

# STIs 102: Persistent NG(C)U and genital ulcer disease

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# Disclosures

- The author and their spouses/partners wish to disclose they have no financial interests or other relationships with the manufacturers of commercial products, suppliers of commercial services, or commercial supporters.

# PTC Disclaimer

**Some terms in this presentation may have been modified to align with executive order requirements that this CDC-funded grant has received.**

# Objectives

- Revisit the differential diagnosis of non-gonococcal urethritis
- Touch briefly on workup for NGU
- Describe Mycoplasma genitalium (Mgen) presentation and testing
- Discuss treatment for Mgen, including rationale, resistance concerns, and options for treatment failure
- Briefly review the differential and workup for genital ulcer disease
- Describe genital herpes presentation and epidemiology
- Discuss HSV diagnosis, treatment strategies, and transmission prevention

# Burning Questions



- Jeremy is a 35 year old man who presents to sexual health clinic where he is seen regularly for PrEP
- C/o 5 days of dysuria and mucopurulent penile discharge
- Sexual hx: chlamydia urethritis 3 months ago treated at outside clinic with 1 gram azithromycin x 1. 5 male and 2 female sex partners in the last 3 months, reports condomless anal (receptive and insertive), vaginal, and oral sex
- Exam: no active discharge, trace meatal irritation
- You order comprehensive STI/HIV testing including urine GC/CT, and prescribe empiric CTX 500 mg IM x 1 + doxycycline 100 mg BID x 7 days
- Urine G/C is negative, and symptoms continue...

# What workup would you send?

## Choose all that apply:

- A: RPR with reflex to FTA-ABS
- B: HSV-1/HSV-2 PCR (urine)
- C: Repeat gonorrhea/chlamydia NAAT
- D: Urinalysis (first catch specimen) with microscopy
- E: Trichomonas NAAT
- F: Mycoplasma genitalium NAAT
- G: Urine bacterial culture
- H: PET-CT of the entire body

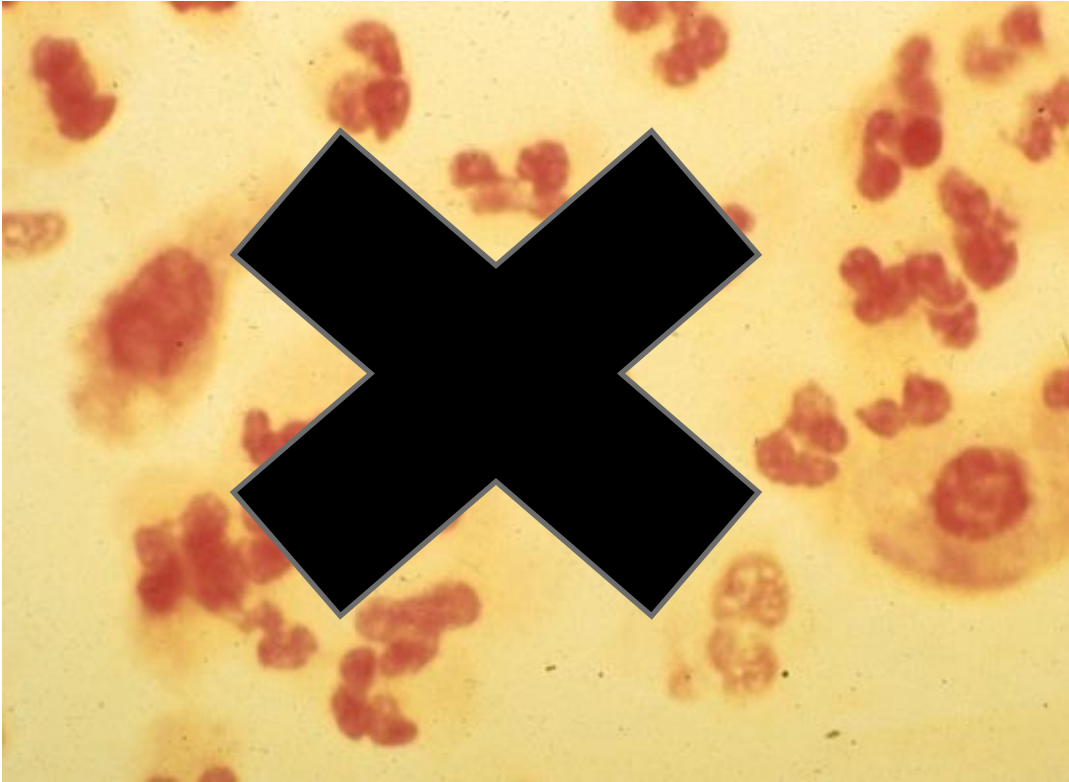


# Non-Gonococcal Urethritis

Non-chlamydial ^

## Etiology

- ~~C. trachomatis (15-40%)~~
- M. genitalium (15-25%)
- T. vaginalis (1-8%)
- HSV (3%)
- N. meningitidis
- Other bacteria (i.e. H. influenzae)
- Other viruses (i.e. adenovirus, EBV)
- UNKNOWN (~50%)!



# Non-GC, Non-CT Urethritis Workup

- Trichomonas urine NAAT (if MSW)
  - Mycoplasma genitalium NAAT
  - Consider HSV PCR
- } Less likely cause of isolated urethritis in females. Consider pelvic exam
- NOT M hominis, Ureaplasma spp
- Confirm urethritis!
    - Mucoid/purulent urethral discharge on exam
    - First-void urine : +LE or >10 WBC/HPF
- ↓
- If neither: consider e.g. chronic prostatitis/CPP, interstitial cystitis

# What Workup Would You Send?

- A: RPR with reflex to FTA-ABS
- B: HSV-1/HSV-2 PCR (urine)
- C: Repeat gonorrhea/chlamydia NAAT
- D: Urinalysis (first catch specimen) with microscopy
- E: Trichomonas NAAT
- F: Mycoplasma genitalium NAAT
- G: Urine bacterial culture
- H: PET-CT of the entire body



