



MSS

MICROBIOLOGY

#2



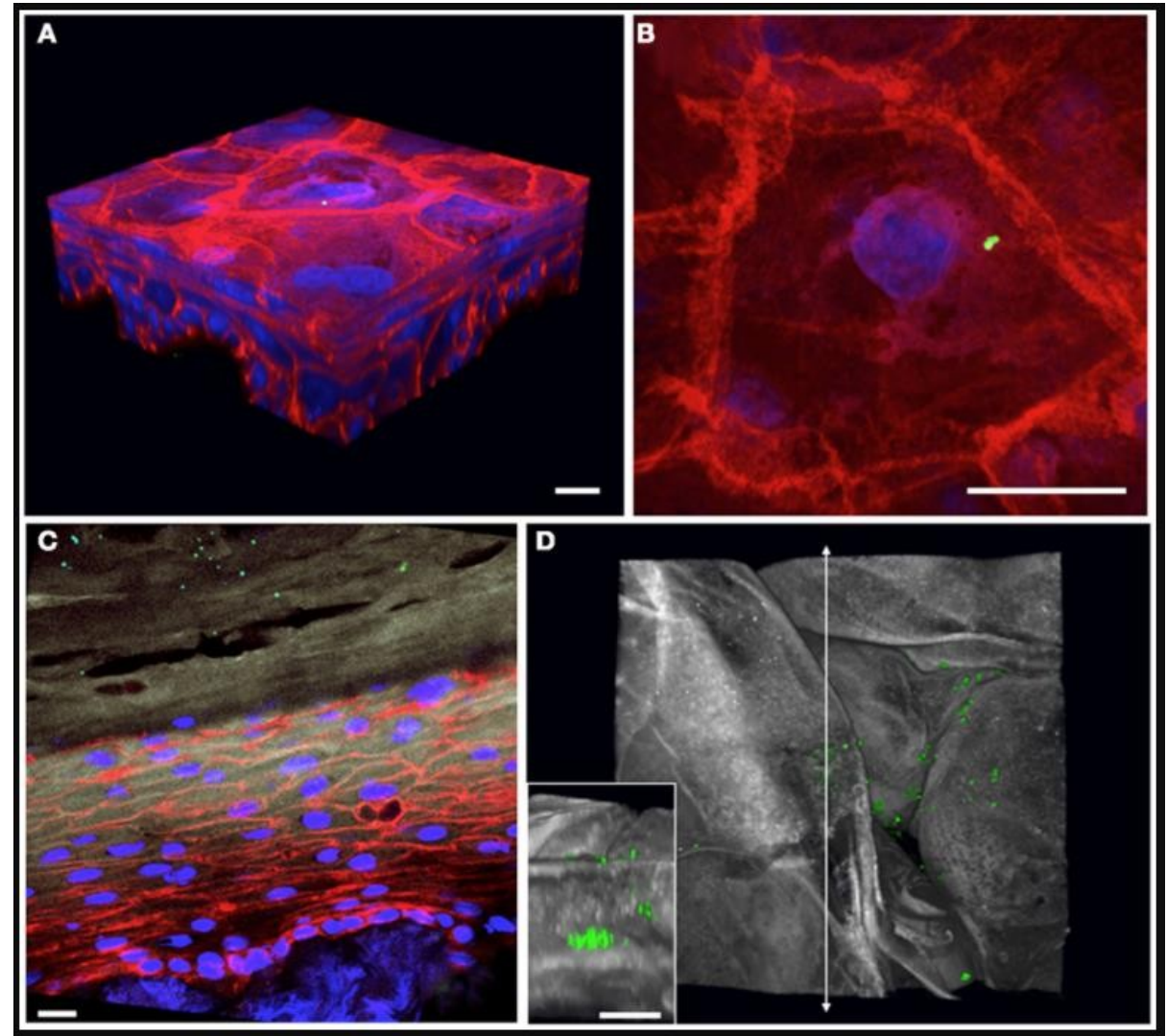
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Musculoskeletal System Microbiology

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Necrotizing soft tissue infections

Overview

In this lecture we will discuss:

- Necrotizing fasciitis .
- Clostridial myonecrosis
- Pyomyositis
- Diabetic foot infections

In this lec we will talk about the infections that can spread to the subcutaneous fat n fascia that surrounds the muscles n infections of the muscles themselves.

