



Host Defense: Where the Rubber of Immunology Hits the Road of Life

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Disclosures of Financial Relationships with Relevant Commercial Interests

- List of disclosures or “None”



Concepts of Hell

Host Immune Defense

Humoral

- Complement
- Mannose binding lectin
- Antibody

Cellular

- Neutrophils
- Monocytes
- Eosinophils
- Lymphocytes (NK, T, B)
- Other (erythrocytes, platelets)

Basic Principles

Patients with impaired inflammation:

- May be unable to tell you they are sick (feel fine)
- Are often sicker than they look
- Often have more extensive disease than is apparent
- May require longer treatment than normals
- May have unusual infections

In vitro testing is tricky and variable, genetics is not

Who's Got a Problem?

Abnormal frequency of infections

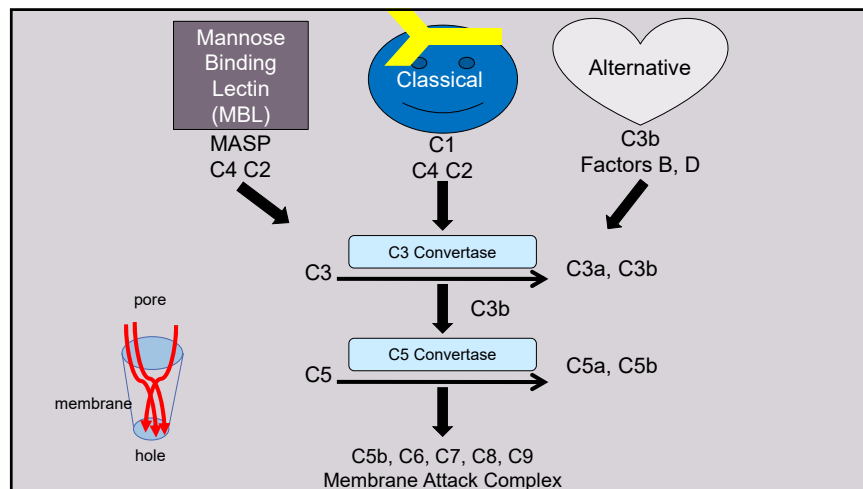
- *Recurrent Neisseria* bacteremia, pneumonia

Abnormal presentation of infections

- Necrotic cutaneous ulcers (not anthrax)
- *Aspergillus* pneumonia

Unusual infections

- *Pneumocystis jirovecii*
- *Burkholderia cepacia* complex
- *Nontuberculous mycobacteria*



Complement Deficiencies

Classical Pathway (C1-C9) (Recessive)

- Antibody dependent bacterial lysis
- Deficiency leads to recurrent bacteremia and meningitis

Alternative Pathway (Factors I, H, Properdin, C3)
(Properdin X-linked, others AR)

- Antibody independent bacterial lysis
- More severe than classical defects

Mannose Binding Lectin (MBL) Pathway

- Very modest IF ANY defect, mild effect in infancy

Complement Defects

C5-C9 Defects

- Recurrent *Neisseria* bacteremia and meningitis
- Average age of onset 17 y, milder CNS sequelae
- High rates of relapse and reinfection

C1-C4 Defects

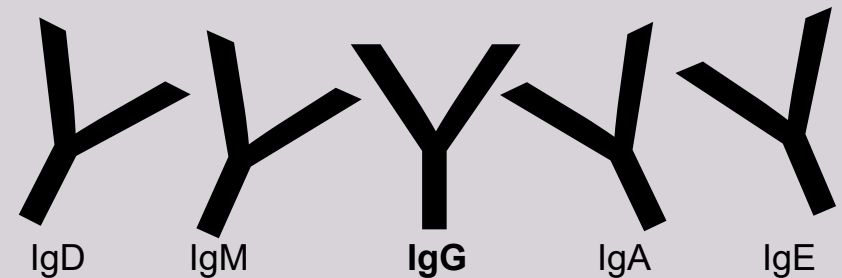
- Autoimmune disease (SLE, DLE) more common

Dx - CH₅₀ (Classical), AH50 (Alternative)

Rx - treat infections, prophylaxis if needed, hypervaccination?

J Clin Immunol 2020 May;40(4):576-591

Let's Talk About Antibodies!!! Five Flavors



Antibody Deficiencies

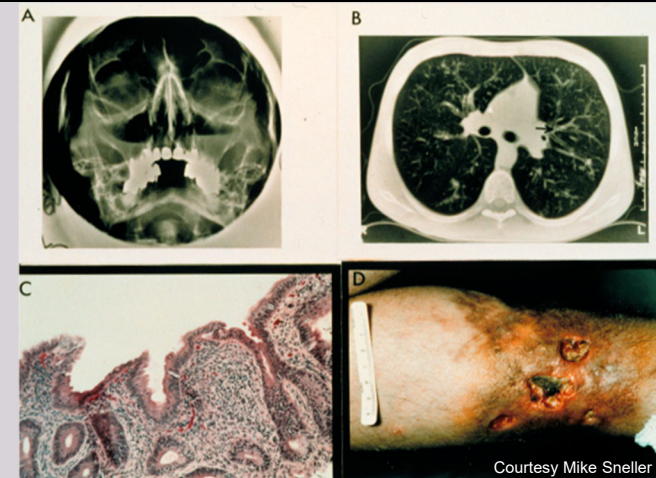
IgA Deficiency (AR)

- Common (1/700 adults)
- Probably not a pathologic condition per se
- Frequently associated with other deficits, such as common variable immunodeficiency (CVID), Ig subclass deficiencies, allergies, etc.

Dx - Low IgA

Rx - None

J Transl Autoimmun 2019 Nov 23;2:100025



Courtesy Mike Sneller

Common Variable Immunodeficiency (CVID)

Recurrent sino-pulmonary bacterial infections

Chronic enteric *Giardia*, *Campy*, *Salmonella*, *Shigella*

Severe enteroviral meningitis/encephalitis/myositis

Nodular regenerative hyperplasia of liver

Dx -  IgG (total and subclasses 1,3 or 2,4),
IgA, IgM, isohemagglutinins, DTH,
Impaired response to new or recall immunization

Rx - Autoimmunity and cancer
Treat infections, Ig replacement

Cunningham-Rundles C. Immunol Rev. 2019 Jan;287(1):145-161.

Question #1

47-year-old woman

- Recurrent episodes of bronchitis, recently more exacerbations; tired
- One episode of documented bacterial pneumonia and sinusitis
- Immunoglobulin levels:
 - IgG 500 (normal 523-1482)
 - IgA <10 (normal 51-375)
 - IgM 165 (normal 37-200)

Question #1

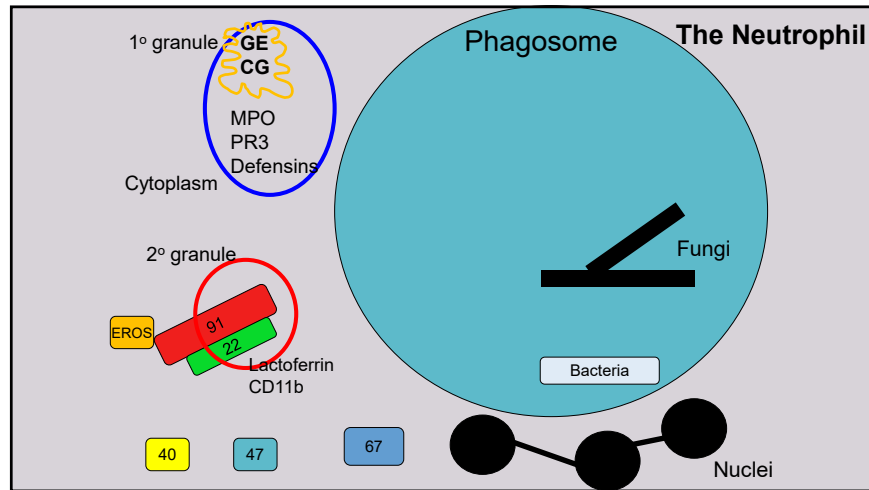
What is the next step?

