Course Objectives

1. Become familiar with current epidemiologic trends of common STIs and screening recommendations from the US Centers for Disease Control and Prevention (CDC)

2. Review common presentations, available diagnostic modalities, and recommended treatment regimens for common STIs

Overview

- Introduction
- STIs to be reviewed today:
  - *Chlamydia trachomatis* (CT)
  - *Neisseria gonorrhoeae* (GC)
  - *Trichomonas vaginalis* (TV)
  - Syphilis (*Treponema pallidum*; TP)
  - Herpes Simplex Virus (HSV)
  - Human Papillomavirus (HPV)
- STIs not covered: HIV, Hepatitis A, B, and C, mycoplasmas, ureaplasmas, chancroid, and bacterial vaginosis
- Speakers report no conflicts of interest or financial disclosures
Introduction

• STIs are “hidden epidemics” of enormous health and economic consequence in the United States

• STIs can be disruptive, painful, and lead to long-term negative health consequences; they also increase the risk of other detrimental reproductive morbidities

• All communities are impacted by STIs and all individuals directly or indirectly pay for the costs of these diseases


Chlamydia trachomatis (CT)

• Small gram-negative rod and obligate intracellular organism

• Most common reportable bacterial STI in the US
  – U.S. → 1,307,893 cases (Rate: 426.0 / 100,000)
  – Ohio → 51,150 cases (Rate: 443.1 / 100,000)
  – Columbus → 9,545 cases (Rate: 529.7 / 100,000)

• As of January 2000, regulations in all 50 states and D.C. require reporting all CT cases to CDC


Case 1 – “Urethritis”

• 27 y/o African-American male presents with complaints of mild dysuria and clear penile discharge x 3 days. Denies constitutional symptoms.

• Patient reports multiple sex partners and unprotected vaginal and rectal intercourse with women only.

• Gram stain of discharge shows > 5 WBC (PMNs) / HPF. No organisms seen.


CT Rate by Gender, US, 1990-2010

Women >> Men (ages 15-24 years)

**Neisseria gonorrhoeae (GC)**

- **Gram-negative diplococci**
- **Second most common bacterial STI in US:**
  - U.S. → 309,341 cases (Rate: 100.8 cases / 100,000)
  - Ohio → 16,496 cases (Rate: 142.9 cases / 100,000)
  - Columbus → 3,351 cases (Rate: 186.0 cases / 100,000)
- **As of January 2000, regulations in all 50 states and D.C. require reporting all gonorrhea cases to CDC.**

**GC Rate by Race/Ethnicity, US, 2001-10**

![Graph showing GC rate by race/ethnicity from 2001 to 2010.](http://www.cdc.gov/std/stats10/figures/22.htm)


**Risk factors: CT and GC**

- Young age (adolescent into early adulthood)
- New or multiple partners, or partner with other partners in last 3 months
- Infrequent/non-use of barrier contraception
- Cervical ectopy
- Single (unmarried)
- Prior history of STIs (especially CT/GC)
- Low or intermediate education
- Low socioeconomic status
- Ethnic minority (especially African-American)
- Illicit drug use
- Commercial sex work


**GC Prevalence by Sex and Reporting Source, US, 2010**

![Bar chart showing GC prevalence by sex and reporting source.](http://www.cdc.gov/std/stats10/figures/24.htm)

*HMO = health maintenance organization; HD = health department

These categories represent 69.5% of cases with known reporting source. Of all cases, 13.2% had missing or unknown reporting source.


**Screening Recommendations: CT**

- All sexually active women ≤ 25 y/o, annually
- All sexually active women ≥ 26 y/o with risk factors (i.e. new or multiple sex partners), annually
- All pregnant women at 1st prenatal visit
- Sexually active young men? Not enough evidence to suggest routine screening, except in high risk settings (i.e. adolescent clinics, prison/jail, STD clinics)
- Primary focus of CT screening is to prevent complications in women (i.e. pelvic inflammatory disease)

