33. Antimicrobial Guide

This clinical pathway is intended to supplement, rather than substitute for, professional judgment and may be changed depending upon a patient's individual needs. Failure to comply with this pathway does not represent a breach of the standard of care.

For detailed guidelines and other conditions not listed below, refer to your hospital's guidelines for antimicrobial use

Condition	Comments/Caveats	Recommended Therapy		
URTI/Sinusitis AVOID PRESCRIBING ANTIBIOTICS FOR UPPER RESPIRATORY TRACT INFECTIONS SINCE MOST ARE VIRAL.	The most common cause of URTIs is viral and thus no antibiotics are necessary A clinician should diagnose Acute Bacterial Rhinosinusitis (ABRS) when a) symptoms or signs of Acute Rhinosinusitis (ARS) (purulent nasal drainage accompanied by nasal obstruction, facial pain/pressure/fullness, or both) persist without evidence of improvement for at least 10 days beyond the onset of upper respiratory symptoms or b) symptoms or signs of ARS worsen within 10 days after initial improvement (double worsening). DO NOT ORDER A CT SCAN TO DIAGNOSE SINUSITIS	Amoxicillin/Clavulanate 1gm PO BD x 5-10 days is the first-line therapy for most adults who meet the criteria for ABRS In Penicillin-Allergic Patients: Azithromycin 500mg PO OD x 3 days Supportive therapy; • Decongestants (a-adrenergic) - xylometazoline hydrochloride for 3 days. • Saline irrigation - Nasal saline irrigation, alone or in conjunction with other adjunctive measures, may improve quality of life, decrease symptoms, and decrease medication use for ABRS, particularly in patients with frequent sinusitis. • Mucolytics • Antihistamines have no role in the symptomatic relief of ABRS in non-atopic patients.		
Pharyngitis/Tonsillitis AVOID PRESCRIBING ANTIBIOTICS FOR UPPER RESPIRATORY TRACT INFECTIONS SINCE MOST ARE VIRAL.	The most predictable clinical parameter for GABHS pharyngitis is reported to be the Centor Score (available on MDCalc) a) Age < 15 years (+1) or ≥ 45 years (-1) b) History of fever > 38°C c) Absence of cough, d) Swollen and tender anterior cervical lymph nodes e) Tonsillar exudates or swelling	Adult patients with acute exudative adult pharyngitis who report ≥ 4 Centor Score ONLY Benzathine penicillin G 1.2MU IM stat OR Amoxicillin/Clavulanate 1gm PO BD x 5-10 days Consider - Single-dose Prednisone 60 mg PO or Dexamethasone 8 mg IM therapy added to the standard treatment has a more rapid improvement of pain in adult patients with acute exudative adult pharyngitis who report ≥ 4 Centor Score Patients who are allergic to Penicillin Azithromycin: 500 mg PO on day 1 followed by 250 mg PO OD for 4 days		
Laryngitis	Mostly viral	No Antibiotics necessary		
Acute Gastroenteritis AVOID PRESCRIBING ANTIBIOTICS FOR ACUTE GASTROENTERITIS WITHOUT SYSTEMIC DISEASE OR DYSENTERY	Any diarrhoeal illness lasting > 1 day, especially if accompanied by the following features should prompt evaluation of a faecal specimen; bloody diarrhoea moderate-severe disease (systemically ill/toxic appearing patients) symptoms lasting >7 days immunocompromised patients recent use of antibiotics A Stool Culture is NOT NECESSARY OR COST-EFFECTIVE in most cases of diarrhoea without systemic disease or dysentery unless an unusual bacterial cause is suspected Typhoid - Bone marrow culture is the most sensitive routinely available diagnostic tool. Stool culture is positive only in up to 30-40% of cases but is often negative by the time that systemic symptoms bring patients to hospital. Blood cultures are positive in 40-80% of patients. Serologic tests e.g. the Widal test are of limited clinical utility because positive results may represent a previous infection.	Food-borne toxigenic diarrhoea usually requires only supportive treatment, not antibiotics. Treatment of salmonellosis with antibiotics (including quinolones) can prolong the carrier state and lead to a higher clinical relapse rate. Treat ONLY patients with; • bloody diarrhoea • moderate-severe disease (systemically ill/toxic appearing patients) • symptoms lasting >7 days • immunocompromised patients • recent use of antibiotics Ciprofloxacin 500 mg PO BD x 3 days. The duration of treatment may be extended by 2-3 days for moderate-to-severe cases. The antimotility agent loperamide (imodium) may reduce the duration of diarrhoea when given with antibiotics for traveller's diarrhoea. A loperamide/simethicone combination has demonstrated faster and more complete relief. Loperamide may cause dangerous prolongation of illness in patients with some forms of bloody or inflammatory diarrhoea and, therefore, should be restricted to patients with non-bloody stool.		