Introduction

- Necrotizing soft tissue infections (NSTIs) include necrotizing forms of **fasciitis, myositis, and cellulitis**. These infections are characterized clinically by **fulminant tissue destruction, systemic signs** of toxicity the patient may have sepsis ; over exaggerated immune response to this infection leading to organ damage eventually death , **and high mortality**.
- NSTI can include involvement of **the epidermis, dermis, subcutaneous tissue, fascia, and muscle**.
- Necrotizing infection may be categorized based on microbiology (pathogen that involved)and the presence or absence of gas in the tissues(as the case in clostridium perfringens which cause gas gangrene).
- Sometimes referred to in the press as flesh-eating bacteria .(not a scientific name journalist use it for attention)

Necrotizing fasciitis

- Necrotizing fasciitis is an infection of the deep soft tissues that results in **progressive destruction of the muscle fascia and overlying subcutaneous fat**. muscle tissue is frequently spared (usually not affected, it's rare) because of its generous blood supply which means a lot of immune cells decreasing the chance of getting infected)
- Initially, the overlying tissue **can appear unaffected**; therefore, necrotizing fasciitis is difficult to diagnose without **direct visualization of the fascia**. (when u look at it u will see some erythema which may look not that bad but once u open n explore it from inside u will notice the massive necrotizing (damage) in the fascia n underlying tissue)
- Necrotizing fasciitis may be divided into two microbiologic categories: **polymicrobial** (type I) as it's name applies 'poly' caused by many type of pathogens , and **monomicrobial** infection (type II) ; caused by GAS (strep pyogenes)

Necrotizing fasciitis/ polymicrobial

- Typically, at least one anaerobic species (most commonly **Bacteroides**, **Clostridium**, or **Peptostreptococcus**) is isolated in combination with **Enterobacteriaceae** (eg, Escherichia coli, Enterobacter, Klebsiella, Proteus) and one or more facultative anaerobic **streptococci** (other than group A Streptococcus [GAS]).
- Necrotizing fasciitis of the perineum, known as **Fournier gangrene**, can occur as a result of a breach in the integrity of the gastrointestinal or urethral mucosa.
- **Fournier gangrene** is a form of **polymicrobial (type I) infection**. Fournier gangrene typically begins abruptly with severe pain and may spread rapidly to the anterior abdominal wall and the gluteal muscles. Men are more commonly affected than women. Involvement in men may include the scrotum and penis.

Necrotizing fasciitis/ polymicrobial

Necrotizing fasciitis of the perineum
Swelling, hematoma , severe pain .

Might spread to other parts, adjacent tissue, blood
n causing systemic infection.

Once u suspect necrotizing fasciitis
Visualization not enough u have to open n explore
the lesion to figure out the extent of the damage,
debridement , obtain specimen to culture it ,
antibiotic sensitivity tests .