*Source: 2010 CDC STD Treatment Guidelines*
### Screening Recommendations: GC

- Target all sexually active women at risk for infection (i.e. previous STI, new or multiple sex partners, inconsistent condom use, commercial sex work, drug use, or living in high prevalence areas), annually
- All pregnant women at risk for infection (see above) at 1st pre-natal visit
- Sexually active young men? → Not enough evidence for routine screening except in high risk settings (i.e. adolescent clinics, jail/prison, STD clinics)
- Similar to CT, primary focus of GC screening and treatment effort is to prevent complications in women

Source: 2010 CDC STD Treatment Guidelines

### Clinical Manifestations: CT and GC

- Non-gonococcal urethritis (NGU)
- Epididymitis / Orchitis / Prostatitis
- Proctitis
- Mucopurulent cervicitis (MPC)
- Pelvic inflammatory disease (PID)
- Lymphogranuloma venereum (LGV; serovars L1, L2, and L3)
- Reactive arthritis (Reiter’s Syndrome)
- Infant pneumonia
- Ophthalmia neonatorum
- Asymptomatic infection >50%!!!
- Gonococcal urethritis
- Epididymitis / Orchitis / Prostatitis
- Proctitis
- MPC / PID
- Pharyngitis
- Perihepatitis (Fitzhugh-Curtis Syndrome)
- Disseminated Gonococcal Infection (DGI)
- Meningitis / Endocarditis / Septic arthritis
- Ophthalmia neonatorum
- Asymptomatic infection


### Screening in Men who have Sex with Men (MSM)

- In MSM who report in last 12 months:
  - Insertive anal intercourse → Urethral CT & GC, annually
  - Receptive anal intercourse → Rectal CT & GC, annually
  - Receptive oral intercourse → Oropharyngeal GC only, annually

- More frequent testing (every 3-6 months) if:
  - Multiple or anonymous sex partners
  - Sex with methamphetamine use
  - Sex partners report any of the above

Source: 2010 CDC STD Treatment Guidelines

### Diagnosis: CT

- Nucleic Acid Amplification (NAAT)
  - High sensitivity (80 – 100%) and specificity (≥94%)
  - Yield by site: Endocervical = Urine / Vaginal
  - Liquid-based cervical cytology and Pap-smear samples have lower sensitivity
  - Endocervical, vaginal, urethral and urine are FDA approved
  - Oropharyngeal and rectal specimens NOT FDA approved
  - Tissue culture, Direct Fluorescent Antibody (DFA), and Enzyme Immunoassay (EIA)

Diagnosis: GC (1)

- **Gram stain** → PMNs with intracellular gram-negative diplococci:
  
  - In symptomatic men
    - Sensitivity >95% and specificity >99%
  
  - In asymptomatic men
    - Low sensitivity; does not rule out infection
    - Not recommended for endocervical, pharyngeal, and/or rectal specimens


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CT Treatment

Uncomplicated urogenital infections

<table>
<thead>
<tr>
<th>Non-Pregnant Patients</th>
<th>Pregnant Patients</th>
</tr>
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</table>

**Recommended Regimens**

- **Ampicillin 1 gram orally in a single dose**
- **Ceftriaxone 1 g intramuscularly in a single dose**

**Alternative Regimens**

- **Erythromycin base 500 mg orally four times a day for 7 days**
  - OR
- **Erythromycin ethylsuccinate 900 mg orally four times a day for 7 days**

**Oropharyngeal, rectal, and/or conjunctival specimens are NOT FDA approved**

Source: 2010 CDC STD Treatment Guidelines

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GC Treatment

Uncomplicated Gonococcal Infections of the Cervix, Urethra, and Rectum

**Recommended Regimens**

- **Ceftriaxone 250 mg IM in a single dose**
- **Cefixime 400 mg orally in a single dose**
- **Single-dose injectible cephalosporin regimens**
- **Azithromycin 1 g orally in a single dose**
- **Doxycycline 100 mg orally twice a day for 7 days**

Uncomplicated Gonococcal Infections of the Pharynx

**Recommended Regimens**

- **Ceftriaxone 250 mg IM in a single dose**
- **Azithromycin 1 g orally in a single dose**
- **Doxycycline 100 mg orally twice a day for 7 days**

- GC/CT co-infection rates ~46%
  - If PCN or cephalosporin allergy
  - Azithromycin (AZM) 2g PO x 1 (must limit use!!)

