Therapy of Infections in Neutropenic Patients

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Introduction

An immunocompromised host is a patient with

defects in host defenses that predispose to infection.

Risk factors include:

- 1. Neutropenia.
- 2. Immune system defects (from disease or immunosuppressive drug therapy).
- 3. Compromise of natural host defenses.
- 4. Environmental contamination.
- 5. Changes in the normal flora of the host.

1. Neutropenia:

- Neutropenia is defined as an abnormally reduced number of neutrophils circulating in peripheral blood.
- An absolute neutrophil count (ANC) of less than
 1,000 cells/mm³ indicates a reduction sufficient to predispose patients to infection.
- The development of infection depends on:

a) the severity of neutropenia. 1
b) the rate of neutrophil decline.
c) the duration of neutropenia.

- All neutropenic patients are considered to be at risk for infection, but those with ANC less than 500 cells/mm³ are at greater risk than those with ANCs of 500 - 1,000 cells/mm³.
- Bacteria and fungi commonly cause infections in neutropenic patients.

Allon - 300 ~ Sever neutropenia 1 100 ~ protound neutrophil Apatients who are less than 500 is at greater Risk for infections.

- 2. Immune System Defects:
- Defects in T-lymphocyte and macrophage function (cell-mediated immunity), B-cell function (humoral immunity), or both predispose patients to infection.

- 3. Destruction of Protective Barriers: Innuous membranes.
- This is a major factor predisposing immunocompromised patients to infection.
- a) Damage to skin and mucous membranes by surgery, venipuncture, IV and urinary catheters, radiation, and chemotherapy.
- b) Chemotherapy-induced mucositis of the oropharynx and GIT establish a portal for subsequent infection by bacteria, HSV, and



- c) Medical and surgical procedures, such as transplant surgery, indwelling IV catheter placement, bone marrow aspiration, biopsies, and endoscopy, further damage the skin & mucous membranes and predispose patients to infection.
- Infections resulting from disruption of protective barriers usually are caused by skin flora such as
 S. aureus, S. epidermidis, and various
 streptococci.

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- 4. Environmental contamination:
- **Contaminated equipment such as nebulizers** a) and ventilators, and contaminated water supplies predispose for outbreaks of P. aeruginosa and Legionella pneumophila infections, respectively.
- b) Fruits and green leafy vegetables are sources of gram negative bacteria and fungal infections in immunocompromised hosts. Non clean water

- 5. Changes in the normal microbial flora of the host:
- a) Administration of broad-spectrum antimicrobial agents disrupts GIT flora and predisposes patients to infection with more virulent pathogens.
- b) Antineoplastic drugs (cyclophosphamide, doxorubicin, and fluorouracil, ...) and <u>acid-</u> <u>suppressive therapy</u> (histamine H₂-receptor antagonists, proton-pump inhibitors, and antacids) also may disrupt GIT flora and predispose to infection.

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Risk Factors and Common Pathogens in Immunocompromised Patients

Risk Factor	Patient Condition	Common Pathogens 11 Page 1
Neutropenia	Acute leukemia Chemotherapy	Bacteria: Staphylococcus aureus, Staphylococcus epidermidis, and other coagulase- negative staphylococci, streptococci, enterococci are most common, followed by Escherichia coli, Klebsiella pneumoniae, Pseudomonas aeruginosa, Fungi: Candida, Aspergillus, Mucorales (Mucor) Viruses: Herpes simplex
Impaired cell-mediated immunity	Lymphoma Immunosuppressive therapy (steroids, cyclosporine, tacrolimus sirolimus, mycophenolate, azathioprine and anti- neoplastic agents	 Bacteria: Listeria, Nocardia, Legionella, Mycobacteria Fungi: Cryptococcus neoformans, Candida, Aspergillus, Histoplasma capsulatum Viruses: Cytomegalovirus, varicella-zoster, herpes simplex Pneumocystis jiroveci Yeast-like fungus
Impaired humoral immunity	Multiple myeloma, Chronic lymphocytic leukemia (have progressive hypogammaglobulinemia Splenectomy Immunosuppressive therapy (steroids, chemotherapy)	Bacteria: encapsulated organisms such as S. pneumoniae, H. influenzae, N. meningitidis Which might produce life-threatening infections a)

