Table 8. Adult parenteral antimicrobial dosage guidelines

Antibiotic	Usual Dosages ^a
ANTIBACTERIAL AGENTS	
Penicillins	
ampicillin	1-2 g q4-6h
cloxacillin	2 g q4-6h
penicillin G	2-4 million units q4-6h
piperacillin-tazobactam	3.375 g q6h
meropenem	500 mg q6h
Cephalosporins	'
cefazolin	1-2 g q8h
cefuroxime	0.75-1.5 g q8h
ceftriaxone	1-2 g q24h
ceftazidime	1-2 g q8h
Fluoroquinolones	
ciprofloxacin	400 mg q12h
levofloxacin	500-750 mg q24h
moxifloxacin	400 mg q24h
Macrolides	
azithromycin	500 mg q24h
Aminoglycosides	
gentamicin or tobramycin	80 mg q8h
Others	·
clindamycin	600 mg q8h
cotrimoxazole (TMP-SMX)	10-20 mg/kg/day trimethoprim in divided doses q6-8h
metronidazole	500 mg q8h
vancomycin	1 g q12h or 15 mg/kg q12h
ANTIFUNGAL AGENTS	
amphotericin B	0.5-1 mg/kg q24h
fluconazole	100-400 mg q24h
caspofungin	70 mg load then 50 mg q24h
ANTIVIRAL AGENTS	·
acyclovir	5-10 mg/kg/dose q8h
ganciclovir	5 mg/kg/dose q12h

a Based on normal renal function in a 70 kg patient.

Table 9. Parenteral to oral conversion suggestions

Parenteral Drug	Oral Therapy Options ^a
ANTIBACTERIAL AGENTS	
Penicillins	
ampicillin	amoxicillin
cloxacillin	cloxacillin or cephalexin
penicillin G	penicillin V
piperacillin-tazobactam	amoxicillin-clavulanate or cotrimoxazole (TMP-SMX) +/- metronidazole or ciprofloxacin +/- metronidazole
Cephalosporins	
cefazolin	cephalexin or cloxacillin
cefuroxime	cotrimoxazole or amoxicillin-clavulanate or azithromycin/clarithromycin
ceftriaxone	amoxicillin-clavulanate or cephalexin or ciprofloxacin/levofloxacin/moxifloxacin
ceftazidime	ciprofloxacin
Fluoroquinolones	
ciprofloxacin	ciprofloxacin
levofloxacin	levofloxacin
moxifloxacin	moxifloxacin
Macrolides	
azithromycin	azithromycin
Others	
clindamycin	cloxacillin +/- metronidazole or cephalexin +/- metronidazole or clindamycin
ANTIFUNGAL AGENTS	
fluconazole	fluconazole
ANTIVIRAL AGENTS	
acyclovir	acyclovir or valacyclovir

^a Patients should be clinically stable, demonstrate clinical improvement, and be able to tolerate oral feeding and medications. Selection of oral therapy should be based on cultures and sensitivities. In absence of useful cultures, oral therapy may be selected based on potential pathogens, community-versus hospital-acquired infection, pharmacokinetics, spectrum of activity, and cost of each oral agent. Oral agents listed above represent those currently on the WRHA Formulary and does not represent all commercially available oral agents.