- A. IgG subclasses and titers against tetanus and pneumococcus. If low, consider IVIG.
- B. Repeat IgG levels. If low, consider IVIG.
- C. Skin tests for DTH. If anergic, consider IVIG.
- D. Titers against tetanus and pneumococcus, immunize, and repeat. If low, consider IVIG.
- E. Check MBL levels. If low, consider IVIG.

Question #1

What is the next step? Refer to Immunology

- A. IgG subclasses and titers against tetanus and pneumococcus. If low, consider IVIG.
- B. Repeat IgG levels. If low, consider IVIG.
- C. Skin tests for DTH. If anergic, consider IVIG.
- D. Titers against tetanus and pneumococcus, immunize, and repeat. If low, consider IVIG.***
- E. Check MBL levels. If low, consider IVIG.



Question #2

Neutrophils: Nobody cares until they are not around.

- Average count 5000/mcl
 - (5,000,000/ml)
 - (5,000,000,000/L)

So, how many do you make a day?

Question #2

How many neutrophils do you make a day?

- 10e8; the half-life is 10 days
- 10e9; the half-life is 5 days
- 10e10; the half-life is 1 day
- 10e11; the half-life is 7 hours
- 10e6; they live until you use them up

Question #2

How many neutrophils do you make a day?

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- 10e10; the half-life is 1 day
- D. 10e11; the half-life is 7 hours***
- 10e6; they live until you use them up

Neutrophils: Always Fresh, Always Ready!

D. 10e11; the half-life is 7 hours*

- Make around 10¹¹/day
- Most are in bone marrow
- Can go up 10-fold in emergency
- Circulating half life 7 hours
- About 50% marginated

Cyclic and Severe Chronic Neutropenias

Cyclic and SCN: *ELANE* mutations (AD)

Kostmann SCN: *HAX1* mutations (AR)

- Digital, oral, perineal infections, usually self-healing with recovery of counts, bacteremia uncommon
- Relatively low baseline PMN count with profound neutropenia, about every 3-4 weeks

Dx - Molecular; periodicity, family history, genetics

Rx - G-CSF, BMT

Hematol Oncol Clin North Am. 2019;33:533-551

Other Causes of Neutropenia

X-linked

WAS

GATA1

TAZ

Dominant

GF11

ELA2

GATA2

DNM2

SRP54

CXCR4

Recessive

G6PC3

HAX1

JAGN

USB1

CSF3R

VPS45

GSD1B

SBDS

Big Causes Drugs

Splenomegaly/ sequestration

Autoimmunity

Case Study

52-year-old man

- Referred from his Family Practitioner
- Recurrent digital and oral ulcers occurring every month or so for the last 4 months
- One CBC showed an ANC of 100, but on repeat several days later was normal
- Previous health good
- Took "some antibiotic for a cold a few months ago"
- Spleen tip felt



Acquired Neutropenia in Adults

- Drugs, lupus, etc.
- Acquired cyclic neutropenia
(Large Granular Lymphocytosis, LGL)
Splénomegaly, often associated with Rheumatoid arthritis (Feltz Syndrome)
- **Dx** - Clonal CD3+/8+/57+ lymphs (LGL)
(Gain of Function mutations in STAT3)
- **Rx** - Treatment of the abnormal clone can be curative
(Cyclosporine, MTX, steroids, alemtuzumab)
G-CSF may lift both nadir and baseline



Hematol Malign Rep. 2020 Apr;15(2):103-112

Myeloperoxidase (MPO) Deficiency (AR)

Most common neutrophil disorder (1/2000)

- Not a pathologic condition per se
- Failure of H_2O_2 -----MPO-----> HOCl
- Compensated by increased H_2O_2 production
- Appears to need another condition to potentiate, such as diabetes

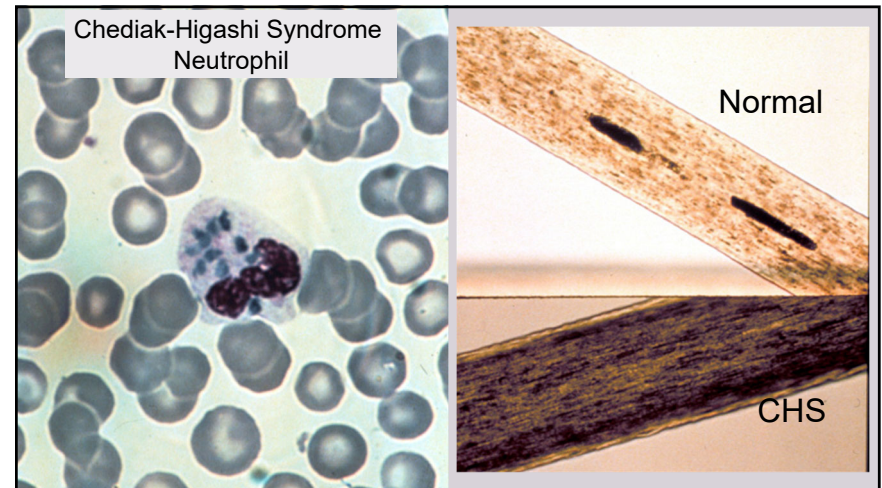
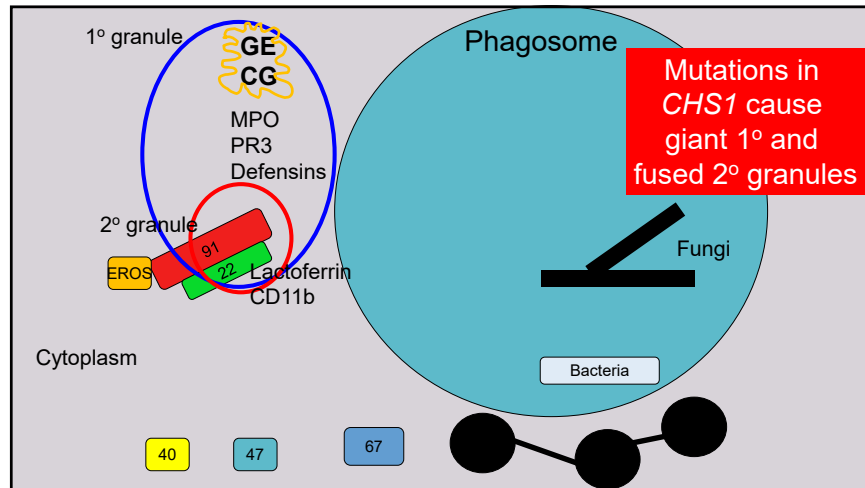
Dx - Absence of peroxidase positive granules due to mutations in *MPO* gene