Loss of protective skin barriers	Venipuncture, bone marrow aspiration, urinary	Bacteria: S. aureus, S. epidermidis, Bacillus spp., Corynebacterium jeikeium	
	catheterization, vascular access devices, radiation, biopsies	Fungi: Candida	
Loss of protective	Respiratory support	Bacteria: S. aureus, S. epidermidis,	
membranes	equipment, endoscopy,	aeruginosa, Bacteroides spp.	
barriers	chemotherapy,		
	radiation	Fungi: Candida	
		Viruses: Herpes simplex	
Surgery	Solid-organ	Bacteria: S. aureus, S. epidermidis,	
	transplantation	Enterobacteriaceae, P. aeruginosa, Bacteroides spp.	
		Fungi: Candida	
		Viruses: Hernes simplex	
		viruses. respes simplex	
Alteration of	Antimicrobial	Bacteria: Enterobacteriaceae, P. aeruginosa,	
flora	шегару	Legionena, 5. aureus, 5. epiderinidis	
	Chemotherapy Acid –lowering agents Hospital environment	Fungi: Candida, Aspergillus	
Blood products,	Bone marrow	Fungi: Candida	
donor organs	transplantation	Viruses: Cytomegalovirus, Epstein–Barr virus	
	Solid-organ	hepatitis B, hepatitis C	
	transplantation	Protozoa: Toxoplasma gondii	

Management of Febrile Episodes in

Neutropenic Patients

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Goals of therapy:

- 1. Protect the patient from early death caused by undiagnosed infection.
- 2. Prevent breakthrough bacterial, fungal and viral infections during periods of neutropenia.
- 3. Effectively treat established infections.
- 4. Reduce morbidity. 7

- 5. Avoid unnecessary use of antimicrobials that contribute to increased resistance.
- 6. Minimize toxicities and cost of antimicrobial therapy while increasing patient quality of life.
- Empirical broad-spectrum antibiotic therapy is effective at reducing early mortality. *p p*-90% *mortality*.

ما سنستنى ادر رحبرد المشاف بالمع من معما Spectrum antibiotics





No withholding

b)	Undiagnosed infection in
	immunocompromised patients can rapidly
	disseminate and results in death.
	Empirical antibiotic therapy is 70-90% effective
	at reducing early morbidity and mortality.
2	Antimicrobial therapy must be appropriate and
	should be initiated promptly in afebrile
	patients with clinical signs and symptoms of
	infection.

3. When designing optimal empirical antibiotic regimens, physicians must consider infection patterns and antimicrobial susceptibility trends in their respective institutions. المحكاة بم بشتعن Patient factors such as, risk of infection, drug allergies, concomitant nephrotoxins, and previous antimicrobial exposure (including prophylaxis) must be considered.

- 5. Risk stratification drives both type and setting of antimicrobial therapy:
- **1)** Low-risk patients:
- a) have an anticipated duration of neutropenia ≤

7 days.

- b) are clinically stable.
- c) have no or few co-morbidities.←
- d) have no bacterial focus or systemic signs of infection other than fever.



Management of Febrile Episodes in Neutropenic Patients High-risk patients should be hospitalized for parenteral antibiotics, whereas low-risk patients may be candidates for oral or outpatient Parantel iSoo antibiotics. **Because of their frequency and relative** pathogenicity, P. aeruginosa and other gramnegative bacilli and *staphylococci* are the primary targets of empirical antimicrobial



- The optimal antibiotic regimen remains controversial.
- All empirical regimens must be: carefully monitored and appropriately revised on the ¹/₂ basis of documented infections, susceptibilities

of bacterial isolates, development of more³

defined clinical signs and symptoms of infection,

or a combination of these factors.

Recognized antibiotic regimens:

- **1.** Monotherapy with an antipseudomonal β
 - lactam (cefepime or ceftazidime), a carbapenem (imipenem-cilastatin or meropenem), or

(2)

- 2. Two-drug combination therapy with an
 - antipseudomonal β-lactam + either an
 - aminoglycoside or an antipseudomonal
 - fluoroquinolone (ciprofloxacin or levofloxacin).