- Fluoroquinolones (ofloxacin, levofloxacin) no longer recommended

CT and/or GC Follow-up (1)

- No sex for 7 days after single dose or until completion of 7-day treatment regimens
- Notify and treat all sex partners from last 60 days and/or most recent partner even if >60 days
- Test of Cure (TOC) at 3 – 4 weeks is NOT recommended in men or non-pregnant women, unless...
  - Compliance with initial treatment is questionable (non-issue with DOT)
  - Symptoms persist
  - Re-infection suspected (common)
- Risk of false (-) and false (+) results with repeat NAAT < 3 weeks after treatment

Source: 2010 CDC STD Treatment Guidelines

Trichomonas vaginalis (TV)

- Motile flagellated protozoan
- Most prevalent curable STI in US and worldwide
- Overall prevalence 3.1% (95% CI, 2.3-4.3%)
- African-American women; 14-49 year-olds (13.3%)
- Women attending STD clinics (13-34%)
- Men attending STD clinics (3-17%)
- Women co-infected with HIV (6.1-52.6%)
- In 2010 → 149,000 initial visits to physician offices due to trichomoniasis


CT and/or GC Follow-up (2)

- Re-testing (women and men) recommended at 3 months after treatment (or whenever medical care is sought within next 12 months) due to high risk of re-infection
- In pregnant women:
  - TOC for CT with NAAT, 3 – 4 weeks after treatment
  - If diagnosed with CT and/or GC during 1st trimester, re-test 3-6 months after treatment (preferably during 3rd trimester)
  - If negative for CT and/or GC during 1st trimester, but remains at high risk, re-test during 3rd trimester

Source: 2010 CDC STD Treatment Guidelines

TV Screening Recommendations

- Women complaining of vaginal discharge, especially if high risk (i.e. new or multiple sex partners, prior STI, commercial sex work, injection drug use)
- All sexually active HIV-positive women (at entry to care, then annually)
  - TV infection increases HIV shedding from vaginal secretions, while TV treatment decreases HIV shedding
- No optimal studies evaluating costs & benefits of screening asymptomatic STD clinic patients

TV Clinical Manifestations

- NGU
- Prostatitis
- Epididymitis
- Vaginitis (diffuse, yellow-green, and malodorous vaginal discharge)
- Cervicitis (“strawberry cervix”)
- Prolonged asymptomatic carriage (women)
- Transient asymptomatic carriage (men)


TV Diagnosis (1)

- Microscopy of vaginal secretions
  - “Wet prep”
  - Low cost, simple
  - Requires rapid examination after collection
  - Sensitivity → 60-70%
- Culture (Diamond’s media): current Gold Standard
  - Sensitivity with vaginal secretions → 75-95%
  - Sensitivity of male urethra samples (swab, urine, or semen) → 28-56%
- Liquid-based cervical cytology / PAP-smears
  - May require confirmatory testing
  - Overall poor sensitivity


TV Diagnosis (2)

- Point-of-Care Tests
  - Rapid Antigen Test & Nucleic-Acid Probe Hybridization
    - Sensitivity → >83%, specificity → >97%
- NAAT: Transcription-mediated amplification (TMA)
  - Sensitivity → 74-98%
  - Specificity → 87-98%
- NAAT: Polymerase chain reaction (PCR)
  - Sensitivity → 88-97%
  - Specificity → 98-99%


TV Treatment and Follow-up

- Notify, test, and treat all sex partners
- Re-screen at 3 months due to high re-infection rate (~17% positive again at 3 months)
- Treatment failure 1 month post-treatment is 7% in HIV (-) women, 10% in HIV (+) women
- Low-level metronidazole (MNZ) resistance: ~2 – 5%
  - High-level resistance is rare.
- No data to support re-screening men diagnosed with TV infection

TV Management: Adverse Reactions

- Allergy or adverse reactions to 5-nitroimidazoles
  - Severe reactions do not always reoccur, but we cannot predict who will react with re-challenge
  - Avoid 5-nitroimidazoles in patients with history of allergic reactions
  - Alternative treatments have poor response rates (<50%)
  - Can consider consult with allergist or immunologist for desensitization if history of immediate-type hypersensitivity reaction