Rx - Treat invasive infections (*Candida*), no specific therapy

J Leukoc Biol. 2013 Feb;93(2):185-98



Chediak-Higashi Syndrome



Chediak-Higashi Syndrome (Recessive)

Recurrent cutaneous, sino-pulmonary infections

- GNR, staph, strep, no fungi
- Mild neutropenia (intramedullary destruction)

Partial oculocutaneous albinism,

Mental retardation, neuropathy (late)

Lymphoma or HLH-like “accelerated phase” (late)

Dx - Giant blue PMN granules or chaotic hair granules
due to mutations in CHS1, encodes LYST

Rx - Prophylaxis, treatment of infections, BMT

Drug Discov Today Dis Models. 2020;31:31-36

Silvery Sheen



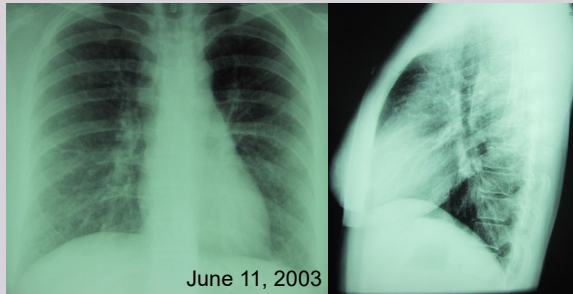
Peripheral Neuropathy



Case Study

23-year-old woman; athletic coach

Previously healthy; short of breath 4 hours after 3-mile run



ER Presentation

- Recent weekend with friends in NYC
- Anxious, chest pressure, febrile
- Acute mononucleosis?

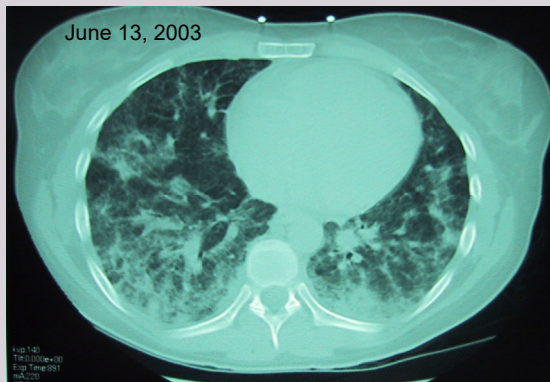
PMH

- Respiratory infections in infancy
- Cat scratch disease 8-year-old: resolved with antibiotics

Family History

- 1 brother with two episodes Cat scratch cervical nodes
- 2 sibs well

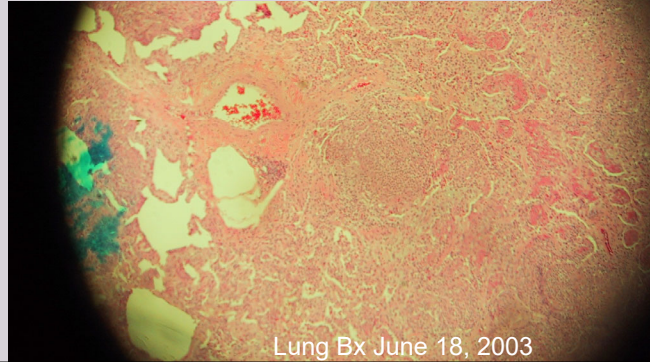
2 Days Later, Hypoxia And Fever



Hospital Course

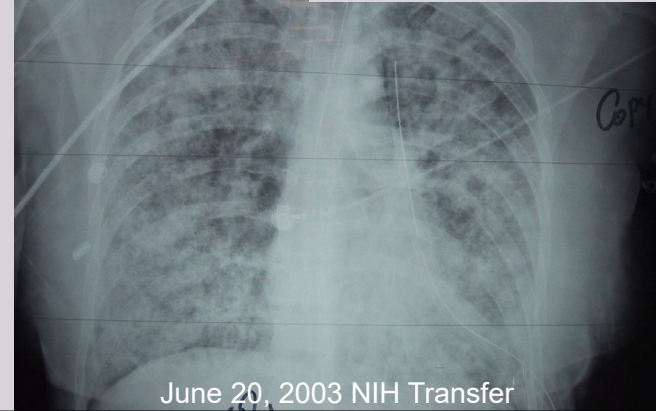
- Progressive dyspnea, fever, leukocytosis
- Refractory to antibiotics and steroids
- Bronchoscopy uninformative
- Visually Assisted Thoracoscopic Surgery (VATS)

8 days after presentation:
Intubation and lung biopsy
necrotizing granulomata and hyphae