Monotherapy or two-drug combination therapy as above, + the addition of vancomycin



- There is no significant difference, overall, between monotherapy and combination therapy (βlactam/aminoglycoside) in rates of survival, response, and bacterial/fungal superinfections.
- A higher rate of adverse effects was observed in *a*minoglycoside-containing combination regimens.
- Cefepime and antipseudomonal carbapenems have good activity against viridans streptococci and pnéumococci but not all gram positive bacteria.

Disadvantages:

Regimen 1:

limited gram positive activity, and high rate of superinfection)

Regimen 2:

- 1. Antipseudomonal β-lactam + aminoglycoside: limited gram positive activity, potential for nephrotoxicity and need of TDM.
- 2. Antipseudomonal β-lactam + fluoroquinolone: limited gram positive activity and development of resistance.

Regimen 3:

Selection of vancomycin resistant enterococci, risk of nephrotoxicity and need for TDM.



- 2) Requires compliant patients with 24-hour access to medical care in case it is needed.
- 3) Requires supporting home environment.

Antimicrobial Therapy After Initiation of Empirical Therapy After 2 to 4 days of empirical antimicrobial therapy, the clinical status and culture results should be reevaluated to determine whether therapeutic modifications are necessary. **During periods of neutropenia**, patients should continue to receive broad-spectrum therapy because of risk of secondary infections or breakthrough bacteremia when antimicrobial Coverage is too narrow. Econdary infection and Superintedictive mining Superinfection is the development of an infection to the breatment of other infection and resulted Secondary infection 's deferent Kind of infection duc to damage - I the tissues by the first infection from broad Spectrum avorage and eradication



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Antimicrobial Therapy After Initiation of Empirical Therapy

- Duration of treatment should be appropriate for the particular organism and site and should continue for at least the duration of neutropenia (until ANC ≥ 500 cells/mm³) or longer if clinically necessary.
 In patients who become afebrile after 2 to 4
 - days of therapy with NO infection identified,
 - continue antibiotic therapy until neutropenia

has resolved (ANC \geq 500 cells/mm³).

- You may switch therapy to an oral regimen (ciprofloxacin plus amoxicillin-clavulanate) after
 2 days of IV therapy, in <u>low-risk</u> patients who become afebrile and who have NO evidence of infection. - You doit Start with oral theory at first
- In <u>high-risk</u> patients, <u>parenteral antibiotic</u> regimens should be continued until resolution of neutropenia.

Fever after 2 or more days of antibiotic therapy can be due to:

- 1) nonbacterial infection
- resistant bacterial infection or infection slow to respond to therapy
- 3) emergence of a secondary infection
- 4) inadequate drug concentrations
 - drug fever
 - 6 infection at a non-vascular site (catheter infection or abscess)
 - 7) **noninfectious causes such as:**
 - a. tumors
 - b. administration of blood products

- Persistently febrile patients should be evaluated carefully, but modifications generally are NOT made to initial antimicrobial regimens within the first 2 to 4 days of therapy unless there is evidence of clinical deterioration.
- Antibiotic regimens may require modification in
 patients experiencing toxicities as well as in
 patients with evidence of progressive disease,
 clinical instability, or documentation of an
 organism NOT covered by the initial regimen.

- Addition of vancomycin should be considered, if NOT already part of the regimen. Kuman _____ add Wantomycin
- If vancomycin was included in the initial empirical regimen and the patient is still febrile after 2 to 3 days of therapy without isolating a gram-positive pathogen, discontinuation of vancomycin should be considered to reduce the risk of toxicities or resistance.

- Neutropenic patients who remain febrile despite > 4 - 7 days of broad-spectrum antibiotic therapy are candidates for antifungal therapy.
- A significant percentage of febrile patients who die during prolonged neutropenia have evidence of invasive fungal infection on autopsy, even when they have NO evidence of fungal disease before death.



- Persistence of fever or development of a new fever during broad-spectrum antibiotic therapy may indicate the presence of a fungal infection, most commonly *Candida* or *Aspergillus* spp.
- Blood cultures for fungi are positive in < 50% of neutropenic patients with invasive fungal infections, and waiting for isolation of fungal organisms is associated with high morbidity and mortality.