Fournier's gangrene in a patient with diabetes

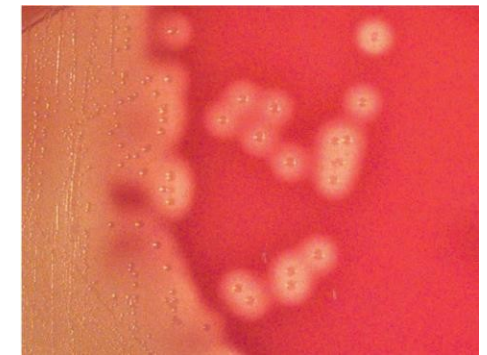
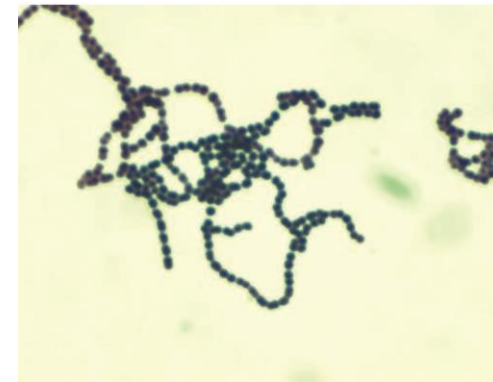
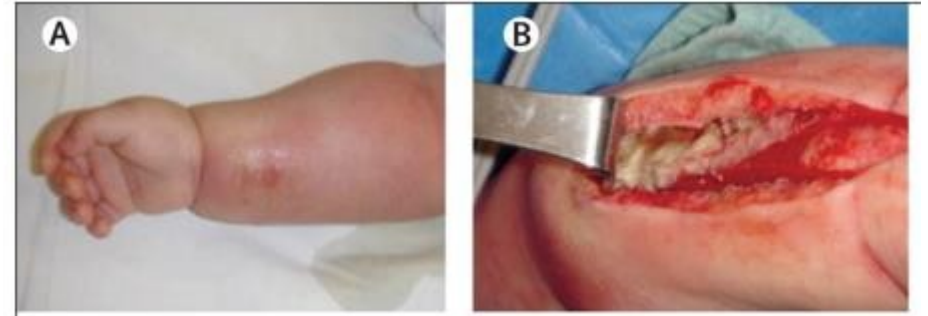


Necrotizing fasciitis of the perineum (Fournier's gangrene) can involve the scrotum. The infection can begin abruptly with severe pain and may spread rapidly.

Reproduced with permission from Lawrence B Stack, MD.

Necrotizing fasciitis/ monomicrobial

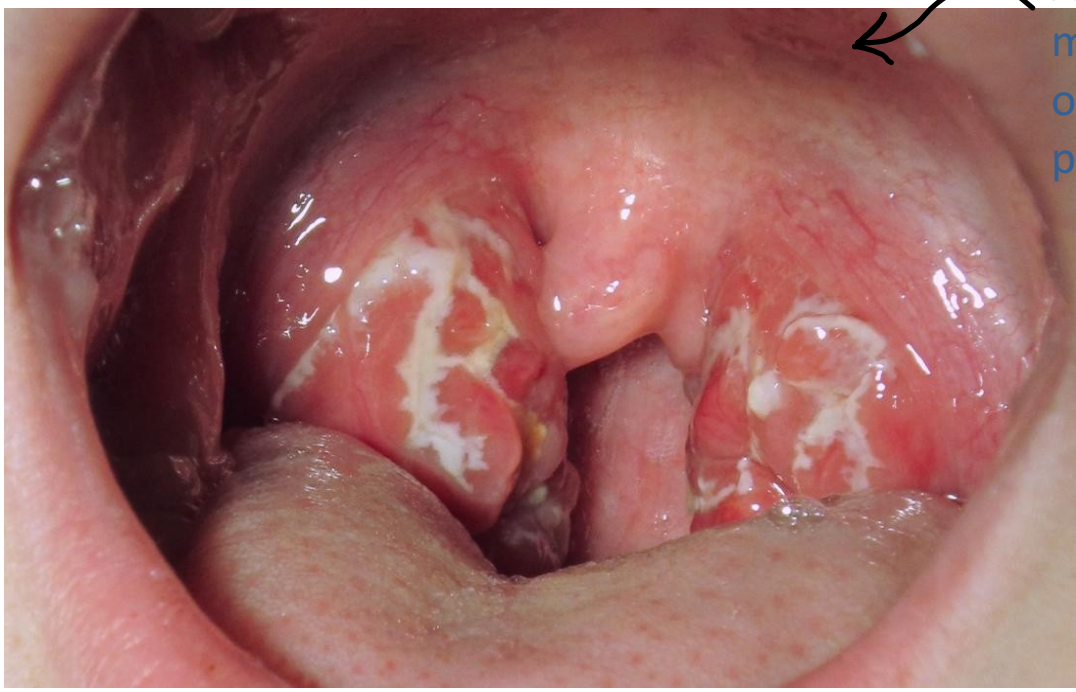
- Monomicrobial (type II) necrotizing infection is usually caused by **GAS or other beta-hemolytic streptococci** : GBS , but more often caused by GAS). Infection may also occur as a result of **Staphylococcus**. Infection with no clear portal of entry occurs in about half of cases; in such circumstances, the pathogenesis of infection likely consists of **hematogenous translocation of GAS** from the throat (asymptomatic or symptomatic pharyngitis) to a site of blunt trauma or muscle strain.
- **M protein** is an important virulence determinant of **GAS**. Necrotizing infection caused by GAS strains with M types 1 and 3 is associated with streptococcal toxic shock syndrome in about 50 percent of cases.