Condition	Comments/Caveats	Recommended Therapy		
Urinary Tract Infection (UTI)	Cloudiness of the urine is most often due to protein or crystal presence, and malodorous urine may be due to diet or medication use. A urinalysis with quantitative urine WBC counts should NOT be used alone to support a diagnosis of UTI or start antimicrobial therapy in any patient population. A negative Leukocyte Esterase AND a negative urine Nitrate largely rule out infection in pregnant women, elderly patients, family medicine, and urology patients. The combination of a negative leukocyte esterase and negative nitrite test demonstrated a UTI negative predictive value of 88% (95% confidence interval [CI] 84–92%). Pyuria in a urine specimen, in the absence of symptoms (Asymptomatic Bacteriuria), is NOT AN INDICATION for antimicrobial therapy. Urine cultures are NOT RECOMMENDED in most cases of uncomplicated UTIs in adult women. Urine Cultures ONLY for; In patients suspected of having pyelonephritis, a urine culture and susceptibility test should always be performed, and initial empiric therapy should be tailored appropriately based on the likely infecting uropathogen. A urine specimen should be obtained for culture and susceptibility testing before initial antimicrobial therapy for complicated UTIs.	Uncomplicated Cystitis Ciprofloxacin 500 mg PO BD x 3 days OR Nitrofurantoin 100mg TDS x 3 days Uncomplicated Pyelonephritis, Outpatient Therapy Ceftriaxone 1 g IV stat PLUS Ciprofloxacin 500 mg PO BD x 7 days UTI during Pregnancy, Outpatient Therapy Cefuroxime 500 mg PO BD for 7 days OR Nitrofurantoin 100mg TDS x 3 days		
	Complicated UTI Male gender Structural or functional anatomic abnormalities Renal stones Indwelling catheters Renal transplant Neurogenic bladder Recent urologic procedure	Complicated UTI Ciprofloxacin 500 mg PO BD x 14 days		
	Inpatient therapy • Sepsis • Pregnancy • Urinary tract obstruction • Persistent vomiting • Poor outpatient follow-up	Uncomplicated Pyelonephritis, Inpatient Therapy Ceftriaxone 1g IV OD 10-14 days OR Ciprofloxacin 400 mg IV BD x 10-14 days UTI during Pregnancy, Inpatient Therapy Ceftriaxone 1-2 g IV OD		
Sepsis & Septic Shock	See Sepsis & Septic Shock Algorithm	Give ANTIBIOTICS as an EMERGENCY (within the FIRST HOUR or recognition of Sepsis/Septic Shock) Ceftriaxone 2gm IV stat For probable Neutropenic patients or if patient has been admitted in hospital in the last 3 months (Hospital Acquired Infection) Imipenem 500 mg IV infusion over 3 hrs then QID for General sepsis OR Meropenem 1 gm IV infusion over 3 hrs then TDS for possible CNS infections		



Condition Comments/Caveats Recommended Therapy Community-Acquired In addition to a constellation of suggestive clinical **Outpatient Treatment** Pneumonia features, a demonstrable infiltrate by chest Amoxicillin/Clavulanate 1gm PO BD x 7 - 10 days radiograph or other imaging technique, with or In Penicillin-Allergic Patients: without supporting microbiological data, is required for the diagnosis of pneumonia. Azithromycin: 500 mg PO on day 1 followed by 250 mg PO OD for The strongest indications for blood cultures are severe CAP and in immunocompromised patients or those with significant co morbidities, as these patients are more likely to be infected with pathogens other than S pneumoniae. Co morbidities: . Chronic heart, lung or renal disease · Diabetes mellitus Alcoholism Malignancy Asplenia · Immunosuppressant condition or drugs Inpatient Treatment Inpatient Therapy Amoxicillin/Clavulanate 1.2gm IV T x 7 - 10 days CURB65 ≥ 2 (available in MDCalc) PLUS · Patient factors requiring hospitalization Azithromycin 500mg IV OD x 7 - 10 days Healthcare Associated Pneumonia (HCAP) **HCAP** risk factors? Antipseudomonal beta-lactam . Hospitalization for 2 or more days of the past 90 Imipenem 500mg IV infusion over 3 hours QID • Resides in nursing home or long-term care facility Received chemotherapy, IV antibiotics, or wound care within the prior 30 days · Attended a hospital or haemodialysis clinic in the last 30 days Malaria **Defining Criteria for** Finding Uncomplicated Malaria Severe Malaria Artemether + Lumefantrine - Coartem® 80/480 1 tablet at 0, 8, 24, Impaired A Glasgow coma score < 11 36, 48 and 60 hours (six doses). consciousness in adults or a Blantyre coma (cerebral malaria) score < 3 in children Body weight (kg) Dose (mg) of artemether + lumefantrine given twice daily for 3 days Generalized weakness so 5 to < 15 20 + 120 Prostration that the person is unable to 15 to < 25 40 + 240 sit, stand or walk without 25 to < 35 60 + 360assistance 2.35 80 + 480Multiple > 2 episodes within 24 h convulsions **Acidosis** A hase deficit of > 8 mFg/I IV Artesunate 2.4mg/kg at 0, 12 and 24 hours and daily until or, if not available, a plasma patient can take oral. Children weighing < 20 kg should receive a bicarbonate level of < 15 higher dose of artesunate (3 mg/kg bw per dose) to ensure mmol/L or venous plasma equivalent exposure to the drug. lactate ≥ 5 mmol/L. Severe acidosis manifests clinically as respiratory distress (rapid, deep, laboured breathing). Hypoglycaemia Blood or plasma glucose < 2.2 mmol/L (< 40 mg/dL) Severe malarial Haemoglobin concentration ≤ 5 g/dL or a haematocrit of ≤ 15% anaemia in children < 12 years of age (< 7 g/dL and < 20%, respectively, in adults) with a parasite count > 10 000/µL Renal impairment Plasma or serum creatinine > 265 µmol/L (3 mg/dL) or blood urea > 20 mmol/L Jaundice Plasma or serum bilirubin > 50 umol/L (3 mg/dL) with a parasite count > 100 000/ µL