Table 10. Adult dosing recommendations in renal impairment^a

Drug	Creatinine Clearance (CrCl) in mL/min ^b (suggested dosage adjustment based on normal dose)						
Penicillins	(***55	, , , ,					
ampicillin	> 30 (q6h)	10-30 (q6-12h)	< 10 (q12h)				
cloxacillin		NO CHANGE	NECESSARY				
penicillin	> 50 (q4-6h)	10-50 (q6-8h)	< 10 (20-50% of usual dose) ^a				
piperacillin- tazobactam	> 40 (q6h)	20-40 (q8h)	< 20 (q12h)				
Carbapenems							
meropenem	> 50 (q6h)	30-49 (q8h)	10-29 (q12h)	< 10 (q24h)			
Cephalosporins							
cefazolin	> 50 (q8h)	10-50 (q12h)	< 10 (q24h)				
cefuroxime	> 20 (q8h)	10-20 (q12h)	< 10 (q24h)				
ceftriaxone		NO CHANGE	NECESSARY				
ceftazidime	> 50 (q8h)	30-50 (q12h)	10-30 (q24h)	< 10 (50% q24-48h)			
Aminoglycosides							
gentamicin/ tobramycin/ amikacin		ct the Pharmacist at you	ur facility for dosing ass	sistance			
Fluoroquinolones							
ciprofloxacin	> 30 (a12b)	< 30					
	(q12h)	(q24h)					
levofloxacin (e.g. CAP)	> 50 (q24h)	(q24h) 20-49 (500 mg load, then 50% q24h)	10-19 (500 mg load, then 50% q48h)				
	> 50	20-49 (500 mg load, then 50% q24h)	(500 mg load,				
(e.g. CAP)	> 50	20-49 (500 mg load, then 50% q24h)	(500 mg load, then 50% q48h)				
(e.g. CAP) moxifloxacin Macrolides azithromycin	> 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE	(500 mg load, then 50% q48h)				
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents	> 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE	(500 mg load, then 50% q48h) NECESSARY				
(e.g. CAP) moxifloxacin Macrolides azithromycin	> 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE	(500 mg load, then 50% q48h) NECESSARY				
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents	> 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h)	(500 mg load, then 50% q48h) NECESSARY NECESSARY				
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole	> 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h)	(500 mg load, then 50% q48h) NECESSARY NECESSARY < 20 (25% of usual dose q24h)				
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole caspofungin	> 50 (q24h) 5 > 50 (q24h) > 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h)	(500 mg load, then 50% q48h) NECESSARY NECESSARY < 20 (25% of usual dose q24h)	< 10 (50% q24h)			
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole caspofungin Antiviral Agents acyclovir (induction doses)	> 50 (q24h)	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h) NO CHANGE	(500 mg load, then 50% q48h) NECESSARY NECESSARY < 20 (25% of usual dose q24h) NECESSARY 10-25				
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole caspofungin Antiviral Agents acyclovir ganciclovir	> 50 (q24h) s > 50 (q24h) > 50 (q8h) 50-69	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h) NO CHANGE 25-50 (q12h)	(500 mg load, then 50% q48h) NECESSARY NECESSARY 20 (25% of usual dose q24h) NECESSARY 10-25 10-25	(50% q24h) < 10			
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole caspofungin Antiviral Agents acyclovir ganciclovir (induction doses) Miscellaneous clindamycin	> 50 (q24h) s > 50 (q24h) > 50 (q8h) 50-69	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h) NO CHANGE (q12h) 25-49 2.5 mg/kg q24h NO CHANGE	(500 mg load, then 50% q48h) NECESSARY NECESSARY	(50% q24h) < 10			
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole caspofungin Antiviral Agents acyclovir ganciclovir (induction doses) Miscellaneous	> 50 (q24h) s > 50 (q24h) > 50 (q8h) 50-69	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h) NO CHANGE (q12h) 25-49 2.5 mg/kg q24h NO CHANGE	(500 mg load, then 50% q48h) NECESSARY NECESSARY	(50% q24h) < 10			
(e.g. CAP) moxifloxacin Macrolides azithromycin Antifungal Agents fluconazole caspofungin Antiviral Agents acyclovir ganciclovir (induction doses) Miscellaneous clindamycin	> 50 (q24h) > 50 (q24h) > 50 (q8h) 50-69 2.5 mg/kg q12h	20-49 (500 mg load, then 50% q24h) NO CHANGE NO CHANGE 20-50 (50% q24h) NO CHANGE (q12h) 25-49 2.5 mg/kg q24h NO CHANGE	(500 mg load, then 50% q48h) NECESSARY NECESSARY	(50% q24h) < 10 1.25 mg/kg 3x/wk			

^a Suggested dosages – for individualized dosage modifications or more information contact the Pharmacy Department at your facility.



Seven Oaks General Hospital Antibiogram for 2024

(Based on data from 2023)

Provided by: Shared Health, Clinical Microbiology Discipline

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 $^{^{}b}$ To estimate creatinine clearance (CLc_R) (mL/min) use the following calculation normalized for a 72 kilogram person. CL_{CR} male = $(140-age) \times 88.4$ CL_{CR} female = $0.85 \times$ CL_{CR} male S_{CR} (µmoles/L)

Monitor serum concentrations.

DISCLAIMERS

This guide is provided as an educational resource for physicians and other healthcare professionals caring for patients at the Seven Oaks General Hospital. The authors of the guide have made every effort to ensure that the information contained in it was accurate at the time of publication. Users of the guide are encouraged to consult other references to confirm the information presented in it. The authors are not responsible for errors, omissions, inaccuracies, or the continued completeness of the information contained in the guide. The information in the guide should not be used or relied upon to replace the skill and professional judgment required to determine appropriate patient care and treatment. Also, the guide is not intended to replace or to be used as a substitute for the complete prescribing information prepared by each pharmaceutical manufacturer for their anti-infective agents. Because of possible changes in anti-infective indications, changes in dosage information, differences in patients' responses to therapy, newly described toxicities, drug-drug interactions, and other items of importance, reference to complete prescribing information is recommended before any of the anti-infective agents described in the guide are used.