↖ GAS : Beta (complete)
hemolysis on blood agar

Pic A : there is a swelling and some erythema which may not reflect the amount of extensive damage that appears when u open it as in Pic B.

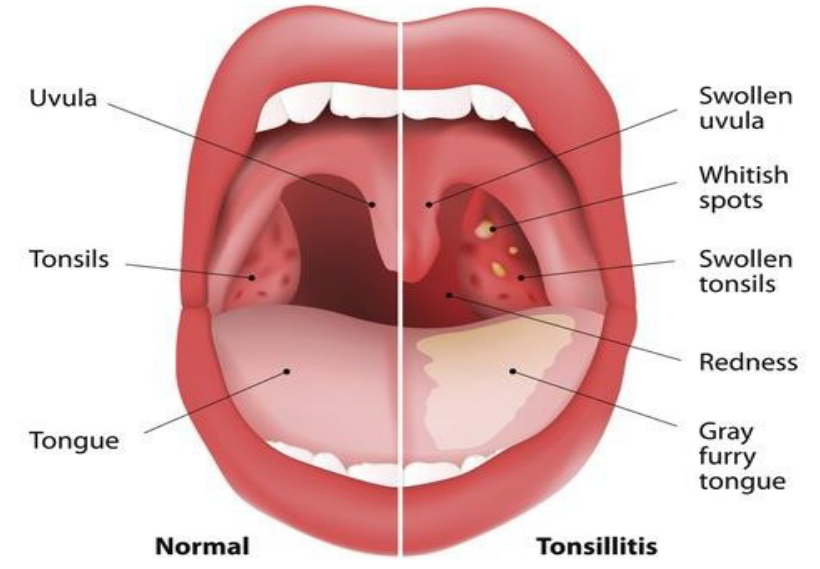
Streptococcus pyogenes



As u remember the most common cause of bacterial pharyngitis is GAS

Introduction to Microbiology and Immunology

BACTERIAL TONSILLITIS



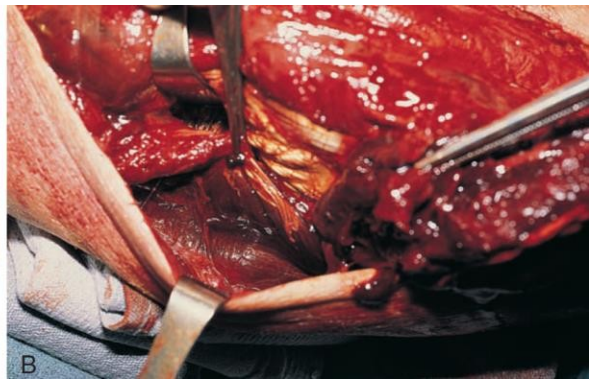
Cellulitis



Erysipelas



Necrotizing fasciitis



Necrotizing fasciitis/ risk factors

- Necrotizing infection can occur among healthy individuals with no past medical history or clear portal of entry in any age group .

Risk factors for NSTI include:

- Penetrating trauma.
- **Recent surgery** (including colonic, urologic, and gynecologic procedures as well as neonatal circumcision) and **Mucosal breach** (hemorrhoids, rectal fissures, episiotomy).
- Immunosuppression (**diabetes**, cirrhosis, **neutropenia**, **HIV infection**, malignancy)

Diabetes is a particularly important risk factor for necrotizing infection involving the lower extremities, perineum, and head and neck region.