omments/Cave		Kecommend	Recommended Therapy		
Defining Criteria for Severe Malaria Finding Pulmonary oedema Radiologically confirmed or oxygen saturation < 92% on		Uncomplicated Malaria Artemether + Lumefantrine - Coartem* 80/480 1 tablet at 0, 8, 24 36, 48 and 60 hours (six doses).			
	room air with a respiratory rate > 30/min, often with	Body weight (kg)	Dose (mg) of artemether + lumefantrine given twice daily for 3 days		
	chest in-drawing and crepitations on auscultation	5 to < 15	20 + 120		
gnificant bleeding	Including recurrent or	15 to < 25	40 + 240		
	prolonged bleeding from	25 to < 35	60 + 360		
	the nose, gums or	≥ 35	80 + 480		
yperparasitemia	venepuncture sites; haematemesis or melena Compensated shock is defined as capillary refill 2 3 s or temperature gradient on the leg (mid to proximal limb), but no hypotension. Decompensated shock is defined as systolic blood pressure < 70 mm Hg in children or < 80 mm Hg in adults, with evidence of impaired perfusion (cool peripheries or prolonged capillary refill). mia P. falciparum parasitaemia > 10% Severe Malaria IV Artesunate 2.4mg/kg at 0, 12 and 24 hours and daily until the patient can take orally. Children weighing < 20 kg should receive higher dose of artesunate (3 mg/kg bw per dose) to ensure equivalent exposure to the drug.				
Most abscesses are Staph aureus. Most cellulitis is Group A beta-haemolytic streptococcus (although some are Staph aureus)		Oral Therapy			
		Beta-haemolytic Streptococcus coverage: Amoxicillin/Clavulanate 1gm PO BD x 7 days			
	treptococcus pyogenes (beta-	0	R		
haemolytic streptococcus) is recommended Clindamycin is bacteriostatic, potential for cross- resistance and emergence of resistance in		Clindamycin 450 mg PO QID x 7-10 days			
ythromycin-resistar	nt strains; inducible resistance	Parenteral Therapy (Inpatient)			
in MRSA		Beta-haemolytic Streptococcus and MSSA Coverage			
Effective treatment of abscesses entails incision, thorough evacuation of the pus, and probing the cavity to break up ovulations. Gram stain, culture, and systemic antibiotics are rarely indicated unless there is extensive surrounding cellulitis, fever, multiple lesions, severely impaired host defences, or cutaneous gangrene.		-			
		Cefazolin 1gm IV q8 hours for 7-10 days OR Clindamycin 600 mg IV q8 hours for 7-10 days			
Surgical intervention is the major therapeutic modality in cases of necrotizing fasciitis. Necrotizing fasciitis falls into two groups; The spontaneous extremity cellulitis is usually Group A Streptococcus and sometime Staph aureus. The second group includes head and neck,		Consult a Surgeon			
ultiple cutar odalit ecroti The sp Group aureus	e lesions, seveneous gangrei lintervention y in cases of n zing fasciitis fa contaneous ex A Streptococis. econd group in	e lesions, severely impaired host defences, neous gangrene. Intervention is the major therapeutic y in cases of necrotizing fasciitis. zing fasciitis falls into two groups; nontaneous extremity cellulitis is usually A Streptococcus and sometime Staph s.	e lesions, severely impaired host defences, neous gangrene. Initervention is the major therapeutic yin cases of necrotizing fasciitis. zing fasciitis falls into two groups; nontaneous extremity cellulitis is usually A Streptococcus and sometime Staph s. e.		