Lung Bx June 18, 2003

10 days after presentation:
Biopsy growing *A. fumigatus*



June 20, 2003 NIH Transfer

Question #3

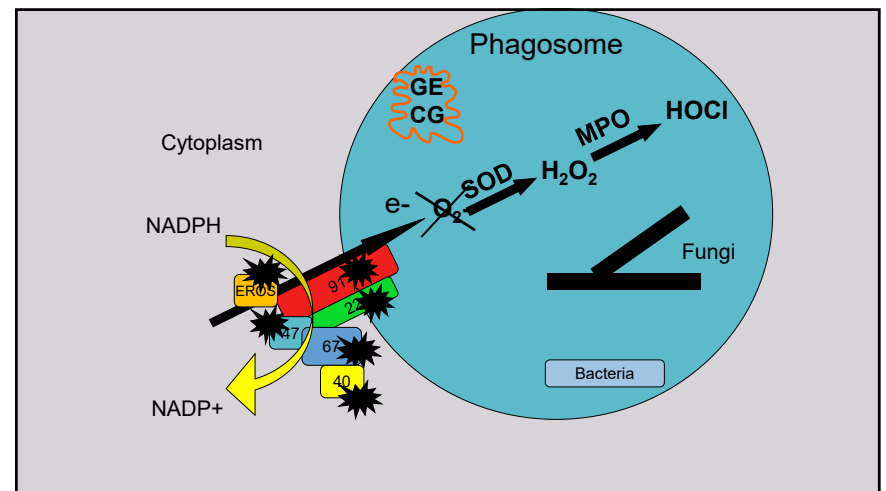
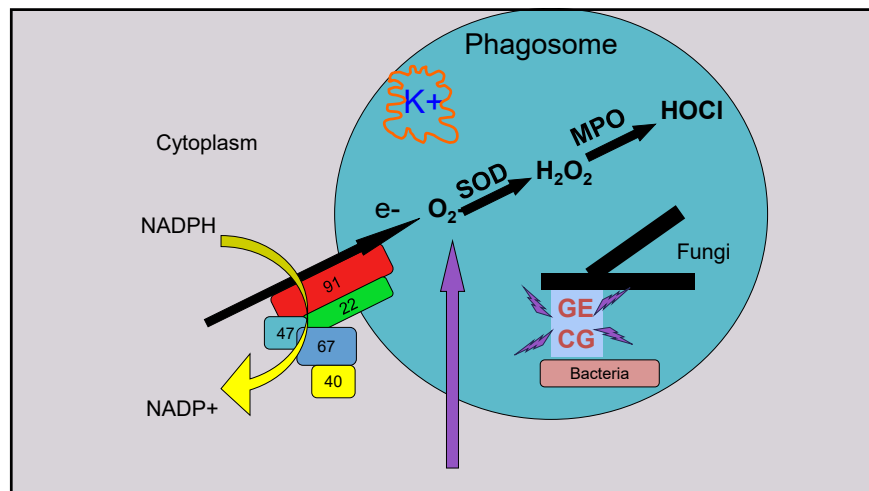
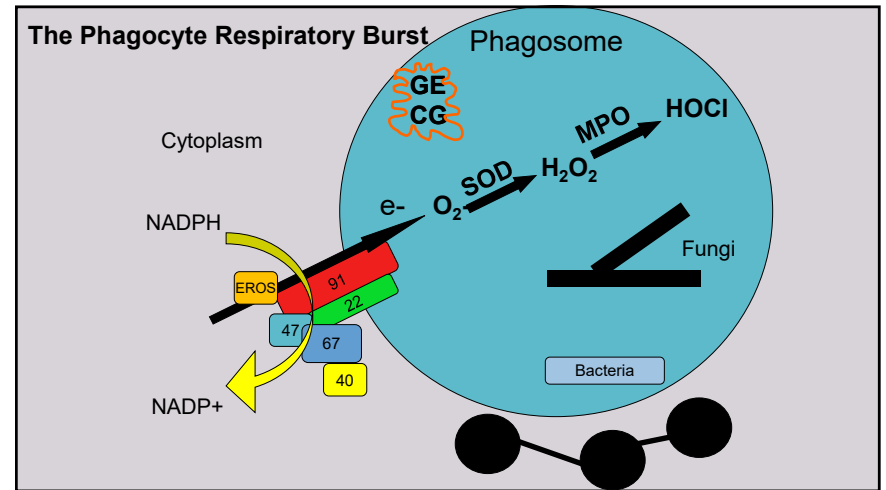
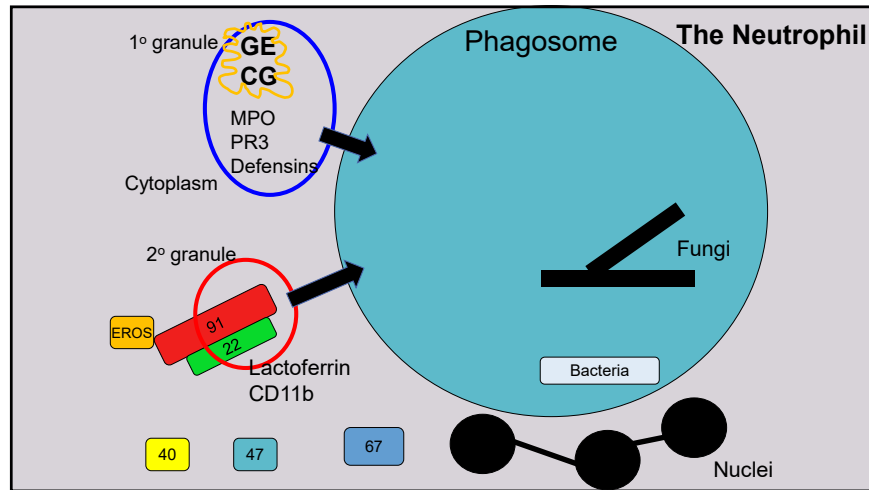
Which of the following is a cause of invasive aspergillosis in an otherwise normal host?

- A. Allergic bronchopulmonary aspergillosis
- B. Cystic fibrosis
- C. Lymphocyte dysfunction (SCID)
- D. Phagocyte defect
- E. Acute HIV

Question #3

Which of the following is a cause of invasive aspergillosis in an otherwise normal host?

- A. Allergic bronchopulmonary aspergillosis
- B. Cystic fibrosis
- C. Lymphocyte dysfunction (SCID)
- D. Phagocyte defect***
- E. Acute HIV



Chronic Granulomatous Disease (X, AR)

Failure to make the phagocyte respiratory burst

Frequency 1/100,000 - 1/200,000 live births

- Presentation usually in childhood; more adults recognized

Recurrent life-threatening infections

- Catalase-positive bacteria, fungi (this is nuanced)
- Tissue granuloma formation

Infections: Lung, liver, lymph nodes, skin, bone

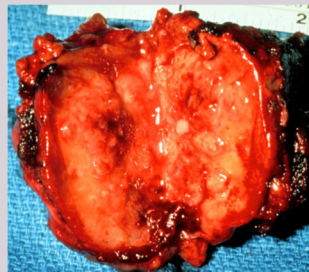
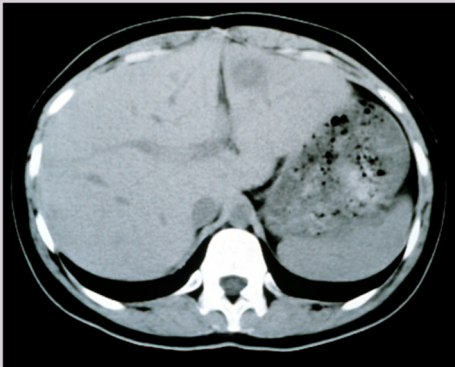
Bacteremia: Uncommon but bad

Infections in CGD

<i>S. aureus</i>	(liver, lymph nodes, osteo)
<i>S. marsecens</i>	(skin, lung, lymph nodes)
<i>B. cepacia</i>	(pneumonia, bacteremia)
<i>Nocardia spp.</i>	(pneumonia, brain, liver)
<i>Aspergillus spp.</i>	(lung, esp. miliary, spine)
<i>Salmonella</i>	(enteric, bacteremia)
<i>BCG</i>	(local/regional infections)
<i>Chromobacterium violaceum</i>	(warm brackish water, soil, e.g., Disney World)
<i>Francisella philomiragia</i>	(brackish water, Chesapeake Bay, Sounds)
<i>Burkholderia gladioli</i>	(causes onion rot)
<i>Granulibacter bethesdensis</i>	(necrotizing LN, hard to grow, likes CYE)
<i>Paecilomyces spp.</i>	

Pediatric Health Med Ther 2020 Jul 22;11:257-268

Staphylococcal Liver Abscess in CGD



CID 2018;66:1427-1434

Staphylococcal Lymphadenitis in CGD



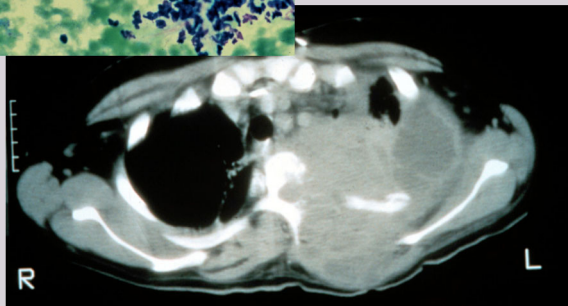
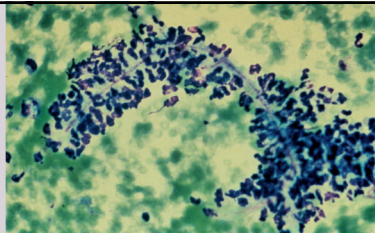
***Staph aureus* Osteomyelitis in CGD**



