDAL | Strontococi | 0071 | IV/PO | |
| "Gram positive or Respiratory" | Streptococci, Staphylococcus | SSTI CAP/community- | IV/PO IV=PO (bioequiva- | UNKNOWN |
| Quinolone | MRSA | associated respir- | lent) | 0.11110111 |
| | Pseudomonas | atory infections | | |
| \$\$ | Legionella | | Above | |
| | Gut anaerobes | | | |
| IV/PO | Atypical myco- | | | |
| delafloxacin | bac/TB | | | |
| | | | | |
| CIDAL | | | | |

Nitroimidazole

| DRUG | COVERAGE | USES | TOXICITY/MISC | CSF |
|--------------------------------|---|--|---|-----|
| \$\$ IV/PO metronidazole | Gram – anaer- obes incl. Bac- teroides fragilis and all Clostridia Entamoeba coli | "Infections below the diaphragm" Intraabdominal ab- scess, peritonitis, diverticulitis, etc Endometritis/ | IV/PO Disulfiram-like re- action (vomiting) if ethanol con- sumed within 3 days of therapy | YES |
| CIDAL | | Bacterial vaginosis Clostridium difficile colitis | Aseptic meningi- tis/ neuropathies, | |
| | | Entamoeba liver abscess/ dysentery | rare | |
| | | NOT to be given alone for lung ab- scess/ENT infec- tions | | |

Tetracyclines/Glycylcycline (INTRACELLULAR ACTIVITY)