Case 1 – Urethritis

- Presumptive diagnosis: NGU
- Urine CT NAAT – Positive
- Urine GC NAAT – Negative
- Urine TV culture – Negative
- Rapid HIV test – Negative
- Treatment: Azithromycin 1g PO x 1
- Notification and treatment of sex partners
- Re-testing in 3 months or whenever care is sought within next 12 months
- Counseling regarding safer sex practices

Sources: 2010 CDC STD Treatment Guidelines

TV Management: Treatment Failure

- Exclude re-infection
- If no response to MNZ 2g PO x 1, treat with MNZ 500mg PO BID x 7 days
- If no response, then MNZ or TNZ 2g PO daily x 5 days
- If no response, consult with ID specialist and CDC for susceptibility testing of TV isolate: 404-718-4141 or http://www.cdc.gov/std
- Treat male sex partners of women who experience treatment failure with MNZ 500mg PO BID x 7 days OR TNZ 2g PO x 1

Sources: 2010 CDC STD Treatment Guidelines

Case 2 – “Rash and Sores”

- 36 y/o white HIV + gay male presents with 3-week history of generalized malaise, low-grade fevers, oral ulcers, and whole body rash
- Has had a steady partner (also HIV+) for 4 months, but reports that they both engage in unprotected anal and oral intercourse with each other and with anonymous partners met online

Syphilis

- Etiologic agent is *Treponema pallidum*
- Microaerophilic or anaerobic gram-negative rod
- Cannot survive outside host, difficult to grow in vitro
- Very slow growth rate (doubling time 30-36 hours)
- Stages of infection:
  - Primary (PS)
  - Secondary (SS)
  - Early Latent (ELS; < 1 year)
  - Late Latent (> 1 year)
  - Tertiary
  - Neurosyphilis (can occur at any stage)
  - Syphilis of "Unknown Duration"

"Early Syphilis," (ES) (Increased risk of transmission to susceptible partners)


Syphilis Rates, US, 2010

- All Stages: 45,834 cases (Rate: 14.9 cases / 100,00)
- Primary & Secondary Syphilis (PS & SS):
  - U.S. → 13,774 cases (Rate: 4.5 cases / 100,000)
  - Ohio → 528 cases (Rate: 4.6 cases / 100,000)
  - Columbus → 120 cases (6.7 cases / 100,000)
- Early Latent Syphilis (ELS):
  - U.S. → 13,604 cases (Rate: 4.4 cases / 100,000)
  - Ohio → 193 cases (Rate: 1.7 cases / 100,000)*
  - Columbus → 37 cases (Rate 4.8 cases / 100,000)*

Primary and Secondary Syphilis Cases by Sex, Behavior and Race/Ethnicity, US, 2010

- Of the reported male cases of P&S syphilis, 18.3% were missing sex of sex partner; 2.0% of male cases with sex of sex partner data were missing race/ethnicity data.
- No imputation done for race/ethnicity
- MSW = men who have sex with women only; MSM = men who have sex with men

Primary and Secondary Syphilis Cases by Sex, Behavior, and Reporting Sources, US, 2010

* Of reported male cases of P&S syphilis, 18.3% were missing sex of sex partner; 2.7% of male cases with sex of sex partner data were missing source.
† HMO = health maintenance organization; MSW = men who have sex with women only; MSM = men who have sex with men

Sources: http://www.cdc.gov/std/stats10/figures/46.htm

Syphilis Screening

- Routine screening of asymptomatic adolescents is NOT recommended
- Certain groups should be screened:
  - Pregnant women
  - MSM (especially with risk factors present)
  - HIV-positive patients
  - Inmates at correctional facilities, depending on local prevalence of infectious ES

Sources: 2010 CDC STD Treatment Guidelines

Risk Factors for Early Syphilis (ES)