***Burkholderia cepacia* Complex Bacteremia in CGD**



**CGD
Aspergillus nidulans
Pneumonia**

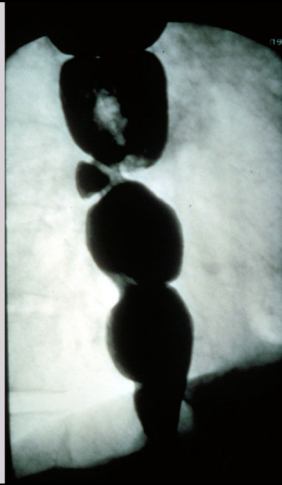


Thoracotomy 10 Days Postop



CGD Granulomatous Wound Dehiscence

CGD Granulomatous Esophageal Obstruction



CGD Inflammatory Bowel Disease



Chronic Granulomatous Disease

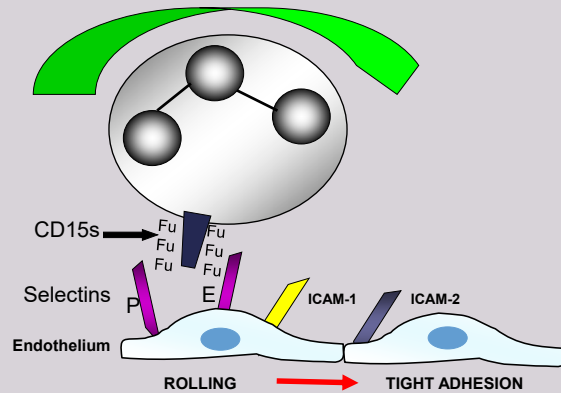
- X-linked, chr. Xp21 (70% of US cases)
 - Carrier females are mosaic (Lyonization)
 - 1/2 of offspring of carrier Mom will receive the gene
 - ~1/3 of carriers are sporadic, from sperm
 - X-linked male: all daughters carriers, no sons affected
 - Autosomal recessive (30% of cases)
- Dx** - PMN dihydrorhodamine 123 oxidation (DHR)
[PMN nitroblue tetrazolium reduction (NBT) is the old test]
(MPO Deficiency gives a FALSE ABNORMAL DHR)
- BE CAREFUL ABOUT THE LAB AND HOW YOU DISCUSS IT!

CGD Management and Treatment

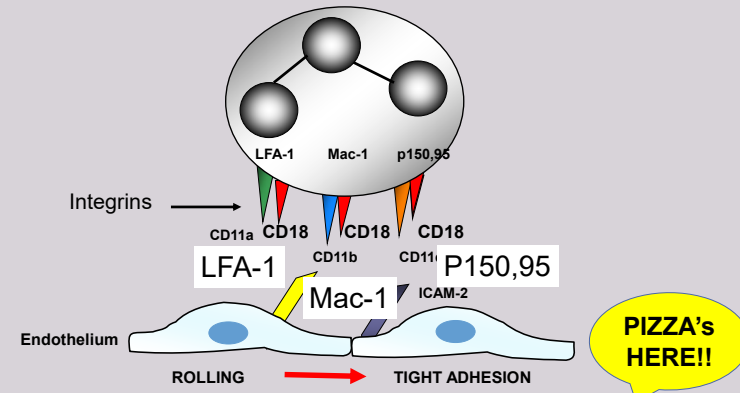
- 90% overall long-term survival
- Follow CRP, radiographs
- Prophylactic antibiotics and antifungals
 - TMP/SMX, itraconazole
- Prophylactic interferon gamma
 - 50 µg/m² subcutaneously three times weekly
- Aggressive search for and treatment of infections
- BMT (gene therapy)

Hematol Oncol Clin North Am. 2013 Feb;27(1):89-99

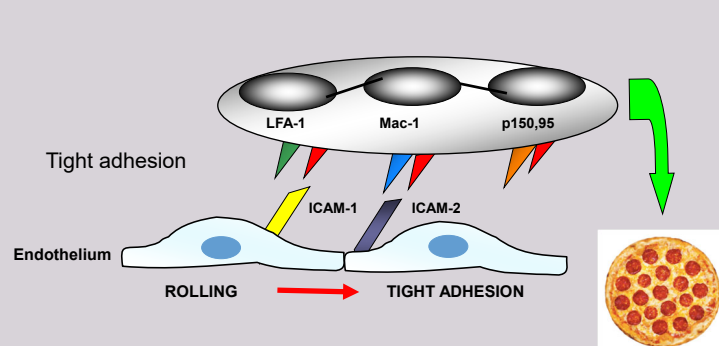
Neutrophil Rolling



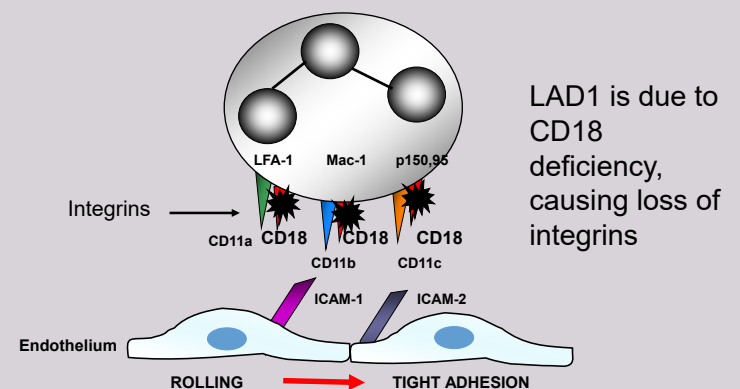
Neutrophil Adhesion



Neutrophil Tissue Entry (Diapedesis)



Leukocyte Adhesion Deficiency Type 1



Leukocyte Adhesion Deficiency Type 1 (AR)

- Failure to attach to the endothelium due to mutations CD18
- Recurrent necrotizing infections: skin, perineum, lung, gut
- Enteric GNR, GPC, NOT fungi or Candida
- Baseline leukocytosis, further WBC increase to infection
- Rare, consanguinity common

Dx - FACS for CD18, genetics

Rx - Treatment of infections, BMT

Leukocyte Adhesion Deficiency I

- Delayed umbilical stump separation
- Dystrophic, “cigarette paper” scars
- Gingivitis with tooth loss, alveolar ridge resorption
- Biopsies show no neutrophils at sites of infection, rare monocytes and eosinophils
- Severe and moderate forms of disease

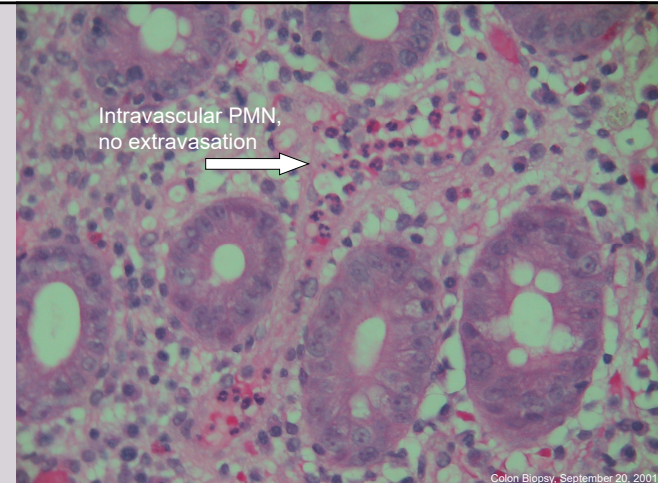
Almost Universal Tooth Loss in LAD1 By Adulthood



Impaired Wound Healing in LAD1



Cigarette Paper Scarring



Question #4

19-year-old boy with Pneumonia

- Admission WBC 43,000, looked OK
- Ceftriaxone, good response
- Medical student: WBC never <11,000/mcl
- Left shin ulcer not inflamed
- Not healed in > 2 mos
- She raises the possibility of Leukocyte Adhesion Deficiency (LAD1)

Question #4

Which of the following would rule against LAD1?