- Empirical antifungal therapy, thus, should be initiated after 4 to 7 days of broad-spectrum antibiotic therapy in persistently febrile patients if the duration of neutropenia is expected to be greater than 1 week.
- Antifungal therapy must be adequate to treat
 undiagnosed fungal infection and prevent fungal
 superinfection in high-risk patients.

 Empirical coverage for both Candida spp. and Aspergillus should be considered because these organisms are responsible for more than 90% of fungal infections in neutropenic patients.

- Aspergillus is particularly common in patients with hematologic malignancies and amphotericin B is the preferred agent.
 - Lipid-associated amphotericin B (LAMB) products are similar in efficacy to conventional amphotericin B while causing fewer toxicities, and can be used at higher doses (3 mg/kg).
 - LAMB products have significantly higher cost.



- من المحمدة والجد المحمدة • <u>Posaconazole</u> has extended activity against some
- Mucorales, in addition to Candida and

Aspergillus, but is only approved for prophylaxis of Aspergillus and Candidal infections in

neutropenic patients.

TDM is recommended for some azole antifungals given the potential for interpatient variability, therapeutic failure associated with subtherapeutic concentrations, and toxicities associated with supratherapeutic concentrations.

- The echinocandin antifungals (caspofungin, micafungin, and anidulafungin) have broad spectrum of antifungal activity and favorable adverse effect profiles. (1) Caspofungin is as effective as and better

 - tolerated than, liposomal amphotericin B for

empirical treatment of neutropenic patients with persistent fever. Therefore, it is considered

an appropriate alternative to LAMB and

voriconazole.

· Amphotoricin (asparagetters) + Variconazole (- 2), i - i - i Jeis - Eigeburgin (

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Monitoring of Antifungal Agents

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Drug	Adverse Reaction	Monitoring	Comments
		Parameters	
Amphotericin	Nephrotoxicity, ¹	Serum_	Liposomal preparations
B	hepatotoxicity, 2	c <u>reatinine</u> ,	associated with less renal
(lipid-	electrolyte (2)	electrolytes,	toxicity, similar efficacy to
associated)	disturbances,	LFTs, blood	standard preparation.
LAMB	infusion reactions	pressure, heart	Electrolyte disturbances
boen		rate	occur before creatinine
0ú.	. Lea		alterations. Pretreatment
have to	risicol		and slow infusion may
Yeaghror	lore		decrease
nor	por fr		incidence of infusion reaction
mean	don		
Jones and	atinin		40

			I
Posaconazole	Hepatotoxicity,	LFTs, skin,	Poor absorption with
	rash; interactions	posaconazole	suspension, goals of >1
	with CYP3A4	serum	µg/mL for treatment and
		concentrations	>0.7 µg /mL for prophylaxis.
	7		Parenteral formulation not
			recommended for patients
	well we ve o		with CrCL <50 mL/min
			Multiple interactions with
	TPM		drugs motobolized by CVP
			arugs metabolized by C 11
			SA4, including
			immunosuppressants; close
•			monitoring needed.
V riconazole	Mental status	Mental status,	Mental status/visual changes
	changes, headache,	visual function,	associated with elevated
	hallucinations	LFTs, ECG,	troughs > 5.5 μg /m; goal
	v sual disturbances,	voriconazole	trough 1-5.5 μg/mL for
	hepatotoxicity, QTc	serum	treatment and prophylaxis,
	prolongation;	concentrations	target trough of $> 2 \ \mu g/ml$ in
	interactions with		disease with poor prognosis.
	ond 344		rarenteral formulation not
	anu JA4		with CrCI <50 mI /min
	Tom		Multiple interactions
			manipic meracions



Initiation of Antiviral Therapy

- Acyclovir and the newer antivirals valacyclovir and famciclovir may be used.
- Routine use of antiviral agents in the management of patients without mucosal lesions or other evidence of viral infection is NOT recommended.

* Symptome of Viral intertions usually appear after the Medication has ended - but success rate is higher in prevention rather than treatment.