- HIV-positive status (OR 7.3)
  - Co-infection ~20 – 60%
- Anonymous sex partners (OR 2.0)
- “Barebacking” (OR 2.6)
- recreational illicit and/or prescription drug use
  - Crystal meth (OR 3.2)
  - Viagra® (OR 2.1)
  - Crystal meth + Viagra® (OR 6.2)
- Oral sex (“safe sex”?)
  - Risk for syphilis vs. HIV transmission (13.7% vs. 0.31%)
  - Availability of HAART = False sense of security
  - Patients with HIV and syphilis report being on HAART (69%) and having an undetectable VL (58%)


Primary & Secondary Syphilis

Tissue Diagnosis

- Dark-field microscopy examination and/or Direct Fluorescent Antibody (DFA) testing of lesion


Serologic Diagnosis: Screening Tests

- Non-Treponemal tests
  - Rapid Plasma Reagin (RPR)
  - Venereal Disease Research Laboratory (VDRL)
  - Reported as highest dilution titer at which agglutination is visibly appreciated (i.e. NR, 1:1, 1:2, 1:4, ... 1:256, etc.)
  - Markers of disease activity
  - Used to monitor response to treatment
- False (+) can occur! (i.e. autoimmune conditions, pregnancy, older age, IVDU, other spirochetal infections, etc.)
- False (-) can also occur! (i.e. early PS, “prozone” reaction, late latent untreated infection; early treatment)


Serologic Diagnosis: Confirmatory Tests

- Specific Treponemal tests
  - Fluorescent Treponemal Antibody Absorbed assay (FTA-ABS)
  - Treponemal pallidum particle agglutination assay (TP-PA)
  - Immunoassays (EIA, CIA, and MBIA)
  - Once positive, usually remain so for life
- False (+) can occur! (i.e. low prevalence populations, autoimmune conditions, endemic treponematoses, etc.)
- False (-) can also occur! (i.e. early treatment)

**Treatment of Early Syphilis**

- **PS & SS** → Benzathine PCN-G 2.4MU IM x 1
- **ELS** → Benzathine PCN-G 2.4MU IM x 1
- **PCN-Allergy**
  - Doxycycline 100mg PO BID x 14 days
  - Tetracycline 500mg PO QID x 14 days
- **Additional non-traditional regimens**
  - Ceftriaxone 1g IM/IV QD x 10-14 days
  - Azithromycin 2g PO x 1 (not recommended in MSM or pregnant women)
- **Pregnancy** – only use PCN!!
- **Recommended Benzathine PCN-G regimens are same for HIV (-) and HIV (+) patients**

**Sources:** 2010 CDC STD Treatment Guidelines

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**Early Syphilis Follow-Up**

- **PS & SS** → Repeat RPR/VRL at 6 and 12 months for HIV (-) and 3, 6, 9, 12, and 24 months for HIV (+) patients
- **ELS** → Repeat RPR/VRL at 6, 12, and 24 months for HIV (-) and 6, 12, 18, and 24 months for HIV (+) patients
- **Appropriate serologic response to treatment:**
  - PS & SS → ≥4-fold decrease in RPR/VRL titer between 6 and 12 months
  - ELS → ≥4-fold decrease in RPR/VRL titer between 12 and 24 months.
- **Response to treatment is variable and a slower decrease in titers may be seen in patients with prior history of syphilis and/or HIV infection**

**Sources:** 2010 CDC STD Treatment Guidelines

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**Need for Lumbar Puncture in ES**

- **Patients with NEUROLOGIC SYMPTOMS** (i.e. cognitive dysfunction, motor or sensory deficits, ophthalmic or auditory symptoms, CN palsies, symptoms of meningitis)
- **Patients who meet criteria for suspected treatment failure:**
  - Recurrence or persistence of symptoms
  - **PS & SS** → Lack of ≥4-fold decrease in RPR/VRL titer
  - **ELS** → If pre-treatment RPR ≥1:32 and lack of ≥4-fold decrease in titer
  - ≥4-fold increase in titer at any time after treatment
- **No data to support routine LP of HIV-positive patients with ES and with RPR ≥1:32 and/or CD4 count ≤ 350 cells/mL and no neurologic symptoms**