| DRUGCOVERAGEUSESTOX- ICITY/MISCCSFTetracyclinesMRSA/MSSA Pneumococcus Enterococci (incl VRE. faeca- lis/faecium) +/- GASAcne/rosacea SSTI CAP, esp under age 40IV/PO permanent teeth in childrenYES (n roborel syphilisIV/PO Minocycline DoxycyclineIis/faecium) +/- GAS E. coli Legionella N. meningitidis Hemophilus Moraxella Brucella ActinomycesAcne/rosacea CAP, esp under age 40 Dog/cat bite prophy as alterna- tive to amox/clav Tickborne dis- easesEsophageal ul- cerYES (n roborel teeth in childrenSTATICMoraxella Mycoplasma Chlamydia Listeria Brucella ActinomycesProstatitisPseudotumor cerebri, esp minocycline! (watch for head- ache) | lliosis, |
|---|----------|
| Pneumococcus SSTI Discoloration of permanent YES (n roborel \$ Pneumococcus SSTI Discoloration of permanent YES (n roborel IV/PO lis/faecium) Dog/cat bite bite bite Minocycline +/- GAS prophy as alterna- tive to amox/clav Esophageal ul- cer Esophageal ul- cer STATIC Hemophilus Moraxella Prostatitis Pseudotumor cerebri, esp Moraxella Prostatitis Prostatitis Pseudotumor cerebri, esp Brucella Brucella ache) | lliosis, |
| \$ Enterococci (incl VRE. faeca- lis/faecium) CAP, esp under age 40 permanent teeth in children roborel syphilis IV/PO lis/faecium) Dog/cat bite prophy as alterna- tive to amox/clav Esophageal ul- cer Esophageal ul- cer Esophageal ul- cer STATIC Hemophilus Moraxella Chlamydia Listeria Brucella Prostatitis Pseudotumor cerebri, esp minocycline! (watch for head- ache) | lliosis, |
| IV/PO VRE. faeca- age 40 teeth in children syphilis IV/PO lis/faecium) | |
| IV/PO Minocycline Doxycycline lis/faecium) +/- GAS Dog/cat bite prophy as alterna- tive to amox/clav Esophageal ul- cer STATIC Legionella Tickborne dis- eases Hepatotoxicity STATIC Moraxella Mycoplasma Chlamydia Listeria Prostatitis Pseudotumor cerebri, esp minocycline! Brucella Brucella ache) | >) |
| Minocycline Doxycycline +/- GAS prophy as alterna- tive to amox/clav Esophageal ul- cer STATIC Hemophilus Moraxella Chlamydia Listeria Brucella Prostatitis Pseudotumor cerebri, esp minocycline! | |
| Initicity clinic E. coli tive to amox/clav cer Doxycycline E. coli tive to amox/clav cer STATIC Hemophilus rickborne dis- eases Hepatotoxicity STATIC Hemophilus Prostatitis Pseudotumor cerebri, esp Chlamydia Listeria (watch for head- ache) | |
| Docysystem Legionella Tickborne dis- eases N. meningitidis eases STATIC Hemophilus Moraxella Prostatitis Mycoplasma Prostatitis Chlamydia minocycline! Listeria Gwatch for head- ache) | |
| STATIC Hemophilus Moraxella Mycoplasma Chlamydia Listeria Prostatitis Pseudotumor cerebri, esp minocycline! (watch for head- ache) | |
| Moraxella Prostatitis Pseudotumor Mycoplasma Prostatitis cerebri, esp Chlamydia minocycline! Listeria (watch for head- Brucella ache) | |
| Moraxella Prostatitis Pseudotumor Mycoplasma Prostatitis cerebri, esp Chlamydia minocycline! Listeria (watch for head- Brucella ache) | |
| Chlamydia minocycline! Listeria (watch for head- Brucella | |
| Listeria (watch for head- Brucella ache) | |
| Brucella àche) | |
| | |
| | |
| Borrelia burgdorferi Sun sensitivity | |
| Rickettsia | |
| Vibrio C.difficile | |
| Treponema | |
| Anaerobes: Fuso- Inhibit lipopol- | |
| bacterium, Cuti- ysa-ccharide-in- | |
| bacterium, Pepto- duced proin- | |
| streptococcus, flammatory | |
| Clostridium, some products | |
| Bacteroides fra- gilis/melano- | |
| genicus | |
| Glycylcycline Above, plus SSTI IV only | |
| Staph. epidermidis Intraabdo-inal in- Above UNKNO | OWN |
| \$\$ Enterococci fections | |
| Corynebacterium CAP/HAP 20% tige, 6.5% | |
| IV ESPL + vomiting | |
| U U LODE + Voliniting | |
| tigecycline >** E.coli/Klebs Prostatus eravacycline* (NOT KPC) Inhibit lipopoly- | |
| Storestonbornence | |
| IV/PO Acinete Severe C.difficile duced proin | |
| omadacycline Solmonollo Y alveolar, soft tis- | |
| B. fragilis/ | |
| sarecycline (acne anaerobes Poor bone/joint, | |
| Clostridia incl. CNS | |
| C.difficile Ampicillin/ | |
| ** Bacteriostatic - Allovicium | |
| NOT Pseudomo- nas, Burkholderia ube aware of this ferred in VRE | |
| when empirically that is amplify | |
| *erovoovolino oddo liteaung senious in- | |
| ESBL carban-R | |
| Asingtoheaster > Increased mor- | |
| tality vs. compar- ators in after-mar- | |
| ators in after-mar- ket review of | |
| pooled clinical tri- | |
| als, incl in FDA- | |
| approved indica- | |
| tions. | |