# Jeremy's Results

UA with micro: + LE, -nitrites, 20 WBCs

Trichomonas urine NAAT: neg

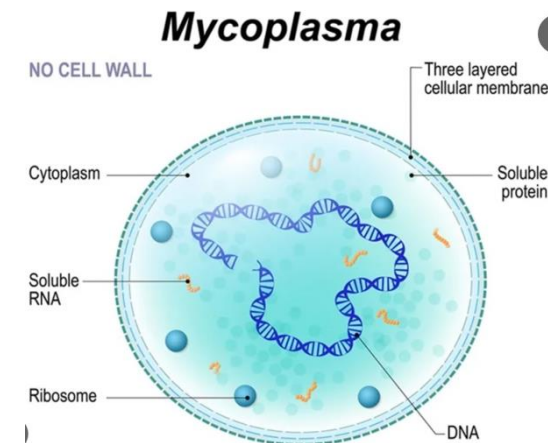
M. genitalium NAAT: positive

HSV urine PCR: neg



# Molli-Not-So-Cute

- Mollicutes: class of bacteria distinguished by lack of a peptidoglycan cell wall
  - medically significant genera include Mycoplasma and Ureaplasma
- Largely parasitic and intracellular, very small genome size—some of the smallest and simplest living things
- Difficult to culture, diagnosis depends on molecular testing
- Simplicity confers intrinsic resistance to many antibiotics
  - No peptidoglycan – no beta lactams, glycopeptides, or fosfomycin
  - No enzymes for folic acid metabolism – no TMP-SMX
  - Intracellular – poor activity of most aminoglycosides
  - Mutations in RNA polymerase – no rifampin



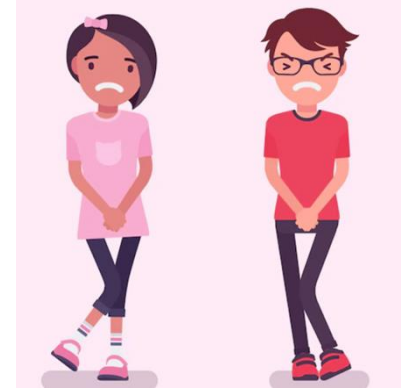
# M genitalium (Mgen) Epidemiology

- 2017-2018 NHANES: overall prevalence about 1.7% among people in the US aged 14-59
  - Other series with prevalence in the general population closer to 5%
  - In series from US STI clinics, prevalence 26% among women, 28.7% among men
  - Higher prevalence among people with HIV
  - Prevalence not increased among MSM
    - In this group, M genitalium detected in rectal >urine >> pharyngeal specimens
  - Concordance among couples is high, approximately 40-50% in heterosexual couples. One Australian study showed concordance among MSM of 27%

# Syndromes Associated with Mgen

## Male:

- Urethritis: 20-25% of non-gonococcal, non-chlamydial urethritis, and 40% of persistent or recurrent urethritis
- Proctitis: conflicting association in MSM between rectal Mgen detection and symptoms; weak or not present
- Pharyngitis: Mgen has not been demonstrated as a cause
- No clear association with chronic complications e.g. prostatitis



## Female:

- Cervicitis – Mgen detected in 10-30% of women with clinical cervicitis. Co-infection common, but also studies showing cytokine normalization after treatment
- PID – multiple studies demonstrate greater frequency of Mgen in women with PID, but generally cross-sectional. No clear prospective evidence of cause, or trial data showing that treatment of Mgen cervicitis prevents PID.
- Pregnancy/fertility-related complications– separate discussion!

# Back to Jeremy. What's Our Treatment?

A: Minocycline 100 mg BID x 14 days

B: Doxycycline 100 mg BID x 7 days, followed by azithromycin 1 gram x 1, then 500 mg daily x 3 days

C: Azithromycin 1 gram x 1

D: Doxycycline 100 mg BID x 7 days followed by moxifloxacin 400 mg daily x 7 days

E: Moxifloxacin 400 mg daily x 14 days

F: Bloodletting to rebalance his humours



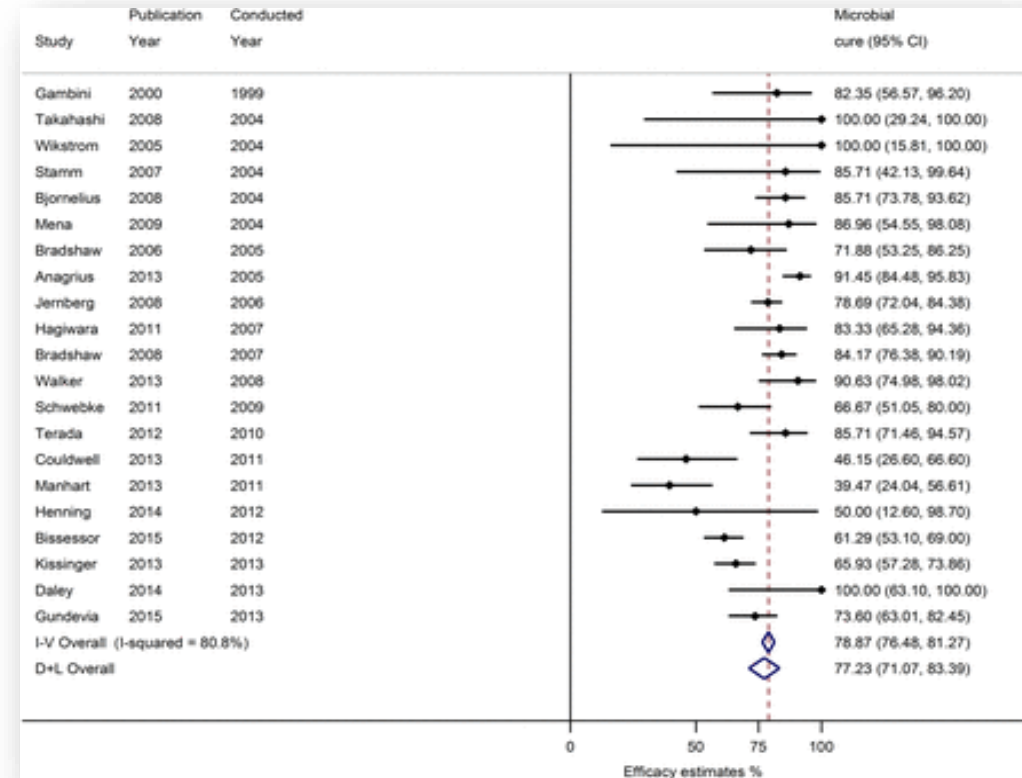
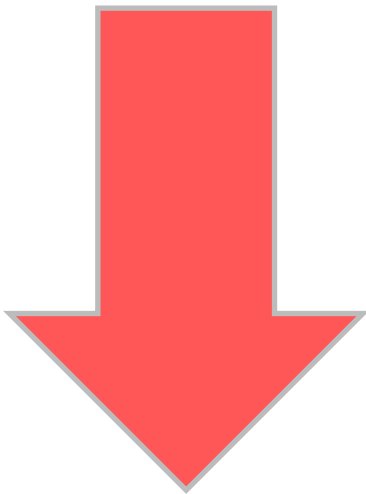
# Treatment – Why So Complicated?