- Gingivitis, tooth loss, and alveolar ridge resorption
- FACS showing 5% of normal expression of CD18 and CD11a-c on granulocytes
- He is the product of a first cousin union
- Extensive neutrophil infiltration in the left shin ulcer
- Multiple dystrophic scars over the legs from previous ulcers

Question #4

Which of the following would rule against LAD1?

- A. Gingivitis, tooth loss, and alveolar ridge resorption
- B. FACS showing 5% of normal expression of CD18 and CD11a-c on granulocytes
- C. He is the product of a first cousin union
- D. Extensive neutrophil infiltration in the left shin ulcer***
- E. Multiple dystrophic scars over the legs from previous ulcers

27-year-old Woman with Boils

Referred for recurrent boils with *S. aureus*

- IgE of 12,376 IU
- “Bronchitis and sinusitis at least once a year”
- Persistent eczema requiring topical steroids
- Never hospitalized but having “more trouble” lately

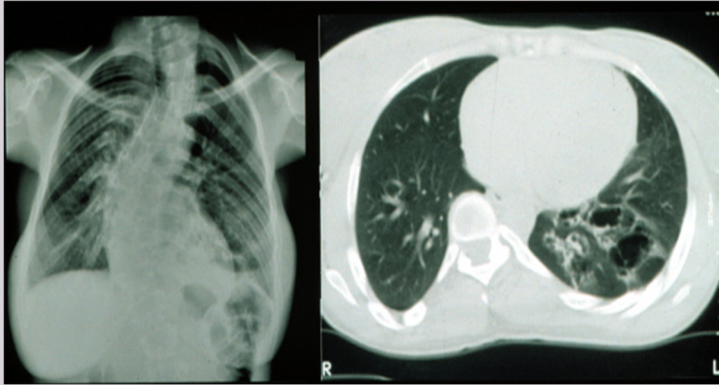
Job's Syndrome



N Engl J Med. 1999;340:692-702

HIE (Job's) Syndrome History and Exam

Eczema	100%
Facies	100% (>16y)
Boils	87%
Pneumonia	87%
Mucocutaneous Candidiasis	83%
Pulmonary Cysts	77%
Scoliosis	76% (> 16y)
Delayed dental deciduation	72%
Coronary artery aneurysms	65%
Pathologic fractures	57%



Pulmonary Pathogens in HIE

Primary Pathogens:

- *Staphylococcus aureus*
- *Streptococcus pneumoniae*
- *Hemophilus influenzae*

Secondary Pathogens:

- *Pseudomonas aeruginosa*
- *Aspergillus fumigatus*

Others:

- *Pneumocystis jirovecii*, *M. avium* complex

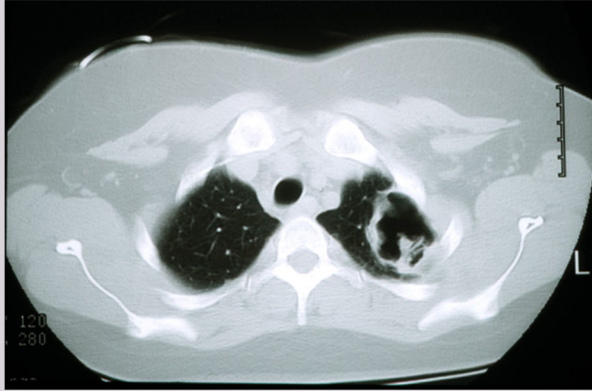
Candida



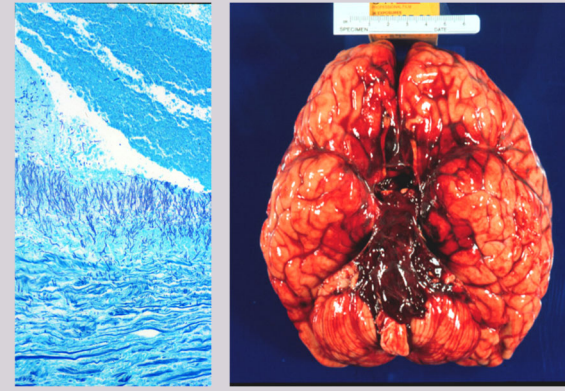
Group A Strep



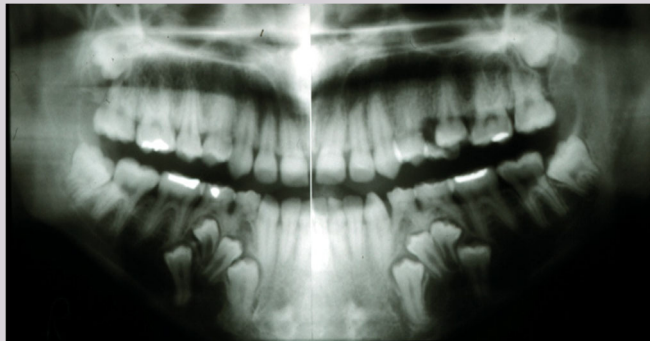
Aspergillus Lung Infection



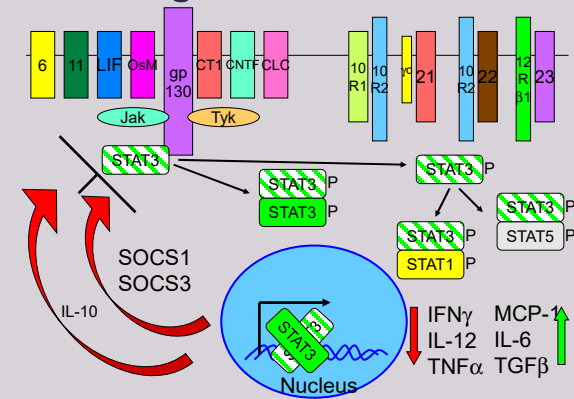
Metastatic Aspergillus Brain Infection



Failure Of Primary Dental Deciduation



Dominant Negative Mutant STAT3



Hyper IgE Recurrent Infection (Job's)

- Recurrent sinopulmonary *S. aureus*, *S. pneumo*, *H. flu*
 - Post-infectious pulmonary cyst formation
 - Recurrent *S. aureus* skin abscesses
 - Characteristic facies, eczema, scoliosis, fractures
 - Very elevated IgE (>2000 IU), modest eosinophilia
- DDx** - Atopic dermatitis is a close mimic
Job's: pneumonia, lung cysts, skeletal, mutations in STAT3
- Rx** - Treatment of infections, prophylactic antibiotics, antifungals increasingly BMT

J Clin Immunol. 2021;41:864-880

Close Mimic: DOCK8 Deficiency

Autosomal Recessive hyper IgE syndrome

Eczema, allergies, asthma, high IgE
Staph, *Strep*, *H. flu*, *Acinetobacter*, *Pseudomonas*
Candida, *Cryptococcus*, *Histoplasma*
HPV, **HSV**, **molluscum**
Squamous cell carcinomas, lymphoma