Glycopeptides, lipoglycopeptides

| DRUG | COVERAGE | USES | TOXICITY/ MISC | CSF |
|--------------------------------|--|---|---|-----|
| \$-\$\$\$ | Gram + cocci ex- cept VRE/VRSA | SSTI due to MRSA | Vanc IV≠PO – PO not absorbed from gut | YES |
| IV/PO vancomy- cin | <i>Most</i> Gram + rods (but see below) Corynebacterium | HAP/CAP due to MRSA | Vanc requires a central IV line , due to phlebitis (which may cause fevers, | |
| CIDAL, ex- cept | Listeria C.diff (only PO) | Infections due to VRE | unnecessary antibiot- ics/cultures/increased lengths of stay) | |
| STATIC for Entero- cocci | Increasing vanco- mycin MICs > 1 assoc with treat- ment failures ("MIC creep") | | "Red man syndrome" with vanc (histamine re- lease) if infused too rap- idly—infuse over 1-2 hours | |
| | Instrinsic re- sistance in: Leuconostoc | | Leukopenia Thrombocytopenia Rare interstitial necrosis | |
| | Lactobacillus Propionobacte- rium | | Ototoxicity (abrupt, irre- versible, usually elderly) | |
| | Pediococcus Erysipelothrix Clostridia(non- diff.) | | Rarely, DRESS | |
| \$\$\$ | MSSA, MRSA, Group A, B strep- tococci, Strep an- | SSTI, osteomyeli- tis/prosthetic joint in- fection, endocarditis, | Nausea, headache, diar- rhea | |
| IV dalba- vancin | ginosus group | bloodstream infection | Hepatotoxicity | |
| CIDAL | | 1500mg IV x1 OR 1000mg IV then 500mg in 7 days | "Red man syndrome" with vanc (histamine re- lease) if infused too rap- idly—infuse over 1-2 hours | |
| | | | | |
| \$\$\$ IV | MSSA, MRSA/VISA/VRS | SSTI | N/V, foamy urine | |
| televancin | A, Group A, B streptococci, Strep anginosus | HAPneumonia due to MRSA/VISA | Nephrotoxicity QTc prolongation | |
| CIDAL | group, VSEntero- coccus | | Mortality > with mod/sev renal impairment com- pared with vanco | |
| | | | Possibly teratogenic— avoid in pregnancy un- less maternal benefit ex- ceeds fetal risk | |

| | | | "Red man syndrome" with vanc (histamine re- lease) if infused too rap- idly—infuse over 1-2 hours Interferes with coag tests but not coagulation |
|--|---|---|---|
| \$\$\$ IV oritavancin CIDAL in- cluding <i>En-</i> <i>terococci</i> | MSSA, MRSA, Group A, B, C streptococcus, Streptococcus an- ginosus group, VSEnterococcus | SSTI **FAILED for osteo- myelitis** 1200mg IV x1, over 3 hr | Headache, N/V "Red man syndrome" with vanc (histamine re- lease) if infused too rap- idly—infuse over 1-2 hours Hepatotoxicity 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 Coadministration with warfarin may result in higher exposure of warfa- rin and increase risk for bleeding; monitor fre- quently for signs of bleeding |

Cyclic Lipopeptides

| DRUG | COVERAGE | USES | TOXICITY/ MISC | CSF |
|------------------|--|---|--|--------------|
| \$\$\$\$ | All Gram + cocci incl. Vanc-/Amp- resistant* Entero- | SSTI Bacteremia Osteomyelitis, Joint | IV only | UN- KNOWN |
| IV daptomycin | coccus | infections | False Prothrombin Time prolongation | |
| CIDAL | MRSA/VRSA | May be active in biofilms (which usu- ally inactivate anti- biotics) | Nausea/vomiting Rhabdomyolysis & associated renal in- sufficiency (weekly creatinine, CPK) Rare asthmatic pul- | |
| | | | NOT for primary | |
| | | | preumonia because it is inactivated in al- veolar 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-suscepti- ble. | |

Streptogramins

| DRUG | COVER-AGE | USES | TOXICITY/ MISC | CSF |
|---------------------|----------------|-------------------|-------------------------------|---------|
| quinupristin- | Vanc-/Amp- | SSTI/non-MRSA | IV only | |
| dalfopristin | resistant* En- | D () | | UNKNOWN |
| **** | teroco-ccus | Bacteremia | Needs central IV line | |
| \$\$\$\$ | faecium | Endocarditis due | due to frequent pain, | |
| | MSSA | to VRE faecium | phlebitis, fever | |
| CIDAL | Group A | | | |
| | Strep | Extremely limited | >30+% Myalgias/ | |
| _ . /. | | use. | Arthralgias | |
| Discontin- | NOT Enter- | | Nausea/ | |
| ued by | oco-ccus fae- | | Vomiting/ | |
| | calis or | | Diarrhea | |
| Pfizer 2022 | MRSA | | | |
| | | | *ampicillin/ | |
| | | | amoxicillin (CIDAL) | |
| | | | preferred in VRE that | |
| | | | is amp-susceptible. | |