- Intrinsic resistance leading to reliance on ribosomal agents with good intracellular activity
  - primary classes with activity include **macrolides, tetracyclines, and quinolones**
- Azithromycin 1 gram x 1 was previously the standard for syndromic treatment of non-gonococcal urethritis, and had good efficacy against *M genitalium*, but...

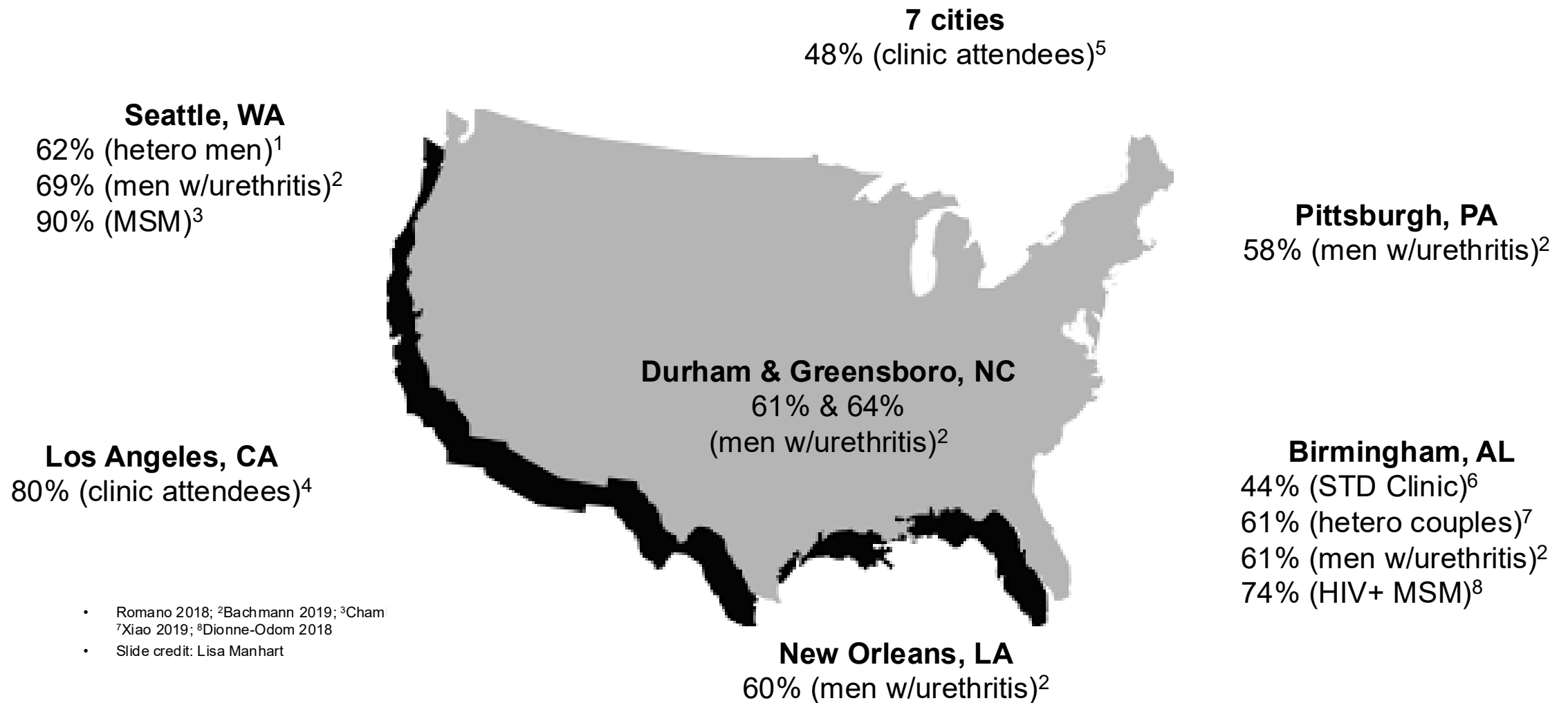
# Macrolide Resistance in *M. genitalium*

Driven by 5 SNP mutations in the 23S rRNA gene

- Pooled microbial cure rate 77.2%
  - Prior to 2009 – 85.3%
  - Since 2009 – 67%

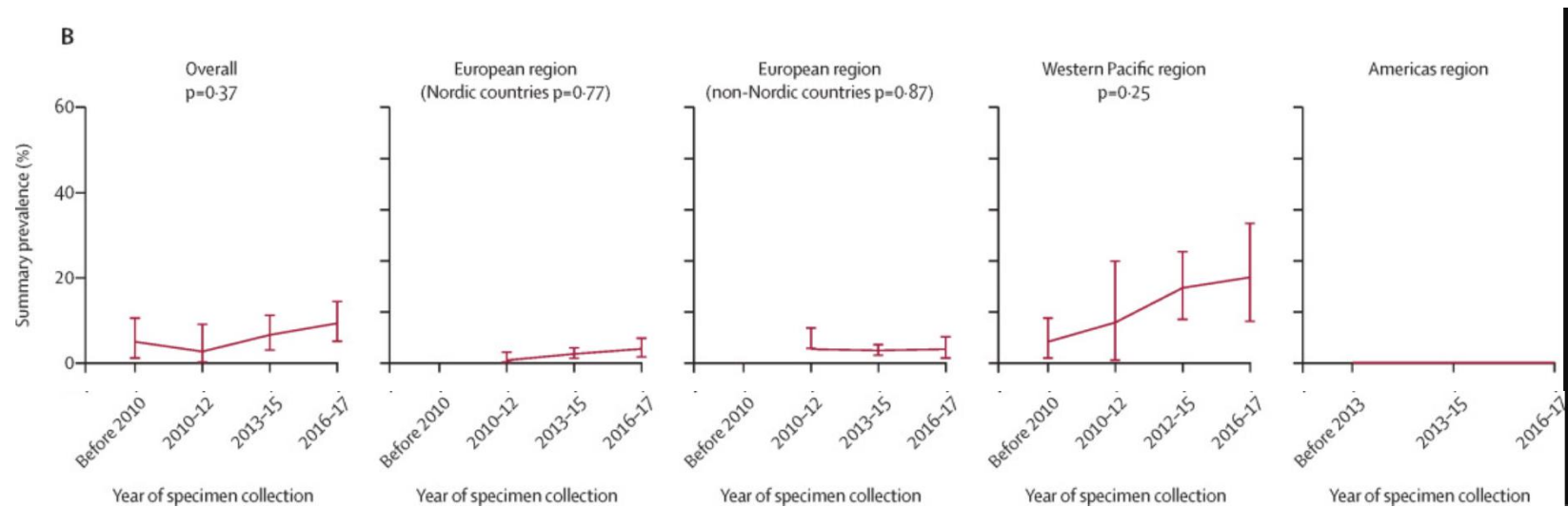


# Macrolide Resistance in M. genitalium



- Romano 2018; <sup>2</sup>Bachmann 2019; <sup>3</sup>Cham <sup>7</sup>Xiao 2019; <sup>8</sup>Dionne-Odom 2018
- Slide credit: Lisa Manhart