HOW TO USE THE ANTIBIOGRAM PORTION OF THE GUIDE (Tables 1-6)

- The information presented in the antibiogram is intended only to guide initial empiric anti-infective agent therapy at the Seven Oaks General Hospital.
- Initial broad-spectrum empiric therapy should be focused to the most appropriate narrow-spectrum agent(s) based on the laboratory identification of pathogen(s) and known susceptibility patterns/results, if the situation permits.
- Consideration should be given to equally efficacious but less expensive anti-infective
 agents for empiric therapy or when streamlining of therapy is desired, if the situation
 permits.

SUGGESTED CRITERIA FOR IV TO ORAL ANTIBIOTIC CONVERSION IN ADULTS

- Clinical improvement of infectious signs and symptoms (e.g., temperature defervescence, decreased white blood cell count).
- Patient is clinically stable (excludes patients in the intensive care unit, patients with febrile neutropenia, or patients with life threatening infections).
- Patient can tolerate oral feeding and medications (bowel sounds, no diarrhea/nausea/ vomiting).
- For rapid step-down, choose agents with high bioavailability (e.g., clindamycin, cotrimoxazole (TMP-SMX), fluoroquinolones).
- If anti-infective agent susceptibilities are known, anti-infective therapy should be tailored based on available data.

Table 1. In vitro activity of selected anti-infective agents tested against Gram-negative bacillia

		Percent Susceptible													
Organism (number tested): January through December 2023														- azole	۶
= Not tested, not routinely reported, or not recommended	Ampicillin	Amoxicillin- Clavulanate	Piperacillin- Tazobactam	Cefazolin	Cephalexin	Cefuroxime	Ceftriaxone	Ceftazidime	Ertapenem	Meropenem	Gentamicin	Tobramycin	Ciprofloxacin	Trimethoprim- Sulfamethoxazole	Nitrofurantoin
Citrobacter spp. (36)			78				67	75	97	100	97	94	86	89	94
Enterobacter cloacae complex (30)			70				57	60	93	100	97	90	87	83	50
Escherichia coli (54) systemic	48	83	100	63			81	89	100	100	89	85	59	74	
Escherichia coli (430) urine	48	80	98	73	n.d.		84	90	100	100	89	89	56	72	97
Haemophilus influenzae (158) ^d	80	n.d.				n.d.								65	
Klebsiella pneumoniae (104)		85	95	81	n.d.		90	91	99	99	91	94	78	80	31
Klebsiella/Raoultella spp. (51) ^c		84	90	22			96	96	100	100	96	96	94	94	84
Proteus mirabilis (37)	70	97	100	n.d.	n.d.		100	100	100	100	95	97	92	84	
Pseudomonas aeruginosa (93)			90					95		94		98	87		

- Isolates tested and reported are from all sources combined, with the exception of Escherichia coil (subdivided into systemic isolates and urine isolates); Isolates were collected from Jan 1 to Dec 31, 2023 with the exception of Cirobacter spp. and Klebsiella oxytoca (collected from Jan 2022 to Dec 2023); data compiled according to the recommendations of the Clinical and Laboratory (Standards Institute (CLSI) in their document M39, 5th ed. (2022).
- b Cephalexin is only indicated for the treatment of uncomplicated lower urinary tract infections.
- Nitrofurantoin is only indicated for acute cystitis.
- d H. influenzae data obtained from isolates tested at Health Sciences Centre, Jan 1 to Dec 31, 2023. Only 134 isolates were tested for Trimethoprim-Sulfamethoxazole. Data from adult and pediatric patients.
- The current laboratory identification system is unable to differentiate Klebsiella oxytoca from Raoultella spp.
- n.d. = no data absence of data for certain drug-organism combinations reflects limitations of the testing method currently used by Shared Health Clinical Microbiology laboratories.