J Clin Immunol 2021 May 1. doi: 10.1007/s10875-021-01051-1

DOCK8 Deficiency



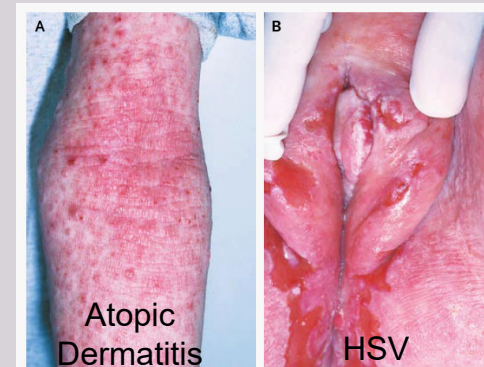
HPV



Molluscum
contagiosum

N Engl J Med. 2009;361:2046-55

DOCK8 Deficiency



Atopic
Dermatitis



HSV

DOCK8 vs. STAT3 Hyper IgEs

	DOCK8 (Recessive)	STAT3 (Dominant)
Pneumonia	+	+++
Pneumatocoles	-	+++
Retained teeth	-	+++
Fractures	-	+++
Viral infections	+++	-
Fungal infections	+	++
Allergies	+++	-
IgM	Low	Normal
Eosinophils	+ to ++	+

Question #5

15-year-old girl with recurrent infections

- Infancy: eczema, recurrent pneumonias, skin infections
- IgE 14,574 IU/ml
- Allergist: use bed covers to avoid dust mites

Going over the allotted 15 minutes, you elicit points trying to establish whether she has hyper-IgE recurrent infection syndrome (Job's).

Question #5

Which one of the following is not supportive of the diagnosis of Job's (STAT3DN)?

- A. Pneumatocoles
- B. Scoliosis
- C. Severe warts
- D. Retained baby teeth
- E. Recurrent fractures

Question #5

Which one of the following is not supportive of the diagnosis of Job's (STAT3DN)?

- A. Pneumatocoles
- B. Scoliosis
- C. Severe warts***
- D. Retained baby teeth
- E. Recurrent fractures

Clinical Spectrum of NTM Infections

Disseminated

Severe, Young
IFN γ /IL-12 defects
NEMO, STAT1



IMMUNE

Skin

Exposure
Inoculation



EXPOSURE

Pulmonary

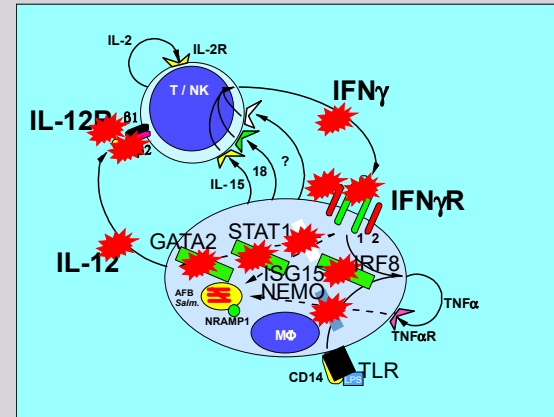
Chronic, Older
Bronchiectasis
Cystic fibrosis (CF)
Ciliary dyskinesia (PCD)



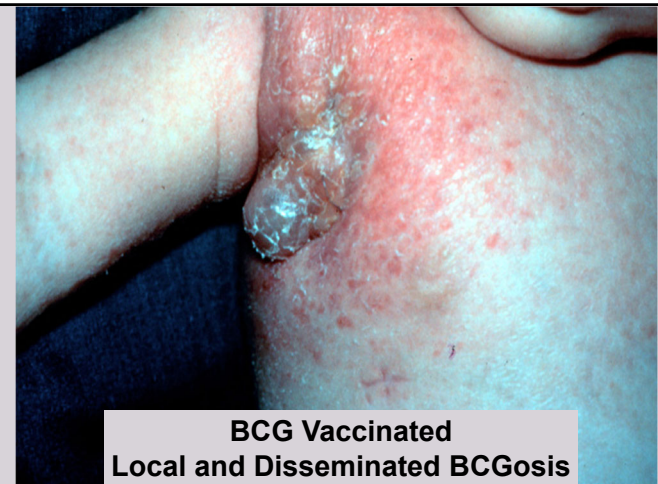
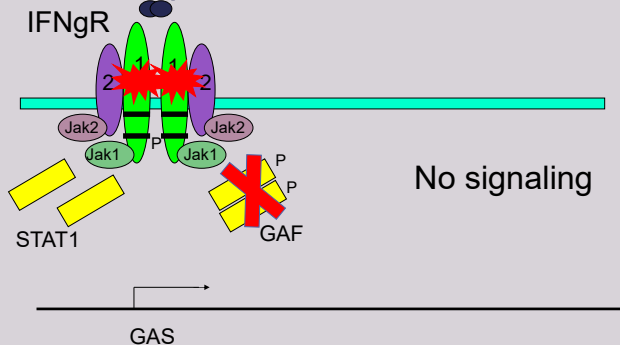
EPITHELIAL

Lancet Infect Dis. 2015;15:968-80

Disseminated NTM Only

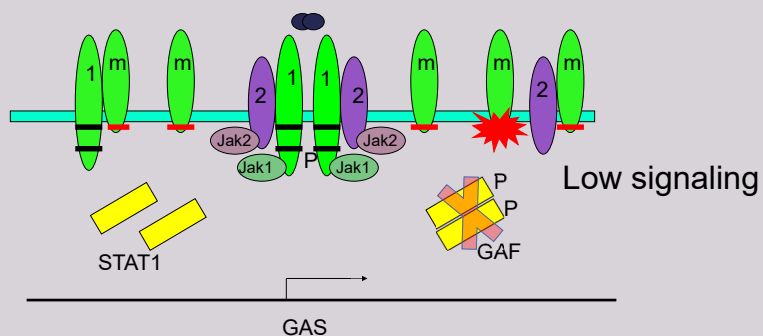


Autosomal Recessive IFNGR1 (Both Alleles)

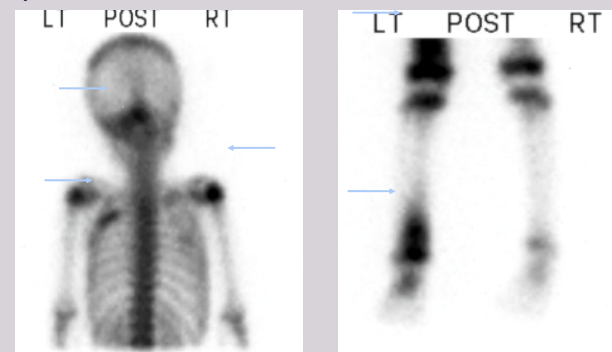


BCG Vaccinated
Local and Disseminated BCGosis

Autosomal Dominant IFNGR1 (One Allele)



Mycobacterial Osteomyelitis in Dominant IFN γ R1 Deficiency



Ann Allergy Asthma Immunol 2002;89:239-43

Pathogens in Human IFN γ R Deficiencies

<i>M. avium</i>	Salmonella
<i>M. intracellulare</i>	Listeria
<i>M. chelonae</i>	
<i>M. abscessus</i>	CMV
<i>M. smegmatis</i>	HSV
<i>M. fortuitum</i>	VZV
<i>M. tuberculosis</i>	RSV
<i>Bacille Calmette Guerin</i>	HHV-8
<i>Coccidioides</i>	
<i>Histoplasma</i>	

NOT JUST MYCOBACTERIA!!