# Quinolone Resistance in *M genitalium*



- Meta-analysis included 25 studies reporting SNPs associated with quinolone resistance
- Global rate 2016-2017: 9.3%
- Americas region rate: 10.1% (insufficient data for temporal trend)

# Doxycycline Treatment failures in M. Gen

Gossé et al. *BMC Infectious Diseases* (2021) 21:1225  
<https://doi.org/10.1186/s12879-021-06910-1> BMC Infectious Diseases

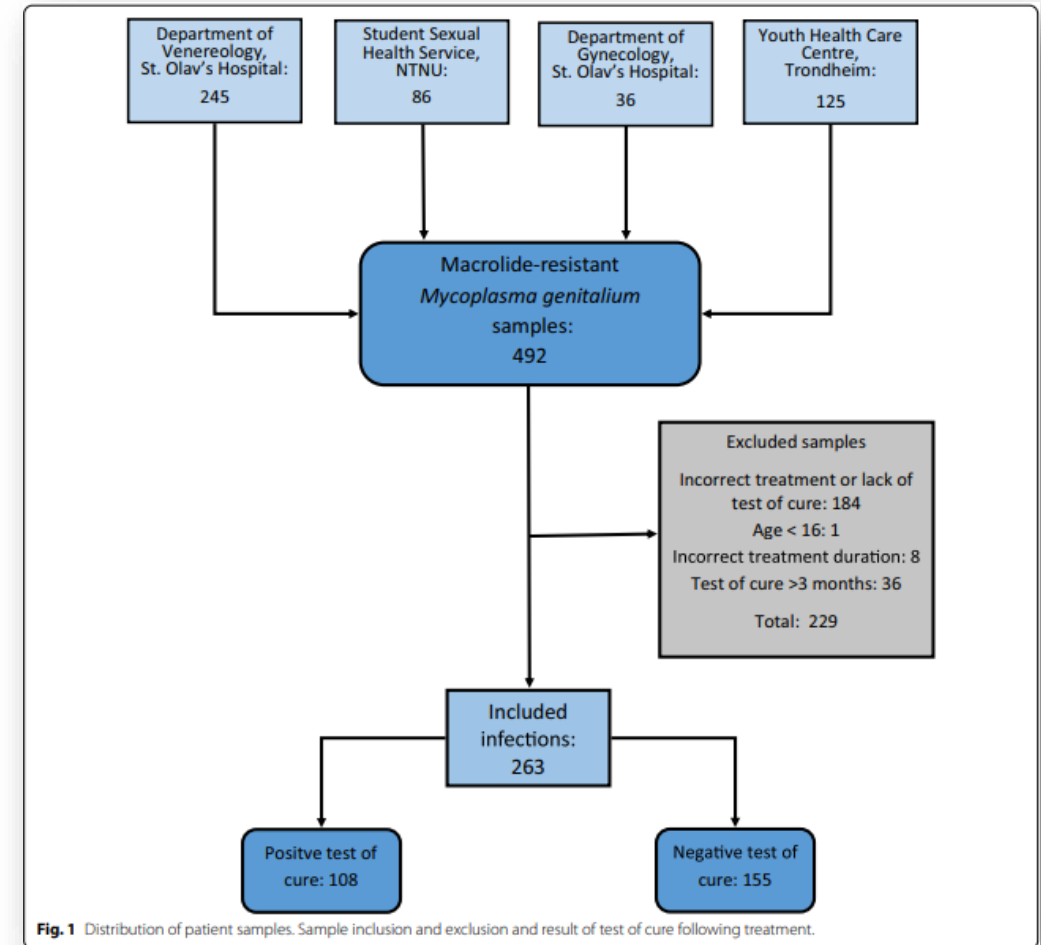
RESEARCH ARTICLE Open Access

Evaluation of treatment with two weeks of doxycycline on macrolide-resistant strains of *Mycoplasma genitalium*: a retrospective observational study

M. Gossé<sup>1\*</sup>, S. A. Nordbø<sup>1,2</sup> and B. Pukstad<sup>1,3</sup>

Check for updates

- 59% had a negative test of cure following treatment
- An additional 35% of symptomatic patients without negative test of cure experienced resolution of symptoms



# But Doxy Has Its Advantages

JOURNAL ARTICLE

## Identification of 16S rRNA mutations in *Mycoplasma genitalium* potentially associated with tetracycline resistance *in vivo* but not selected *in vitro* in *M. genitalium* and *Chlamydia trachomatis* <sup>FREE</sup>

Chloé Le Roy, Arabella Touati, Carla Balcon, Justine Garraud, Jean-Michel Molina, Béatrice Berçot, Bertille de Barbeyrac, Sabine Pereyre, Olivia Peuchant, Cécile Bébéar ✉

[Author Notes](#)

*Journal of Antimicrobial Chemotherapy*, Volume 76, Issue 5, May 2021, Pages 1150–1154,  
<https://doi.org/10.1093/jac/dkab016>

**Published:** 04 February 2021 **Article history** ▼

- 106 specimens of *M. genitalium* collected at the French National Reference Centre for Bacterial STIs from 2017-2019
- Samples passaged for 30 generations in subinhibitory concentrations of doxycycline or tetracycline
- No isolates developed elevated MICs to doxycycline at the end of the experiment
- 6 specimens had 16S rRNA mutations associated with doxy resistance in other organisms, but were still *in vitro* susceptible