Necrotizing fasciitis/ CLINICAL MANIFESTATIONS

- Necrotizing infection most commonly involves the **extremities** (lower extremity more commonly than upper extremity), particularly in patients with diabetes and/or peripheral vascular disease. Necrotizing infection usually presents **acutely** (over hours); rarely, it may present subacutely (over days). **Rapid progression** to extensive destruction can occur, leading to **systemic toxicity**, limb loss, and/or death.
- **Erythema** (without sharp margins; 72 percent)
- **Edema** that extends beyond the visible erythema (75 percent)
- Severe **pain** (out of proportion to exam findings in some cases; 72 percent)
- **Fever** (60 percent)
- **Crepitus** (50 percent) : when the infection is deep n gas is formed within the tissue , locked in space once u press on it may cause some popping sounds.
- **Skin bullae, necrosis, or ecchymosis** (38 percent)

Necrotizing fasciitis/ Diagnosis

- NSTI should be suspected in patients with **soft tissue infection** (erythema, edema, warmth) and **signs of systemic illness** (fever, hemodynamic instability) in association with crepitus, **rapid progression** of clinical manifestations, and/or severe pain (out of proportion to skin findings in some cases). Early recognition of necrotizing infection is critical.
- **Surgical exploration** is the only way to establish the diagnosis of necrotizing infection. Followed by debridement (removal of dead , damaged or infected tissue) to improve the healing potential.
- Intraoperative specimens should be sent for Gram stain and culture
- **Radiographic imaging** studies can be useful. The most useful finding is **presence of gas in soft tissues**, which is seen most frequently in the setting of clostridial infection or polymicrobial (type I) necrotizing fasciitis.

- Treatment of necrotizing infection consists of **early and aggressive surgical exploration and debridement** of necrotic tissue, together with **broad-spectrum empiric antibiotic therapy** and **hemodynamic support** : extra inf. (it aims to avoid hypotension , maintain adequate cerebral perfusion pressure..etc . Why we need it ? b/c the patient suffers from systemic manifestations (the pathogen in the blood) if he had sepsis he may end up in hypotension n septic shock that's why we need that kind of support Administration of antibiotic therapy in the absence of debridement is associated with a mortality rate approaching 100 percent.
- In general, empiric treatment of necrotizing infection should consist of **broad-spectrum antimicrobial therapy**, including activity against gram-positive, gram-negative, and anaerobic organism. (e.g. **carbapenem broad spectrum plus vancomycin for MRSA** plus Clindamycin for the antitoxin activity).
- Necrotizing infection **is associated with considerable mortality**, even with optimal therapy.

Clostridial myonecrosis/ INTRODUCTION

- **Clostridial myonecrosis (gas gangrene)** is a life-threatening muscle infection that develops either contiguously from an area of **trauma** or **hematogenously** from the gastrointestinal tract with muscle seeding. Early recognition and aggressive treatment are essential.
- **Clostridium species** are widespread in nature (**ubiquitous**), due to their ability to form **endospores**. They are commonly found in soil and marine sediments as well as human and animal intestinal tracts.
- Myonecrosis (clostridial gas gangrene) is characterized by **rapidly progressive invasion** and **destruction of healthy muscle** and other soft tissues. Traumatic gas gangrene is most commonly caused by *C. perfringens*; spontaneous gangrene is most commonly caused by the more aerotolerant *C. septicum*.

Clostridium species are: gram +ve bacilli, spore-formers n usually anaerobic

~ Deep infections mainly caused by anaerobic bacteria .

Clostridial myonecrosis/ Pathogenesis

- **Traumatic wounds with vascular compromise** (particularly deep penetrating injuries such as knife wounds, gunshot wounds, and crush injuries) create an **anaerobic environment** that is ideal for **proliferation of clostridia**. Traumatic injury accounts for about 70 percent of gas gangrene cases, and about 80 percent of these are caused by **C. perfringens**.
- Gas gangrene was a common infection in the Civil War, World War I, and World War II due to delayed treatment of injuries.
- Many extracellular toxins are produced by C. perfringens; of these, **alpha and theta toxins** have been implicated in pathogenesis.
- **Shock** associated with gas gangrene may be attributable to both direct and indirect effects of alpha and theta toxins.