Initiation of Antiviral Therapy its avoided by good hydration and avoidance of rapid infusion rate

Adverse reactions of acyclovir:

- Nausea, diarrhea, headache
- IV administration may be associated with reversible crystalline nephropathy or interstitial nephritis; or neurologic toxicity (tremors, delirium, seizures).

These are uncommon with adequate hydration and

avoidance of rapid infusion rates.

Drug Interactions:

Probenecid and cimetidine decrease acyclovir

clearnce and increase exposure.

 \rightarrow Acyclovir + zidovudine \rightarrow somnolence and lethargy.

* A V

- The optimal duration of antimicrobial therapy remains controversial.
- Decisions regarding discontinuation of empirical antimicrobial therapy are more difficult than those of initiation of therapy.
- The patient's ANC is the most important factor for the total duration of antibiotic



- If ANC is ≥ 500 cells/mm³ for two consecutive days, if the patient is afebrile and clinically stable for 48 hours or more, and if NO pathogen has been isolated, antibiotics may be discontinued.
- Some clinicians advocate that patients with ANC
 < 500 cells/mm³ be maintained on antibiotic
 therapy until resolution of neutropenia, even if
 they are afebrile.

- Prolonged antibiotic use has been associated with superinfections resulting from resistant bacteria and fungi and increased risk of antibiotic-related toxicities.
- If <u>low-risk</u> patients are stable clinically with negative cultures but the ANC still is < 500 cells/mm³) antibiotics <u>may be discontinued</u> after a total of 5 - 7 afebrile days.

Duration of Antimicrobial Therapy high risk Patients with severe neutropenia (ANC > 100) but < 500 cells/mm³), mucosal lesions, or unstable vital signs or other risk factors should continue to receive antibiotics until ANC becomes \geq 500 cells/mm³, and the patient is

stable clinically.



- receive antimicrobial therapy until the infecting organism is eradicated and signs and symptoms of infection have resolved (at least 10-14 days of كياج اسبرعس therapy).
- Any way, therapy must be individualized based on individual patient parameters and response to therapy.

Granulocyte-macrophage colony-stimulating Factor (Sargramostim)

Granulocyte colony-stimulating factor (filgrastim)

- May be used as adjunct therapy to antimicrobial treatment of febrile neutropenic patients.
 - They reduce total duration and severity of chemotherapy-related neutropenia.
 - They reduce duration of antibiotic use.
 - They reduce hospitalizations, and decrease hospital length of stay.

Overall mortality or infection-related mortality is NOT decreased.

Not toutine.

- CSFs should NOT be routinely used in patients with uncomplicated fever and neutropenia.
- Patients with prolonged neutropenia and documented severe infections who are NOT responding to appropriate antimicrobial therapy may benefit from treatment with CSFs.
- CSFs should be considered in patients who are at high risk of infection-associated complications, or who have factors that are predictive of poor clinical



- 1) Profound neutropenia (ANC <100 cells/mm³)
- 2) Expected prolonged period of neutropenia (>10 days)
- 3) Patient age >65 years
- 4) Uncontrolled primary disease
- 5) <u>Sepsis syndrome, or severe infection manifest by</u> hypotension and multiorgan dysfunction
- 6) Pneumonia ----
- 7) Invasive fungal infection <---
- 8) Other clinically documented infection <---
- 9) Hospitalized at the time of the development of fever

10) Severe complications during previous episode of febrile neutropenia.

Granulocyte CSF (or GM-CSF) Common Adverse Effects:

- 1. Bone pain: because of proliferation of WBCs in bone marrow. Relieved with analgesics.
- 2. Leukocytosis.
- 3. Bruises, bleeding gum and nose bleeding: Due to drop in platelet count.
- 4. Headache displacement -> Planelet development il ilso al lo
- 5. Fatigue: can be prolonged up to one year.
- 6. Back pain.
- 7. Hepatic problems: reversible with discontinuation of the drug
- 8. Diarrhea or constipation.

- 9. Malaise.
- 10. Fever
- 11. Splenomegally
- 12. Splenic rupture is a rare but serious.
- 13. Inflammation around the injection site.
- 14. Abdominal pain
- 15. Edema in hands and feet, peripheral edema and pleural or pericardial effusions due to a capillary leak syndrome.
- 16. Insomnia.
- 17. Arthralgias & myalgias.