# Resistance-Guided Sequential Therapy

Clinical Infectious Diseases

MAJOR ARTICLE



## Outcomes of Resistance-guided Sequential Treatment of *Mycoplasma genitalium* Infections: A Prospective Evaluation

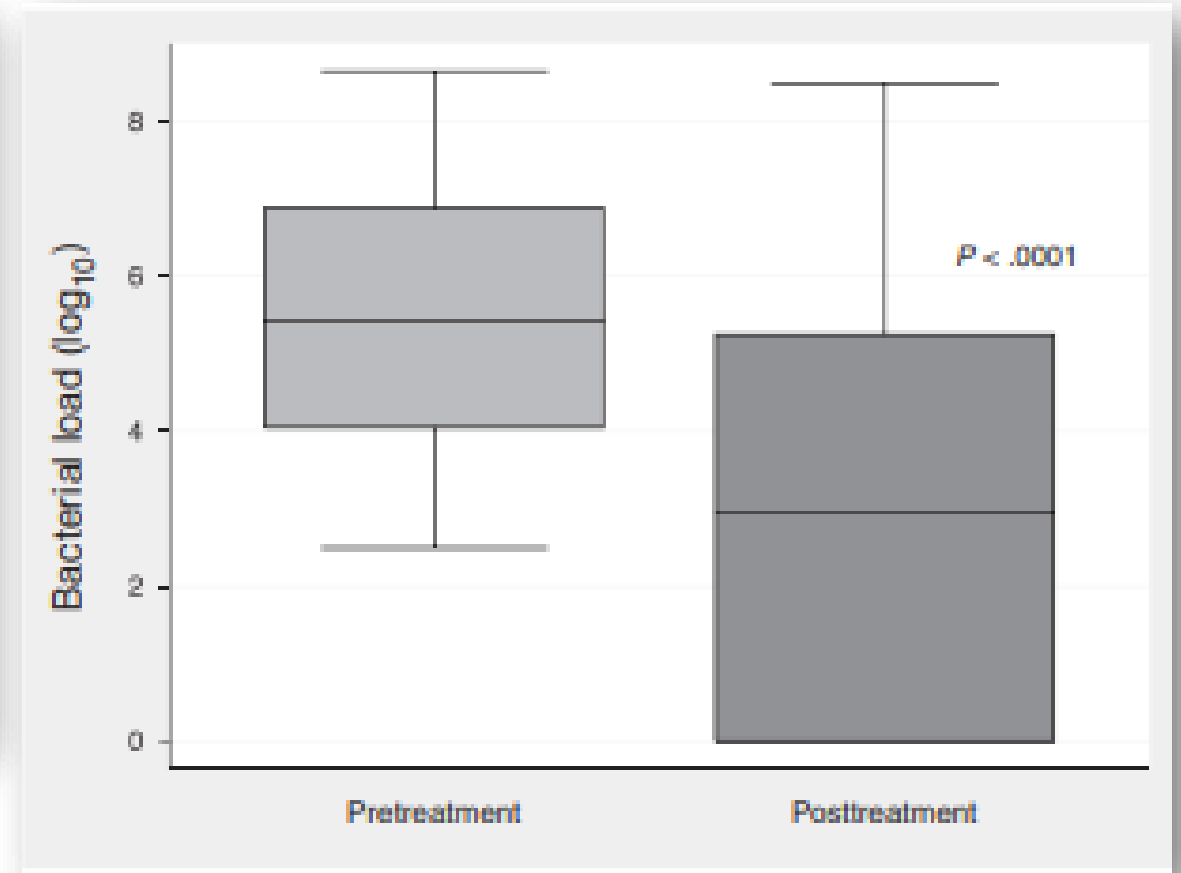
Tim R. H. Read,<sup>1,2</sup> Christopher K. Fairley,<sup>1,2</sup> Gerald L. Murray,<sup>3,4,5,6</sup> Jorgen S. Jensen,<sup>7</sup> Jennifer Danielewski,<sup>3,4</sup> Karen Worthington,<sup>2</sup> Michelle Doyle,<sup>2</sup> Elisa Mokany,<sup>8</sup> Litty Tan,<sup>8</sup> Eric P. F. Chow,<sup>1,2</sup> Suzanne M. Garland,<sup>3,4,5,9</sup> and Catriona S. Bradshaw<sup>1,2</sup>

<sup>1</sup>Central Clinical School, Faculty of Medicine, Nursing and Health Sciences, Monash University, Melbourne, <sup>2</sup>Melbourne Sexual Health Centre, Alfred Health, Carlton, <sup>3</sup>Murdoch Children's Research Institute, Parkville, <sup>4</sup>Department of Microbiology and Infectious Diseases, Royal Women's Hospital, Melbourne, <sup>5</sup>Infection and Immunity Program, Monash Biomedicine Discovery Institute, and <sup>6</sup>Royal Children's Hospital, Melbourne, Victoria, Australia; <sup>7</sup>Statens Serum Institut, Copenhagen, Denmark; <sup>8</sup>SpeeDx Pty Ltd, Eveleigh, New South Wales, and <sup>9</sup>Department of Obstetrics and Gynaecology, University of Melbourne, Victoria, Australia

(See the Major Article by Braun et al on pages 569-76 and Editorial commentary by Sulkowski on pages 577-9.)