Condition	Comments/Caveats	Recommended Therapy
STI – Urethritis, Epididymitis, Orchitis, Proctitis, Cervicitis	Minimum criteria for clinical diagnosis of PID (all 3 should be present): a) Bilateral lower abdominal (uterine) tenderness (sometimes radiating to the legs) b) Cervical motion tenderness - Positive cervical motion tenderness is defined as increased discomfort from a normal pelvic examination, as stated by the patient. Of note, cervical motion tenderness is neither sensitive nor specific for gynaecologic pathology, is a sign of nonspecific peritoneal inflammation, c) Bilateral adnexal tenderness (with or without a palpable mass) One or more of the following additional criteria can be used to enhance the specificity of the minimum criteria and support a diagnosis of PID: • oral temperature >38.3° C; • abnormal cervical or vaginal mucopurulent discharge; • presence of abundant numbers of WBC on saline microscopy of vaginal fluid; and • laboratory documentation of cervical infection with N. gonorrhoea or C. trachomatis.	STI – Urethritis, Epididymitis, Orchitis, Proctitis, Cervicitis Ceftriaxone 250mg IM stat PLUS Azithromycin 1gm PO stat PID Mild-Moderate disease Ceftriaxone 250mg IM stat PLUS Doxycycline 100mg PO BD x 14 days WITH or WITHOUT Metronidazole 500mg PO BD x 14 days Severe disease/In-patient therapy - Suggested criteria: • surgical emergencies (e.g., appendicitis) cannot be excluded; • the patient is pregnant; • the patient does not respond clinically to oral antimicrobial therapy; • the patient is unable to follow or tolerate an outpatient oral regimen; • the patient has severe illness, nausea and vomiting, or high fever; or • the patient has a tubo-ovarian abscess. Amoxicillin/Clavulanate 1.2g IV BD PLUS Doxycycline 100mg IV/PO BD x 14days

Condition	Comments/Caveats		Recommended Therapy		
HIV Post Exposure Prophylaxis (PEP)	Exposed individual must be HIV negative at baseline Exposure must have occurred within the past 72 hours Exposure must be high-risk. Faeces, nasal secretions, saliva, sputum, sweat, tears, urine, and vomitus are not considered to be infectious unless they are visibly bloody.		PEP should be initiated as soon as possible after exposure, but no later than after 72 hours. Consult local guidelines for the recommended regimens		
	Estimated per-unprotected act risk for acquisition of HIV by exposure route				
	Exposure route	% Risk	Regimen	Dose	Comments
	Blood transfusion	90%		ADULTS	
	Needle-sharing injection-drug use	0.67%	Tenofovir/Lamivudine	1 tablet OD	Zidovudine AZT
	Receptive anal intercourse	0.5%	TDF/3TC		
	Percutaneous needle stick	0.3%	(300/300mg)		(300mg) can be used as an
	Receptive penile-vaginal intercourse	0.1%	PLUS	PLUS	alternative when
	Insertive anal intercourse	0.06%			TDF cannot be used
	Insertive penile-vaginal intercourse	0.1%	Atazanavir/Ritonavir (ATV/r)	1 tablet OD with food	
	Receptive oral intercourse	0.01%	(300/100mg)	1000	
	Insertive oral intercourse	0.005%			
	The overall rate of HIV transmission through percutaneous inoculation is reported to be 0 (95% confidence interval [CI] 0.2–0.5); the ris acquiring an HIV infection is greater for percutaneous injuries that involve; - hollow-bore needles that have been in c with an artery or vein, - when blood is visible on the device, - a deep needle stick, and - when the source patient has advanced I disease. Splashes or infectious material to mucous membranes or broken skin may also transmi infection (estimated risk per exposure, 0.099) CI 0.006–0.5). Exposure of intact skin to contaminated blood has not been identified risk for HIV transmission. • Counsel on risks and benefits of PEP and ob verbal consent for testing (HIV, FHG, UEC, L HBV and HCV) • Voluntary HIV testing for source individuals • Offer PEP as soon as high-risk exposure is established and exposed individual tests HI negative at baseline (if HIV testing not avail an provide 1-2 days of PEP to cover until H performed) • Pregnancy testing • Cr (if TDF-containing regimen) and Hb (if AZ containing regimen), however PEP should b offered even when lab tests are not availab not delay administration of PEP while waiti lab results • Hepatitis B vaccination (if not previously immunized & not known HBV positive)		PLUS dosages alternative when		can be used as an alternative when ABC cannot be used as all 28 days of a days after starting hs, if negative, test ing applies PEP