Clostridium perfringens

- *C. perfringens* is responsible for a range of soft-tissue infections including **cellulitis**, fasciitis or suppurative **myositis**, and **myonecrosis** with gas formation (caused by the metabolic activity of the rapidly dividing bacteria) in the soft tissue (**gas gangrene**). The toxin involved in gas gangrene is known as **α -toxin**, which inserts into the plasma membrane of cells, producing gaps in the membrane that disrupt normal cellular function
- **Clostridial food poisoning**, an **intoxication** characterized by (1) a short incubation period (8 to 12 hours), (2) a clinical presentation that includes abdominal cramps. (3) a clinical course lasting less than 24 hours.
- *C. perfringens* produces **enterotoxin**, The enterotoxin is produced during the phase transition from vegetative cells to spores and is released in the alkaline environment of the small intestine when the cells undergo the terminal stages of spore formation (**sporulation**).



Treatment is usually **debridement and excision**, with amputation necessary in many cases. Water-soluble antibiotics (such as penicillin) alone are not effective because they **do not penetrate ischaemic muscles** sufficiently to be effective.

Clostridial myonecrosis/ Diagnosis, treatment, and outcome

- **Pain** at a site of traumatic injury together with signs of **systemic toxicity** and **gas in the soft tissue** support the diagnosis of gas gangrene. Physical evidence of **crepitus** in the soft tissue is the most sensitive and specific finding on clinical examination.
- Radiographic studies can help (to look for gas formation in the tissue) Blood cultures should be obtained.
- Treatment of traumatic gas gangrene consists of **surgical debridement, antibiotic therapy,** and **supportive measures**. Patients with trauma who have not received **tetanus immunization** for 5 years should receive a booster vaccine against tetanus. Use of **hyperbaric oxygen (HBO)** ? (High conc. Of Oxygen on the lesion may help , b/c it's anaerobic bacteria but still we don't know if it really works).
- Antibiotic agents with excellent in vitro activity against C. perfringens include **penicillin, clindamycin,** tetracycline, chloramphenicol, metronidazole (**flagyl**) for anaerobic organisms .
- Patients with associated bacteremia and intravascular hemolysis have the greatest likelihood of progressing to shock and death. Mortality is highest for patients in shock at the time of diagnosis.

Clostridial myonecrosis

Notice in the pic : swelling (edema) , pus , erythema (as we mentioned in the previous lec that in people with darker color erythema appears as change in pigmentation) .

In this video notice that at 1:36 bubbles rising from the lesion upon applying pressure meaning that it contains gas.

3:57 radiograph image shows that there's gas in the soft tissue .



<https://www.youtube.com/watch?v=53FgitG2jl4>

- Ok. Now before we proceed till now we've been talking about necrotizing soft tissue infections .NSTI can include involvement of the epidermis, dermis, subcutaneous tissue, fascia, and muscle (are usually spared b/c the generous blood supply).

* Necrotizing fasciitis (involved in the fascia of the muscle n the overlying subcutaneous fat) .

* Clostridial myonecrosis (when the clostridial infections spread to affect deep muscle tissue. Complications include pain , gas gangrene (gas production n build up within muscle tissue leading to tissue death 'necrosis')).

Pyomyositis/ Introduction

- **Pyomyositis** is a **purulent** infection of skeletal muscle that arises from **hematogenous** spread, usually with **abscess**(collection of pus)formation. It is classically an infection of the tropics, although it has been recognized in temperate climates with increasing frequency.
- Risk factors for pyomyositis include **immunodeficiency** (particularly HIV infection), **trauma**, **injection drug use**, concurrent infection, and **malnutrition**
- **Staphylococcus aureus** is the most common cause of pyomyositis; it causes up to 90 percent of tropical cases and up to 75 percent of temperate cases.
- **Pyomyositis** presents with **fever and pain with cramping localized to a single muscle group**. It develops most often in the lower extremity.