Oxazolidinone (INTRACELLULAR ACTIVITY)

| DRUG | COVER-AGE | USES | TOXICITY/ MISC | CSF |
|--|---|---|---|---|
| \$\$\$\$ IV/PO linezolid STATIC except CIDAL for strepto- cocci | All Gram + cocci incl. ** vanc-/amp-re- sistant* Entero- coccus MRSA/VRSA TB/Atypical my- cobacteria Binds 23S rRNA- blocks formation of 50s/70s ribo- somal initiation complex | SSTI MRSA HAP/CAP due to MRSA Osteomyelitis/ Joint infections (good bone pene- tration) **NOT for bacte- remia without a well-defined and removal or drain- ing focus, NOT for endovascular in- fections | IV=PO (bioequiva- lent) Nausea/ vomiting/ diarrhea Temporary tooth staining Headache Thrombocytopenia/ Neutropenia after 7 days Peripheral/ Optic neuropathies with extended use Lactic acidosis (nau- sea, fatigue) Serotonin syn- drome: Avoid high tyramine food/drink, SSRIs (> 100mg tyramine per meal). E.g. aged cheeses, dried/pro- cessed meats, etha- nol, sauerkraut, soy sauce, or yeast ex- tract/supplements, ferments */**ampicillin/amoxi- cillin (CIDAL) pre- ferred in VRE that is amp-susceptible. **Associated with treatment failure in bacteremia, incl line & endovascular in- fections. | GOOD Myrianthefs et al. Serum and CSF concentrations of linezolid in neurosurgery patients. AAC 2016. 50(12): 3971-6. |

| \$\$\$\$ IV/PO tedizolid STATIC | All Gram + cocci incl. ** VRE, Amp-re- sistant* Entero- coccus, MRSA/VRSA Binds 50s riboso- mal subunit | SSTI | IV=PO (bioequiva- lent) 6 days tedizolid Qdaily = 10 days linezolid BID = higher lipid solubility/higher tissue levels Nausea/headache/di- arrhea Lower throm- bocytopenia than linezolid; similar neu- ropathic events; no longer term data Serotonin syn- drome: Avoid high tyramine per meal). E.g. aged cheeses, dried/pro- cessed meats, etha- nol, sauerkraut, soy sauce, or yeast ex- tract/supplements, ferments | NO DATA – suspect similar to linezolid |
|--|--|------|--|--|
|--|--|------|--|--|

COVERAGE USES TOXICITY/MISC CSF DRUG \$\$\$ Gram - including Intraabdominal in-IV/Aerosol YES Pseudomonas, Acifections IV netobacter UTI/GU infections 30% Nephrotoxicolistin Pneumonia/ city! membrane disrup-Hospital-associated polymixin B colistimethate tion, binds lipopolyrespiratory infec-Peripheral/ saccharide (LPS)/ tions Optic neuropa-Gram - endotoxin thies CIDAL Potent anti-LPS binding/ Neuromuscular neutralizing activity blockade (may exacerbate myasthenia gravis & paralytic agents)

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

Rifamycins

| DRUG | COVERAGE | USES | TOXICITY/MISC | CSF |
|---|--|---|---|-----|
| \$-\$\$ IV/PO rifampin PO rifabutin | COVERAGE Very broad, incl GPC/GNR, myco- bacteria; use is condition-specific RAPID RE- SISTANCE if given alone – Use in combinations | USES Only used alone as prophylaxis against Neisseria meningiti- dis (2 days), He- mophilus influenza b (4 days) in con- tacts/nasal carriage Combination treat- ment in serious S | TOXICITY/MISC IV/PO Red urine, sweat, tears, sa- liva – hold soft contact use Nausea, abd pain | YES |
| Only rifampin is discussed here, in context of use out- side of mycobac- terial infections CIDAL | Inhibits DNA-de- pendent RNA pol- ymerase | ment in serious S. aureus, Streptococ- cal infections Combination treat- ment of Legionella, Anthrax, Brucella, Bartonella, Ana- plasma, Ehrlichia Combination treat- ment of tuberculous and non-tuberculous Mycobacteria | Hepatotoxicity (avoid ethanol & hepatotoxins), hyper-biliru- binemia Type I & Flu-like hypersensitivity Autoimmune re- actions Many drug inter- actions – always check an up- dated reference | |

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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