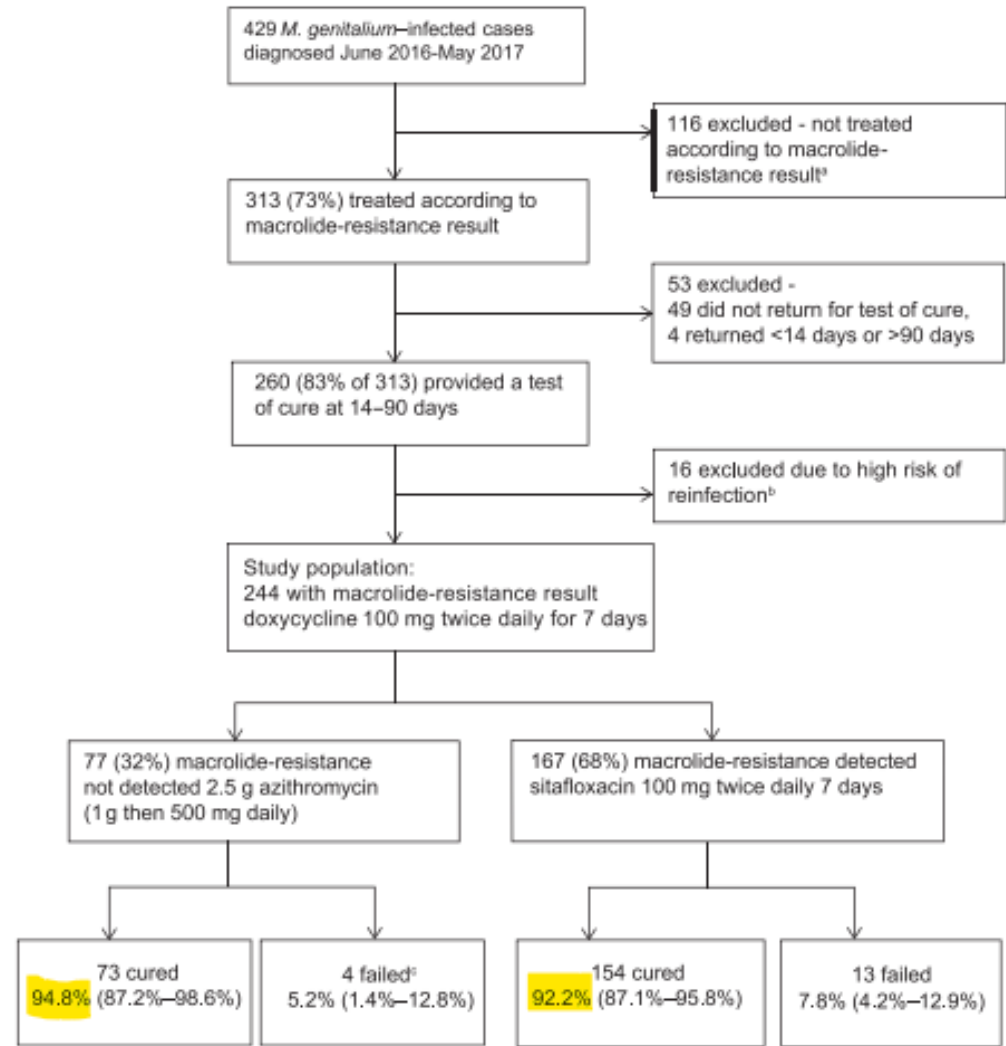
**Background.** Rising macrolide and quinolone resistance in *Mycoplasma genitalium* necessitate new treatment approaches. We evaluated outcomes of sequential antimicrobial therapy for *M. genitalium* guided by a macrolide-resistance assay.

**Methods.** In mid-2016, Melbourne Sexual Health Centre switched from azithromycin to doxycycline (100 mg twice daily for 7 days) for nongonococcal urethritis, cervicitis, and proctitis. Cases were tested for *M. genitalium* and macrolide-resistance mutations (MRMs) by polymerase chain reaction. Directly after doxycycline, MRM-negative infections received 2.5 g azithromycin (1 g, then 500 mg daily for 3 days), and MRM-positive infections received sitafloxacin (100 mg twice daily for 7 days). Assessment of test of cure and reinfection risk occurred 14–90 days after the second antibiotic.



# Resistance Guided, Sequential Therapy (Cont.)

- Success rates of 92-95% with this strategy
- Similar rates (92%) demonstrated by the same group when moxifloxacin was used in place of sitafloxacin
- Low (~5%) treatment-associated macrolide resistance



# Sounds Good, But...



- Molecular tests with detection of macrolide resistance not yet cleared by FDA
- Testing from commercial labs (e.g. LabCorp and ARUP) may be available in your area
- Another option is sending specimens out to University of Alabama at Birmingham
  - Now with quinolone RAM testing!

Test Menu	Acceptable Specimens	Transport/Processing Details	Turnaround Time	CPT Code
<b>PCR- Mycoplasma genitalium</b> Includes detection of macrolide resistance	<ul style="list-style-type: none"> <li>• Cervical swab</li> <li>• Throat</li> <li>• Urine</li> <li>• Urogenital swab</li> <li>• Vaginal swab</li> </ul>	<b>Collection Device:</b> Sterile container (can also be used as transport if received with 48 hours of collection and kept 2-8C)  <b>Transport Media:</b> Mycoplasma Ureaplasma transport media (examples: M4, M5, UTM, UVTM, eSwab)	1-4 days	87581

# Jeremy at Last

A: Minocycline 100 mg BID x 14 days

B: Doxycycline 100 mg BID x 7 days, followed by azithromycin 1 gram x 1, then 500 mg daily x 3 days

C: Azithromycin 1 gram x 1

D: Doxycycline 100 mg BID x 7 days followed by moxifloxacin 400 mg daily x 7 days

- Default treatment for most US M gen cases at this time -

E: Moxifloxacin 400 mg daily x 14 days



# Treatment Failure

- **Minocycline**

- Slightly lower MICs than doxycycline observed
- Largest case series of 90 patients with macrolide resistant M gen – 66.7% cure rate [1]
  - 100 mg BID x 14 days
  - 62 had failed tx with moxifloxacin



- **Pristinamycin**

- 85/114 (75%) of patients with macrolide resistant M gen cured with 10 days treatment
- **Not available in the USA**



Centers for Disease  
Control and Prevention



## **Mycoplasma genitalium Treatment Failure Registry**

**The purpose of this form is to collect clinical information on cases of *Mycoplasma genitalium* that fail antimicrobial therapy. All reported information will be maintained in the strictest confidence.**

# Wasn't There Something About Pregnancy?

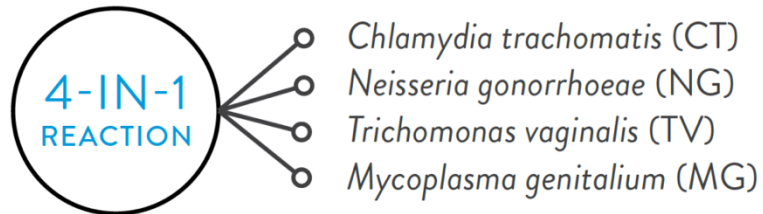
- 2022 meta-analysis assessed risk of adverse pregnancy and peri-natal outcomes
  - Pre-term birth: strongest evidence, OR of approximately 2
    - Oddly greater than OR for this outcome with gonorrhea/chlamydia/trichomonas
    - Unable to assess confounding due to lack of adjustment for variables other than age
  - Spontaneous abortion: OR = 1
  - PROM, low birth weight, perinatal death: minimal data
- Authors conclude that there is insufficient evidence to recommend screening for Mgen in asymptomatic pregnant people

# Mgen in the Pregnant Patient

- Moxifloxacin and doxycycline not routinely recommended for use during pregnancy
- Given lack of clear evidence for harms, reasonable to defer therapy if no sx's
- For patients with symptoms, azithromycin is the only drug routinely recommended for use
  - In this group, reasonable to send resistance testing to UAB
    - If macrolide-susceptible, can treat with azithromycin 4-day course
    - If resistant, risk-benefit conversation with patient, then potential treatment after delivery