Table 2. In vitro activity of selected anti-infective agents tested against Gram-positive coccia

			Percent Susceptible												
Organism (number tested): January through December 2023 = Not tested, not routinely reported, or not recommended	Penicillin	Ampicillin	Oxacillin ^b	Vancomycin	Daptomycin	High-Level Gentamicin*	High-Level Streptomycin°	Erythromycin ^d	Clindamycin	Trimethoprim- Sulfamethoxazole	Rifampin°	Linezolid	Tetracycline	Levofloxacin	Nitrofurantoin
Enterococcus spp. (214)		93		99	n.d.	82	87					n.d.			89
Staphylococcus aureus (404)			66	100	100			46	70	98	100	100	95		99
Staphylococcus epidermidis (35)			40	100	100			43	77	69	100	100	86		100
Staphylococcus lugdunensis (55)			98	100	100			85	85	100	100	100	96		100
Streptococcus pyogenes (100) ⁹ (Group A Streptococcus)	100			100					82					99	
Streptococcus agalactiae (148) ^h (Group B Streptococcus)	100			100					64						

- Isolates tested and reported are from all sources (surveillance isolates excluded), Jan to Dec, 2023 with the exception of Staphylococcus lugdunensis (collected from Jan 2022 to Dec 2023); data compiled according to the recommendations of
- Staphylococcus ingouners (or included in on an azuzz or bec 2023), data compined according to the recommendations of the Clinical and Laboratory Standards Institute (CLSI) in their document M39, 5° ed. (2022).

 Document M39, 5° ed. (2022).
- inhibitor combinationis, cephalosporins, and carbapenems for Staphylococcus aureus and coagulase-negative staphylococci.

 Susceptibility to high-level pentamicin or high-level streptomycin indicates that these agents can be used nombination with a cell wall active agent (e.g., ampicillin or vancomycin) for synergy. Gentamicin and streptomycin should never be used alone as treatment for Entercooccus spor.
- ^d Erythromycin activity predicts the activity of azithromycin and clarithromycin for staphylococci and streptococci.
- ^e Rifampin should NOT be used alone as treatment for infection.
- Nitrofurantoin is indicated for acute cystitis only.
- ⁹ Streptococcus pyogenes isolates were obtained from wound and sterile site specimens submitted to Shared Health Clinical Microbiology laboratories between January and December, 2023.
- Streptococcus agaiactize isolates were obtained from vaginal/rectal swabs submitted for Group B Streptococcus detection to the Health Sciences Centre, St. Boniface Hospital, and Westman Laboratory in 2022. n.d. = no data absence of data for certain drug-organism combinations reflects limitations of the testing method currently used by Shared Health Clinical Microbiology laboratories.

n.d. = no data – absence of data for certain drug-organism combinations reflects limitations of the testing method currently used by Shared Health Clinical Microbiology laboratories.

Table 3. In vitro activity of selected anti-infective agents tested against Streptococcus pneumoniae^a

	Percent Susceptible								
Infection Type (number tested) = Not tested, not routinely reported, or not recommended	Penicillin (oral)	Penicillin (intravenous)	Ceftriaxone	Vancomycin	Levofloxacin	Clarithromycin	Doxycydine	Trimethoprim- Sulfamethoxazole	
Systemic Isolates (Blood + CSF) ^b									
Meningitis (218)		79	95	100				87	
Non-Meningitis infection (218)	79	98	99	100	100	n.d.	n.d.	87	
Respiratory Isolates ^c									
Non-Meningitis infection (40)	78	95	98	100	100	68	83	80	

- For Streptococcus pneumoniae, different susceptibility breakpoints for penicillin and celtriaxone exist depending on whether meningitis or a non-meningitis infection is being trated [CLS, M100, 33° edition]. For penicillin, when treating a non-meningitis infection different breakpoints exist for oral and intravenous dosing. For non-meningitis infections, susceptibility to oral penicillin predicts susceptibility to amoxicillin. Oral a centra sent and concordate for the treatment of bacterial menindistin.
- b Systemic isolates were obtained from the Health Sciences Centre (HSC) and St. Boniface Hospital (SBH) clinical microbiology laboratories between January and December, 2023. CSF = cerebrospinal fluid.
- Respiratory isolates were obtained from patients at the Health Sciences Centre (HSC) and St. Boniface Hospital (SBH) between January and December. 2018.

n.d. = no data.

Table 4. In vitro activity of selected anti-infective agents tested against Methicillin-Susceptible and Methicillin-Resistant Staphylococcus aureus isolates^a

	Percent Susceptible							
Organism (number tested) = Not tested, not routinely reported, or not recommended	Oxacillin ^b	Vancomycin	Trimethoprim- Sulfamethoxazole	Erythromycin	Clindamycin	Tetracycline	Linezolid	Daptomycin
Methicillin-Susceptible Staphylococcus aureus (270)	100		97	57	67	94		
Methicillin-Resistant Staphylococcus aureus (149)	0	100	99	24	73	97	100	100

- ^a Isolates tested and reported are from all sources (surveillance isolates excluded), Jan to Dec, 2023; data compiled according to the recommendations of the Clinical and Laboratory Standards Institute (CLSI) in their document M39, 5th ed. (2022).
- b Oxacillin accurately predicts the activity of all semi-synthetic penicillins, including cloxacillin, beta-lactam/beta-lactamase inhibitor combinations, cephalosporins, and carbapenems for Staphylococcus aureus.