Pyomyositis :infection of the muscle

Pyomyositis

The Dr literally said u can read it by yourself it's not that important really. So...

Pyomyositis can be divided into three clinical stages:

- Stage 1 is characterized by crampy local muscle pain, swelling, and low-grade fever.
- Stage 2 occurs 10 to 21 days after the initial onset of symptoms and is characterized by fever, exquisite muscle tenderness, and edema.
- Stage 3 is characterized by systemic toxicity. The affected muscle is fluctuant. Complications of *S. aureus* bacteremia such as septic shock, endocarditis, septic emboli, pneumonia, pericarditis, septic arthritis, brain abscess, and acute renal failure can occur

Pyomyositis

- **Radiographic imaging** with magnetic resonance imaging is **the most useful tool** for diagnosing pyomyositis, defining the site(s) of infection, and for ruling out other entities. Bacteriologic diagnosis can be made by cultures of drainage specimens and/or blood.
- Although stage 1 pyomyositis **can be treated with antibiotics alone** , (debridement is necessary) most patients present with stage 2 or 3 disease and therefore require both antibiotics and drainage for definitive management.



<https://intjem.biomedcentral.com/articles/10.1007/s12245-008-0067-6>

Diabetic foot infections/ introduction

- Diabetic foot infections are associated with substantial morbidity and mortality.
- Important risk factors for development of diabetic foot infections include neuropathy, peripheral vascular disease, and poor glycemic control.
- In the setting of sensory neuropathy, there is **diminished perception of pain** and temperature; thus, many patients are slow to recognize the presence of an injury to their feet. Autonomic neuropathy can cause **diminished sweat secretion resulting in dry, cracked skin** that facilitates the entry of microorganisms to the deeper skin structures. In addition, motor neuropathy can lead to **foot deformities**, which lead to pressure-induced soft tissue damage.
- **Peripheral artery disease can impair blood flow necessary for healing** of ulcers and infections.
- **Hyperglycemia impairs neutrophil function and reduces host defenses.** Trauma in patients with one or more of these risk factors precipitates development of wounds that can be slow to heal and predispose to secondary infection.

Diabetic foot infections



- Fig. 1. Large forefoot ulcer in patient with diabetes. (Courtesy of William DeCarbo, D.P.M., Columbus, OH.)



Fig. 3. Diabetic foot with abscess undergoing surgical irrigation and debridement. (Courtesy of William DeCarbo, D.P.M., Columbus, OH.)

Diabetic foot infections/ Microbiology

- Most diabetic foot infections are **polymicrobial**, with up to five to seven different specific organisms often involved. The microbiology of diabetic foot wounds is variable depending on the extent of involvement.
- **Superficial diabetic foot** infections (including cellulitis and infected ulcers in antibiotic-naïve individuals) are likely due to aerobic **gram-positive cocci** (including *Staphylococcus aureus*, *Streptococcus agalactiae*, *Streptococcus pyogenes*, and coagulase-negative staphylococci).
- **Ulcers that are deep**, chronically infected, and/or previously treated with antibiotics are more likely to be **polymicrobial**. Such wounds may involve the above organisms in addition to enterococci, Enterobacteriaceae, *Pseudomonas aeruginosa* (resistant to many antibiotics in the diabetic patient has non-healing deep ulcers that needs antibiotics), and anaerobes.
- **Wounds with extensive local inflammation, necrosis, malodorous drainage**, or gangrene with signs of systemic toxicity **should be presumed to have anaerobic organisms** in addition to the above pathogens. Potential pathogens include anaerobic streptococci, **Bacteroides** species, and **Clostridium** species

Diabetic foot infections/ clinical manifestation

- Diabetic foot infections can develop as a result of **neuropathic or ischemic ulcers, traumatic wounds, skin cracks or fissures**, or other defects in the skin of the foot or nail beds (paronychia).
- Thus, infection can present as localized **superficial skin involvement** at the site of a preexisting lesion or as infection of the skin or deeper skin structures that has spread beyond the site of local trauma. Such infections can subsequently extend to **joints, bones, and the systemic circulation**.
- Diabetic foot infections are often accompanied by the cardinal manifestations of inflammation (**erythema, warmth, swelling, and tenderness**) and/or **the presence of pus** in an ulcer or sinus tract.
- **Osteomyelitis** can occur in the setting of a diabetic foot wound with or without evidence of local soft tissue infection.