# And Yet...

## ONE ASSAY, MULTIPLE POSSIBILITIES REDEFINING THE FUTURE OF STI TESTING WITH OPERATIONAL EFFICIENCY



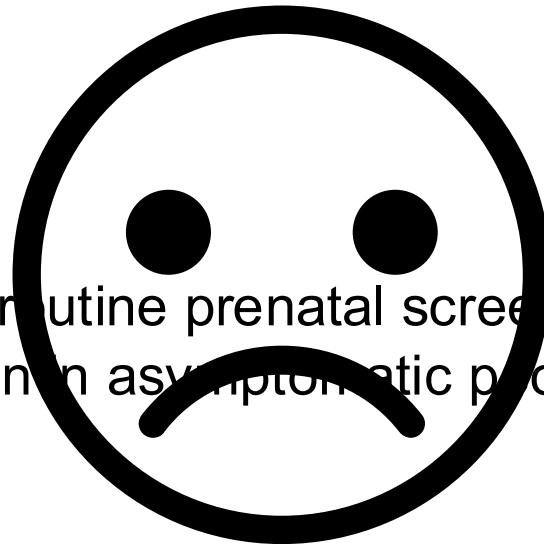
Alinity m STI assay is a 4-in-1 multiplex assay to detect and differentiate CT, TV, MG, and NG to aid in the diagnosis of infection from these organisms.

- Enter the multiplex PCR
- Used at some sites as part of routine prenatal screening rather than G/C alone
- Can result in detection of Mgen in asymptomatic people, especially pregnant people!

## ANALYTES

### Anyplex™ II STI-7e Detection

- *Chlamydia trachomatis* (CT)
- *Mycoplasma genitalium* (MG)
- *Mycoplasma hominis* (MH)
- *Neisseria gonorrhoeae* (NG)
- *Trichomonas vaginalis* (TV)
- *Ureaplasma parvum* (UP)
- *Ureaplasma urealyticum* (UU)
- Exogenous Internal Control



# Take Home

- Mycoplasma genitalium is a relatively common inhabitant of the male and female GU tract
- Clinical syndromes include urethritis in males, possibly proctitis. In females: cervicitis and PID—data are not adequate to demonstrate Mgen causality for preterm birth, spontaneous abortion, perinatal death
- Mgen cell makeup confers intrinsic resistance to many antibiotics
  - Macrolide resistance is prevalent, and fluoroquinolone resistance rising

# Take Home (Cont.)

- Treatment relies on sequential therapy with doxycycline followed by moxifloxacin (unless resistance testing available)
- Options for treatment failure in the US include minocycline, potentially checking for macrolide susceptibility
- Options for treatment of pregnant people are extremely limited—no indication for screening in the absence of symptoms

# Questions?

# Eyes for a Sore Site

- Jasmine is a 29-year-old woman who presents to her PCP for an urgent visit
- Complains of painful genital lesions x 5 days. Mild fever at symptom onset - resolved. Also dysuria, which is improving. No vaginal discharge, pelvic pain.
- No history of anything similar, no new cleansing products/detergents. Has 1 new male sexual partner x 6 months, reports oral sex (gives and receives), and vaginal sex. Stopped using condoms 4 months ago after negative STI screening. Has not seen genital lesions on her partner.
- Exam: many small erythematous erosions of the labia majora and minora as well as surrounding skin, a few have started to crust. Bilateral tender inguinal lymphadenopathy.

# Physical Exam



# What Workup Would You Send for Jasmine?

**Choose all that apply**

- A: RPR with reflex to FTA-ABS
- B: HIV ab/ag
- C: HSV-1/HSV-2 IgG/IgM
- D: HSV-2 lesion PCR
- E: HSV-1/HSV-2 lesion PCR
- F: Urine gonorrhea/chlamydia NAAT
- G: Multiplex PCR panel with ALL STIs



# Genital Ulcer Disease Ddx

Syndrome	Differential	Distinguishing features
<b>Genital ulcer</b>		
	HSV	History of prior outbreaks, systemic symptoms are rare in non-primary infection.
	Primary syphilis	Ulcer is typically painless. Rectal ulcers , however, may be painful.
	Mpox	Systemic symptoms present at some time during course, papular-stage lesions often umbilicated
	LGV	Ulcer typically painless, often resolved at time of presentation. Inguinal LAD pronounced in men. Proctocolitis for rectal infection.
	Chancroid/Granuloma Inguinale	Rare in the USA.
<b>Rash (localized or general)</b>		
	VZV	Dermatomal distribution (shingles), isolated anogenital involvement less common
	Molluscum contagiosum	Lesions typically painless, systemic symptoms and mucosal involvement less common



# What Workup Would You Send for Jasmine?

**Choose all that apply**

- A: RPR with reflex to FTA-ABS
- B: HIV ab/ag
- C: HSV-1/HSV-2 IgG/IgM
- D: HSV-2 only lesion PCR
- E: HSV-1/HSV-2 lesion PCR
- F: Urine gonorrhea/chlamydia NAAT
- G: Multiplex PCR panel with ALL STIs



# Jasmine's Results

- RPR: non-reactive
- HIV ab/ag: non-reactive
- **HSV-1/HSV-2 lesion PCR: HSV-1 +**
- Urine gonorrhea/chlamydia NAAT: neg

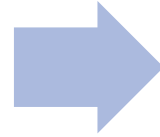


# HSV Virology/Epidemiology

- Transmission via direct contact with lesions or saliva
- Primary infection with higher incidence of systemic sx's, higher severity
- Latency in sensory nerves, followed by episodes of reactivation
  - HSV-1 reactivates more efficiently from trigeminal ganglia (orolabial herpes)
  - HSV-2 reactivates more efficiently from sacral dorsal root ganglia (anogenital herpes)
  - EITHER HSV can cause genital herpes. Rates of HSV-1 genital herpes are rising
  - Recurrence and asymptomatic shedding more common with HSV-2
- From 2015-2016, 11.9% of US adults aged 14-49 were HSV-2+, and 47.8% HSV-1+
- Rate of clinical genital herpes 2019-2021: 236-280 per 100,000 person/years

# Diagnosis: Virologic Tests (When Lesions are Present)

Detection of HSV from genital ulcers or mucocutaneous lesions (PCR or viral culture)



HSV PCR is the preferred diagnostic test

- FDA cleared PCR based HSV tests
  - Sensitive and specific
  - Can distinguish HSV-1 from HSV-2
- Viral culture
  - Low sensitivity (especially for recurrent lesions and in healing lesions)
  - Only way to detect acyclovir resistant HSV