**Sources:** 2010 CDC STD Treatment Guidelines
Follow-Up with Sex Partners

- Notification and screening of sex partners:
  - PS → Last 90 days + symptom duration
  - SS → Last 6 months + symptom duration
  - ELS → Last 1 year
- Presumptive treatment of exposed sex partners:
  - All partners exposed < 90 days before diagnosis of PS, SS, and ELS.
  - All partners exposed >90 days before diagnosis of PS, SS, and ELS, if screening results not available immediately or follow-up is questionable.
  - Partners of patients diagnosed with “Syphilis of Unknown Duration” and RPR ≥1:32, should be managed as ELS

Sources: 2010 CDC STD Treatment Guidelines

Herpes Simplex Virus (HSV)

- Most prevalent ulcerative STI in the US
- Life-long chronic infection with intermittent episodic outbreaks of mucocutaneous and/or genital ulcers.
- Two subtypes:
  - HSV-1 → orolabial infection (but, increasing rates of anogenital infection in MSM and young women)
  - HSV-2 → genital infection
- U.S. → ~50 million people infected w/ genital HSV-2
  - Majority have never been diagnosed due to mild or subclinical infection
  - Viral shedding from genital tract (even when asymptomatic) accounts for majority of transmissions


HSV Risk Factors

<table>
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<tr>
<th>Antibodies, demographics</th>
<th>Odds ratio</th>
<th>95% Confidence interval</th>
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</table>

Sources: Gibson et al. J Infect Dis 1990;162:306

HSV Screening Recommendations

- Serologic screening for HSV-1 and HSV-2 not recommended in general population
- Type-specific serologic screening considered for:
  - Recurrent anogenital ulcers or atypical symptoms with negative HSV cultures
  - Clinical diagnosis of anogenital HSV without lab confirmation
  - Sex partner reports anogenital HSV infection
  - Persons with multiple partners that present for STI evaluation
  - HIV-positive individuals
  - MSM at high risk for HIV infection

Sources: 2010 CDC STD Treatment Guidelines
### HSV Clinical Manifestations

- **Primary infection**
  - Infection of a previously seronegative patient
  - Mild (can be asymptomatic!) to severe
  - Multiple painful ulcers/vesicles
  - Systemic symptoms (i.e. fevers, HAs, lymphadenopathy, etc.)
  - May have neurologic involvement

- **Non-primary infection**
  - Infection with HSV-2 in person already seropositive for HSV-1 or vice-versa
  - Symptoms usually milder than with primary infection

### HSV Diagnosis

- **Clinical diagnosis**
  - Low sens and spec
- **Cytologic detection, i.e. Tzanck Smear**
  - Low sens and spec
- **Cell Viral Culture**
  - Low sensitivity
  - Sensitivity decreases as lesions heal and with recurrent lesions
  - Allows for viral typing (HSV-1 vs. HSV-2)

- **HSV DNA PCR**
  - High sens and spec
  - Allows for viral typing (HSV-1 vs. HSV-2)
  - Useful in late clinical lesions
  - Test of choice in CSF
- **Type specific glycoprotein-G serologic assays (IgG)**
  - Sensitivity 80-98%
  - Specificity ≥ 96%
  - False (-) in early infection
  - False (+) in patients with low likelihood of infection

Testing for IgM is not useful or recommended!!

### HSV Clinical Manifestations

- **Recurrent infection**
  - Reactivation of latent infection (outbreaks)
  - Milder than primary or non-primary infection
  - May be subclinical or asymptomatic (~25%)

### Genital HSV Treatment (Some Examples)

- **First Clinical Episode (always treat, even if mild symptoms)**
  - Acyclovir 400mg PO TID x 7 – 10 days
  - Valacyclovir 1g PO BID x 7 – 10 days

- **Episodic Recurrences (start w/ prodrome or w/in 24hr of vesicle appearance)**
  - Acyclovir 400mg TID OR 800mg BID x 5 days
  - Valacyclovir 500mg PO BID x 3 days

- **Suppressive Therapy (decreases recurrences by ~70-80% and subclinical viral shedding by ~95%)**
  - Acyclovir 400mg PO BID
  - Valacyclovir 500mg* or 1g PO daily (*may be less effective if very frequent recurrences reported; ≥10 episodes/year)