Table 5. In vitro activity of selected anti-infective agents tested against anaerobic isolates collected from hospitals in Winnipega

	Percent Susceptible							
Organism (number tested) = Not tested, not routinely reported, or not recommended	Penicillin	Amoxicillin- Clavulanate	Piperacillin- Tazobactam	Clindamycin	Meropenem	Metronidazole		
Bacteroides fragilis (108)		93	n.d.	44	93	100		
Bacteroides thetaiotaomicron (37)		94	n.d.	14	97	100		
Prevotella bivia (54)	7	100	n.d.	32	100	96		
Prevotella disiens (34)	32	97	n.d.	18	100	100		

a Isolates were obtained from WRHA hospitals between Jan 2019 and Dec 2020; data compiled according to the recommendations of the Clinical and Laboratory Standards Institute (CLSI) in their document M39, 5th ed. (2022)

n.d. = no data – absence of data for certain drug-organism combinations reflects limitations of the testing method currently used by Shared Health Clinical Microbiology laboratories.

Table 6. In vitro activity of selected anti-fungal agents tested against Candida species collected from hospitals in Winnipeg^a

	Percent Susceptible							
Organism (number tested)	Fluconazole°	Voriconazole	Micafungin					
Candida albicans (33)	100	100	100					
Candida glabrata (44)	98	n.d.	100					

- ^a Data obtained by testing a random sample of C. albicans isolates from Health Sciences Centre and St. Boniface Hospital, collected between Jan 2017 and Dee 2018. Susceptibility interpretations are based on updated CLSI breakooints (M27M44. 3" Edition). Isolates tested and reported are from blood only.
- ^b Data obtained by testing C. glabrata isolates from Shared Heath Clinical Microbiology laboratories, collected between Jan and Dec 2023. Susceptibility interpretations are based on updated CLSI breakpoints (M27M44, 3st Edition). Isolates tested and reported are from blood only.
- For fluconazole, there is only a susceptible-dose dependent (SDD) breakpoint for C. glabrata. The percentage of C. glabrata isolates that tested SDD to fluconazole was 98%. Susceptibility of SDD isolates to fluconazole is dependent on achieving the maximum blood level possible (i.e., should use the maximum dosage regimen). Consultation with infectious diseases is recommended for further quidance.
- n.d. = breakpoints have not been defined for voriconazole versus C. glabrata.

Table 7. Adult oral antimicrobial dosage guidelines

A 411 1 41	1	0 (0)		
Antibiotic	Usual Dosages	Cost (\$) per day		
ANTIBACTERIAL AGENTS				
Penicillins				
amoxicillin	500 mg tid	1.10		
amoxicillin-clavulanate	500 mg tid or 875 mg bid	2.75-3.00		
cloxacillin	500 mg qid	1.50		
penicillin V	300 mg qid	0.30		
Cephalosporins				
cephalexin	500 mg qid	1.80		
Macrolides				
azithromycin	250-500 mg daily	1.25-2.50		
clarithromycin	250-500 mg bid	2.25-3.25		
Fluoroquinolones				
ciprofloxacin	250-750 mg bid	1.40-2.50		
levofloxacin	500-750 mg daily	3.50-6.50		
moxifloxacin	400 mg daily	1.50		
Others				
clindamycin	450-600 mg tid	1.50-3.00		
cotrimoxazole (TMP-SMX)	1 DS (double strength) tab bid	0.25		
doxycycline	100 mg bid	1.30		
nitrofurantoin (Macrobid®)	100 mg bid	1.50		
metronidazole	500 mg tid	0.35		
ANTIFUNGAL AGENTS				
fluconazole	100-400 mg daily	5.55- 22.20		
itraconazole	200-400 mg daily	8.00-16.00		
ANTIVIRAL AGENTS				
acyclovir	200-800 mg 5x/day	5.00-16.00		
valacyclovir	1 g tid	5.25		
Approximate cost per inpatient day excl	uding dispensing costs as of Eabruan, 2017 b	ased on the Manitoha		

^a Approximate cost per inpatient day excluding dispensing costs as of February 2017 based on the Manitoba Drug Interchangeability Formulary and Manufacturer's List Prices. Prices have been rounded.