# Guideline Updates – Serologic HSV Testing

- Useful
  - Recurrent or atypical genital symptoms or lesions with a negative HSV PCR or culture result
  - Clinical diagnosis of genital herpes without laboratory confirmation
  - 12 weeks after suspected recent acquisition
  - Patient's partner has genital herpes
- Might be useful
  - Persons at higher risk for infection (presenting for STI evaluation—10 or more lifetime sex partners)
  - Persons with HIV
- Not useful
  - Screening of the general population

## Two-Step Serologic Testing

Step 1: EIA Assay (IgG)\*  
(often falsely positive at  
low index value (<3.0))

### Positive EIA

Step 2: Confirm with a  
second test that uses a  
different antigen  
(Biokit/Western blot)

\*IgM is not recommended for serologic testing

# What Treatment Would You Offer Jasmine?

- A: Valacyclovir 1 g PO twice daily x 7-10 days
- B: Valacyclovir 1 gram PO daily x 5 days
- C: Valacyclovir 2 grams PO twice daily x 1 day
- D: Acyclovir 10 mg/kg IV q 8 hours
- E: Valacyclovir 1 gram daily indefinitely
- F: Transplant of the spinal dorsal nerve root ganglia



# Treatment Options

**All patients with first episodes of genital herpes should receive antiviral therapy**

1. Acyclovir 400 mg orally 3 times/day for 7–10 days
  2. Famciclovir 250 mg orally 3 times/day for 7–10 days
  3. Valacyclovir 1 gm orally 2 times/day for 7–10 days
- Treatment can be extended if healing is incomplete after 10 days of therapy.

# Treating/Preventing Recurrences

- **Episodic/Intermittent therapy** - ameliorate or shorten the duration of lesions
  - Recurrences are less frequent after the first episode of HSV-1 genital herpes, and genital shedding rapidly decreases during the first year of infection
- **Suppressive therapy** - reduce the frequency of recurrences
  - Almost all persons with symptomatic first-episode HSV-2 genital herpes subsequently experience recurrent episodes of genital lesions
  - Suppressive therapy can decrease recurrence rate by 70-80% in those with frequent episodes
  - May confer benefits for preventing transmission (more later)

# Antiviral Options

## Suppressive

### Recommended Regimens

Acyclovir 400 mg orally 2 times/day  
OR  
Valacyclovir 500 mg orally once a day\*  
OR  
**Valacyclovir 1 gm orally once a day**  
OR  
Famciclovir 250 mg orally 2 times/day

\* Valacyclovir 500 mg once a day might be less effective than other valacyclovir or acyclovir dosing regimens for persons who have frequent recurrences (i.e.,  $\geq 10$  episodes/year).

Dose and/or duration are increased for immunosuppressed people: e.g. valacyclovir 1 gram BID x 7-10 days (intermittent), valacyclovir 500 mg BID for suppression

## Intermittent

### Recommended Regimens for Episodic Therapy for Recurrent HSV-2 Genital Herpes\*

Acyclovir 800 mg orally 2 times/day for 5 days  
OR  
Acyclovir 800 mg orally 3 times/day for 2 days  
OR  
Famciclovir 1 gm orally 2 times/day for 1 day  
OR  
Famciclovir 500 mg once, followed by 250 mg 2 times/day for 2 days  
OR  
Famciclovir 125 mg 2 times/day for 5 days  
OR  
Valacyclovir 500 mg orally 2 times/day for 3 days  
OR  
**Valacyclovir 1 gm orally once daily for 5 days**

\*Acyclovir 400 mg orally 3 times/day is also effective, but are not recommended because of frequency of dosing.

# What Treatment Would You Offer Jasmine?

- A: Valacyclovir 1 g PO twice daily x 7-10 days
- B: Valacyclovir 1 gram PO daily x 5 days
- C: Valacyclovir 2 grams PO twice daily x 1 day
- D: Acyclovir 10 mg/kg IV q 8 hours
- E: Valacyclovir 1 gram daily indefinitely
- F: Transplant of the spinal dorsal nerve root ganglia



# Preventing Transmission

- Daily valacyclovir lowers risk of HSV-2 transmission from HIV-negative people with symptomatic genital herpes (approx. 50%)
  - Unknown if true for those without a history of symptoms. Not effective/recommended for people with HIV not on ART
- Condoms decrease, but don't eliminate, risk for HSV-2 transmission
- Male medical circumcision
- Caution against HSV acquisition during pregnancy – avoid genital and/or oral sex with partners who have history of orolabial or genital herpes in 3<sup>rd</sup> trimester, monitor closely peri-delivery
- Pregnant patients with a history of genital herpes should be offered suppression starting at 36 weeks to decrease risk of recurrence during delivery, c-section rate, and asymptomatic shedding

# HSV Patient Counseling

- People with a history of genital herpes are recommended to disclose to prospective sex partners
- Transmission more likely with active lesions, but can occur during asymptomatic periods of viral shedding
- Serology cannot determine whether someone is infected orally, genitally, or both
  - HSV-1 is an increasing cause of genital herpes among young women and MSM
- HSV-2 transmission reduced, but not eliminated, by male condom use and valacyclovir suppression (among people without HIV only)
- Suppression not proven to reduce risk of transmission of HSV-1
- People with HSV-2 are at increased risk of acquiring HIV when exposed via sexual contact

# HSV Takeaway Points

- When possible, HSV diagnosis should be confirmed with PCR testing from an active lesion
- Serology may help support the diagnosis w/o active lesions, but is not conclusive
- All patients with a first episode of HSV should get antiviral treatment
- Subsequent outbreaks can be treated with episodic or suppressive therapy
  - Suppressive typically used for those with frequent outbreaks

# HSV Takeaway Points

- HSV 1 is an increasingly common cause of genital herpes, especially among younger people, but typically causes fewer outbreaks and less viral shedding
- HSV 2 causes more frequent outbreaks and increases the risk of HIV transmission.
- Suppressive therapy can reduce outbreak frequency in all patients, and reduce the chances of HSV-2 transmission among people without HIV

# Resources

- American Society for Sexual Health patient education materials
  - Patient handouts in English and Spanish for most STIs, including Mgen and HSV
  - <https://www.ashapublications.org/patient-education-materials>
- CDC STI 2021 guidelines include patient counseling points for genital herpes
  - <https://www.cdc.gov/std/treatment-guidelines/herpes.htm>

# Questions?