Diabetic foot infections/ management

- The evaluation of a patient with a suspected diabetic foot infection involves three key steps: **1) determining the extent and severity of infection**, **2) identifying underlying factors** that predispose to and promote infection, and **3) assessing the microbial etiology**.
- Clinical examination should note the location of the lesions, extent of infection (eg, involving skin, subcutaneous tissue, muscles, tendons and/or bone) and whether bone is grossly visible or palpable by probing. Although osteomyelitis is highly likely if bone is visible, it may be present in the absence of such findings.
- Clinical examination should also include a neurologic evaluation that documents the extent of sensory loss as well as a vascular evaluation.

Clinical classification of a diabetic foot infection

Infection severity	Clinical manifestations of infection
Uninfected	Wound lacking purulence or any manifestations of inflammation.
Mild	Presence of ≥ 2 manifestations of inflammation (purulence, or erythema, pain, tenderness, warmth, or induration), but any cellulitis/erythema extends ≤ 2 cm around the ulcer, and infection is limited to the skin or superficial subcutaneous tissues; no other local complications or systemic illness.
Moderate	Infection (as above) in a patient who is systemically well and metabolically stable but which has ≥ 1 of the following characteristics: cellulitis extending > 2 cm, lymphangitic streaking, spread beneath the superficial fascia, deep-tissue abscess, gangrene, and involvement of muscle, tendon, joint or bone.
Severe	Infection in a patient with systemic toxicity or metabolic instability (eg, fever, chills, tachycardia, hypotension, confusion, vomiting, leukocytosis, acidosis, severe hyperglycemia, or azotemia).

Foot ischemia may increase the severity of any infection, and the presence of critical ischemia often makes the infection severe.

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Graphic 55551 Version 1.0

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Diabetic foot infections/ management

- **Wound management** —Local wound care for diabetic foot infections typically includes debridement of callus and necrotic tissue, wound cleansing, and relief of pressure on the ulcer.
- **Obtaining samples for culture** —Because microorganisms often colonize lower extremity wounds regardless of the presence of a true infection, cultures should be performed only in selected patients. If the clinical suspicion for infection is low, samples from the wound should not be submitted for culture. The preferred clinical specimens for reliable culture include **aspirate from an abscess or curettage from the ulcer base**.
- **Surgery** —Consultation with a surgeon with experience in diabetic foot infections is important for cases of severe infections and in most cases of moderate infections.
- **Antimicrobial therapy** —Empiric antibiotic therapy should be selected based on the severity of infection and the likelihood of involvement of resistant organisms

Further reading:

- Necrotizing soft tissue infections

[https://www.uptodate.com/contents/necrotizing-soft-tissue-infections?topicRef=3993&source=see link#H3846414689](https://www.uptodate.com/contents/necrotizing-soft-tissue-infections?topicRef=3993&source=see_link#H3846414689)

- *Clostridial myonecrosis*

[https://www.uptodate.com/contents/clostridial-myonecrosis?topicRef=7662&source=see link](https://www.uptodate.com/contents/clostridial-myonecrosis?topicRef=7662&source=see_link)

- Pyomyositis

[https://www.uptodate.com/contents/pyomyositis?topicRef=7662&source=see link](https://www.uptodate.com/contents/pyomyositis?topicRef=7662&source=see_link)

- *Diabetic foot infections*

<https://ezlibrary.ju.edu.jo:2119/contents/clinical-manifestations-diagnosis-and-management-of-diabetic-infections-of-the-lower->

V2

Cellulitis



Erysipelas



Slide number 9
Right pic is erysipelas
Left one is cellulitis ..