### Sources:
Case 2 – “Rash and Sores”

- RPR 1:1,024 and Reactive FTA-ABS
- Diagnosis – Secondary Syphilis with characteristic body rash and mucous patches in tongue
- Benzathine PCN-G 2.4MU IM x 1
- Follow-up RPRs at 3, 6, 9, and 12 months
- Attempt to notify, screen, and treat any sex partners met online within last 6 months with the help of Disease Intervention Specialists (DIS)
- Education and counseling regarding safer sexual practices and modes of syphilis transmission

HPV and Genital Warts (GW):
Diagnosis and Clinical Manifestations

- Majority of HPV infections are subclinical, completely asymptomatic, and clear spontaneously.
- When present, diagnosis of GW is mainly clinical by visual inspection
- GW are highly infectious (64% of contacts develop GW).
- GW are usually asymptomatic, but may be pruritic or bleed.
- Large GW may cause anatomic obstruction

Human Papillomavirus (HPV)

- >50% of sexually active people are infected at least once in life.
- >100 types of HPV and >40 are known to infect the genital area.
- Oncogenic or High-Risk (HR-HPV types 16 and 18)
  - Cause majority of cervical, anal, penile, vulvar, and vaginal CA
  - Oropharyngeal CA (HPV type 16)
- Non-Oncogenic or Low-Risk (LR-HPV types 6 and 11)
  - Cause ~90% of genital warts (GW)
  - Efficient rates of transmission among sex partners
    - Penis → Cervix (4.9 infections / 100 person-months)
    - Cervix → Penis (17.4 infections / 100 person-months)
  - Limited data on transmission via non-intercourse sexual contact

Genital Wart (GW) Treatment

<table>
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<th>Anatomic Location</th>
<th>Podophyllin*</th>
<th>Imiquimod*</th>
<th>cryosurgery</th>
<th>Curettage</th>
<th>TCA/ACCA</th>
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<td>Perianal</td>
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*Adapted from the CDC’s Centers for Disease Control and Prevention: Sexually Transmitted Disease Treatment Guidelines.
*Non-ablative treatments include intralesional interferons, laser therapy, topical antivirals, and other.
This expert recommeneded uses of podophyllin for subclinical low-risk lesions.

HPV DNA Tests: General Overview

- **Recommended:**
  - Triage of women ≥21 years who have ASC-US cytology results
  - Routine adjunctive testing in combination with cervical cytology for women ≥30 years.

- **Not Recommended:**
  - LR-HPV DNA testing (No clinical indications)
  - STD screening for HPV (including sex partners)
  - Deciding whether to administer vaccine or not.
  - Testing adolescents < 21 years old.
  - Screening for primary cervical CA as stand-alone test (i.e. w/o cervical cytology)
  - Triaging Low-grade squamous intraepithelial lesions (LSIL)

Source: 2010 CDC STD Treatment Guidelines

Final Thoughts

- We all share the responsibility of helping to fight the epidemic of STIs that affect this country.
- We must inquire about specific risk factors for common STIs (i.e. Syphilis).
- Implement a guideline-based STI screening protocol for your at-risk patient population.
- Consult with your local health department, sexual health clinic, ID physician, and/or the 2010 CDC STD Treatment Guidelines for the latest diagnostic and treatment recommendations.

HPV Vaccine Recommendations

<table>
<thead>
<tr>
<th>Vaccine</th>
<th>Quadrivalent HPV vaccines</th>
<th>Bivalent HPV vaccines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Location in United States</td>
<td>2008</td>
<td>2009</td>
</tr>
<tr>
<td>HPV vaccine guidance</td>
<td>Routine vaccination 11 or 12 year-old male begins at 9-12 year-old while</td>
<td>Routine vaccination 11 or 12 year-old prior</td>
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<td>W/ V vaccine types</td>
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<tr>
<td>Efficacy against HPV 16/18-related CIN (CAD)</td>
<td>90%</td>
<td>90%</td>
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<td>Efficacy against HPV 6/11-related genital lesions</td>
<td>90%</td>
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