IFNGR1: Dominant vs. Recessive

Characteristic	AD	AR
IFN γ R1 display	High	None
IFN γ R1 responsiveness	Low	None
Clinical presentation	Local	Disseminated
Granulomata	Present	Absent
Osteomyelitis	100%	Rare
Survival	Excellent	Most die

Lancet. 2004;364:2113-21

Interferon γ Receptor Deficiencies

Absent or defective IFN γ R1

- MAC and other NTM, *Salmonella*, TB, viruses
- Complete defects present in childhood
- Partial defects present later in life
- May be misdiagnosed as malignancy!
- NOT a cause of isolated lung disease in adults
- **Dx** - genetics, flow cytometry for IFN γ R1
- **Rx** - antimycobacterials, BMT

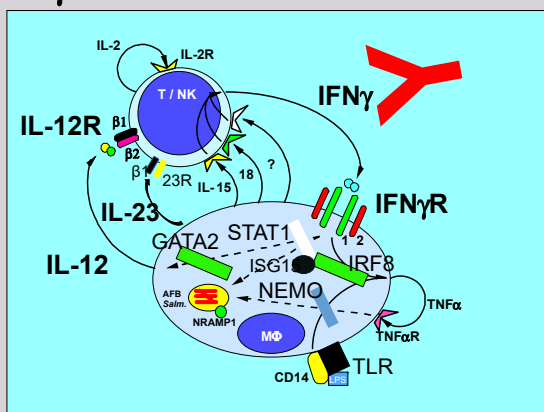
N Engl J Med. 2017;377:1077-1091.

60-year-old Vietnamese Woman

- USA 1970s
- 1 year recurring disseminated *M. avium* complex
- Numerous fistulae



Anti-IFN γ Autoantibodies

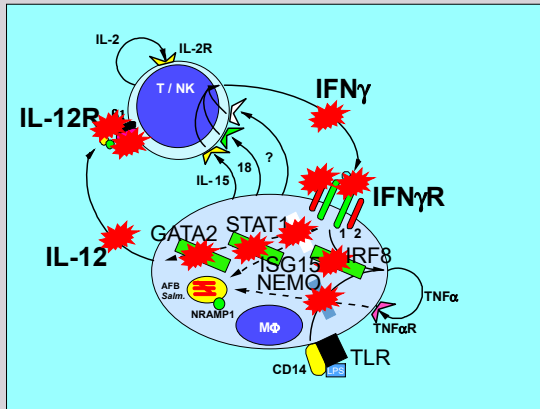


Anti-IFN γ Autoantibody Syndrome

- *Disseminated* NTM later in life also TB, *Talaromyces*, *Burkholderia*, VZV
- Predominantly female, mostly East Asian
- **Dx** - anti-IFN γ autoantibody detection
Quantiferon is often **INDETERMINATE**
- **Rx** - antimycobacterials, possibly rituximab

NEJM 2012;367:725

NOT the Pathway for Lung Disease



Question #6

30-year-old Thai Woman with Back Pain

- 2 months pain and weight loss
- HIV-, normal CBC and chemistries, normal CD4
- Biopsy: osteomyelitis, MAC growing
- Quantiferon indeterminate
- You suspect that she has the anti-interferon gamma autoantibody syndrome

Question #6

Supporting this diagnosis, what should you do?

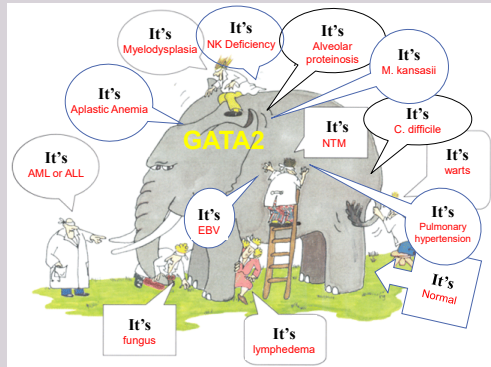
- Check complements and total IgG
- Determine anti-IFN γ antibody levels
- Determine anti-GM-CSF autoantibody levels
- Determine anti-IFN α autoantibody levels
- Determine her cellular response to IFN γ

Question #6

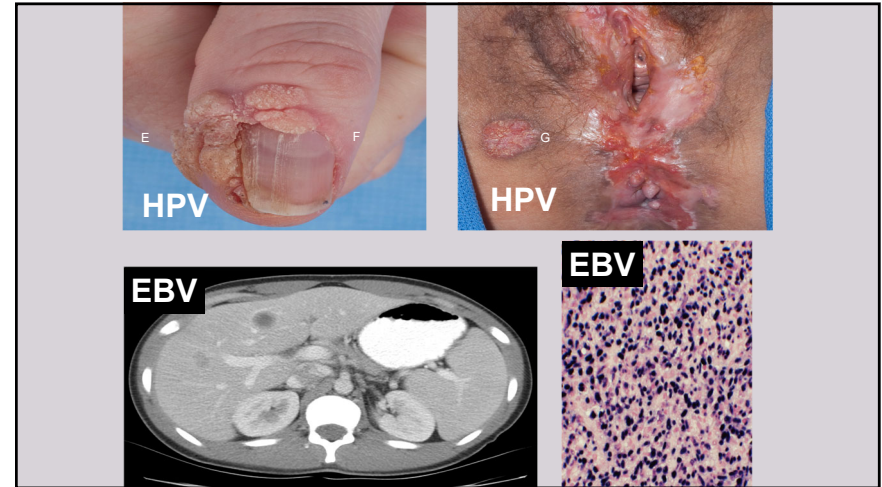
Supporting this diagnosis, what should you do?

- Check complements and total IgG
- Determine anti-IFN γ antibody levels***
- Determine anti-GM-CSF autoantibody levels
- Determine anti-IFN α autoantibody levels
- Determine her cellular response to IFN γ

The Blind



The Elephant



GATA2 Deficiency

- Mutations in GATA2, a critical hematopoietic gene
- Adolescent to adult onset
 - HPV (hands, genitals, cervical, vulvar)
 - Disseminated NTM (mediastinal *M. kansasii*)
 - Pancytopenia
- Labs: profound monocytopenia, low B, low NK
- CT: subpleural blebs
- Autosomal dominant
- Dx: genetics, hypocellular marrow, abnormal megakaryocytes
- Rx: antibiotics, BMT

Blood 2014; 123:809-21

Idiopathic CD4+ T-lymphocytopenia

- Idiopathic CD4+ T-lymphocytopenia (ICL)
 - ≤ 300 CD4+/ μ l
 - Associated with AIDS-like infections (crypto, PCP, MAC)
 - Exclude HIV infection (PCR, bDNA, p24, culture)
 - Often older onset than HIV associated OI
 - Surprisingly stable, consider incident cancers
- **Dx** - Determination of CD4 and CD8 number
 - Often due to an underlying defect, so LOOK
- **Rx** - Treat infections (follow CD4+, ?cytokines)

N Engl J Med. 2023;388:1680-1691

Screening Laboratories

- For Lymphocytes
 - Ig levels
 - Immunization status (tetanus, pneumovax)
 - CD4+ number
 - *Genetics* (exome studies, panels)

Screening Laboratories

- Phagocytes
 - DHR for CGD
 - Genetics for everything else
- Complement
 - CH₅₀ (classical pathway)
 - AH₅₀ (alternative pathway)
 - Think about the gene involved!
 - Use Pubmed OMIM
 - Sequence is faster and cheaper than you think

