## Best Practices in the Diagnosis and Treatment of *Clostridioides difficile* Infections

### Acute Care

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| **Best Practices in the Diagnosis and Treatment of *Clostridioides difficile* Infections**
**Acute Care**

**SAY:**

This presentation will address “Best Practices in the Diagnosis and Treatment of *Clostridioides difficile* Infections.” |

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<tr>
<th><strong>Objectives</strong></th>
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| **SAY:**

By the end of this presentation, participants will be able to—

- Discuss the importance of judicious *Clostridioides difficile* or *C. difficile* laboratory testing
- Discuss management approaches for *C. difficile* infections or CDI
- Discuss the role of antibiotics, gastric acid suppressive agents, and probiotics in inciting or preventing *C. difficile* infections |

1. Discuss the importance of judicious *C. difficile* laboratory testing
2. Discuss management approaches for *C. difficile* infections (CDI)
3. Discuss the role of antibiotics in preventing CDI
4. Discuss the role of gastric acid suppressive agents in inciting CDI
5. Discuss the role of probiotics in preventing CDI
## The Four Moments of Antibiotic Decision Making

**SAY:**

As we discuss the diagnosis and treatment of CDI, we will continue to use the Four Moments of Antibiotic Decision Making framework.

As a reminder, the Four Moments include:

Moment 1: Does my patient have an infection that requires antibiotics?

Moment 2: Have I ordered appropriate cultures before starting antibiotics? What empiric therapy should I initiate?

Moment 3: A day or more has passed. Can I stop antibiotics? Can I narrow therapy or change from intravenous to oral therapy?

Moment 4: What duration of antibiotic therapy is needed for my patient’s diagnosis?

### Slide 3

**The Four Moments of Antibiotic Decision Making**

1. Does my patient have an infection that requires antibiotics?
2. Have I ordered appropriate cultures before starting antibiotics? What empiric therapy should I initiate?
3. A day or more has passed. Can I stop antibiotics? Can I narrow therapy or change from IV to oral therapy?
4. What duration of antibiotic therapy is needed for my patient’s diagnosis?

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**SAY:**

We will begin with Moment 1, “Does my patient have an infection that requires antibiotics?”

### Slide 4

**The Four Moments of Antibiotic Decision Making**

1. Does my patient have an infection that requires antibiotics?
Colonization of the intestinal tract with *C. difficile* occurs via the fecal-oral route. This process is facilitated by antibiotic use. Antibiotics disrupt the barrier function of normal colonic flora, providing a niche for *C. difficile* to multiply and produce toxins. *C. difficile* releases two toxins that mediate disease: toxin A and toxin B. Toxin B appears to be the more clinically important toxin. CDI has not been shown to occur in the absence of toxin B production. These toxins lead to inflammation and diarrhea. Antibodies to the toxins, however, can be protective. Asymptomatic carriers often demonstrate higher serum levels of antibodies against *C. difficile* toxins compared to patients who develop clinical disease. As will be discussed later in the presentation, monoclonal antibodies commercially available against *C. difficile* toxin appear to reduce the recurrence rate of CDI.

The case definition of CDI is as follows as of April 2019: At least three unformed stools within a 24-hour period and either a positive stool test for the *C. difficile* toxin or colonoscopic or histopathologic findings compatible with pseudomembranous colitis. Presence of the *C. difficile* organism in the absence of toxin production should not be considered an infection.
### C. difficile Clinical Spectrum

**SAY:**

Asymptomatic carriage with *C. difficile* is common. In healthy infants under 1 year of age, the prevalence of *C. difficile* colonization has been described as between 10 and 60 percent in various studies. Reasons for the high prevalence of *C. difficile* in infants are unclear. Despite higher rates of colonization, clinically apparent disease in infants is uncommon. Based on animal models, it has been proposed that the absence of toxin receptors is the main reason for the rarity of clinical disease in human infants.

By 2 years of age, the risk of colonization is the same as healthy adults—at about 3 percent. The risk of colonization has been shown to increase linearly with the duration of hospital stay; 10–20 percent of hospitalized patients test positive for *C. difficile* toxin.

The most common clinical presentation of *C. difficile* infection is watery diarrhea. Additional findings include lower abdominal pain, cramping, and nausea. Low-grade fevers occur in approximately 15 percent of patients. Leukocytosis may be present and on average the serum white blood cell count is approximately 15,000 cells per microliter.

Severe or fulminant colitis may present with severe abdominal pain, a possible ileus, lactic acidosis, hypoalbuminemia, and/or significant leukocytosis.

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<td><strong>C. difficile Clinical Spectrum</strong></td>
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| • Asymptomatic colonization  
  – 10–60% of healthy infants under 1 year of age\(^1\)\(^6\)  
  – 3% of healthy adults\(^1\)\(^5\)  
  – 10–20% of hospitalized patients\(^1\) \(\text{\(\text{\textsuperscript{1}}\)}\) |
| • Watery diarrhea\(^1\)  
  – Most common presentation  
  – Lower abdominal pain, cramping, nausea, low-grade fever (15%), and leukocytosis (average ~15,000 cells/microliter) |
| • Severe or fulminant colitis\(^1\) |

\(\text{\(\text{\textsuperscript{1}}\)}\) AHRQ Safety Program for Improving Antibiotic Use – Acute Care

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AHRQ Safety Program for Improving Antibiotic Use – Acute Care

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### Severe and Fulminant CDI

**SAY:**

As of April 2019, there is no consensus for what defines severe versus fulminant CDI. The 2018 Infectious Diseases Society of America and Society for Healthcare Epidemiology of America guidelines recommend considering CDI as severe when a leukocytosis of greater than 15,000 cells per milliliter is present or the serum creatinine is greater than 1.5 mg/dL. Fulminant CDI is generally defined as CDI requiring ICU admission because of hypotension, intestinal perforation, or toxic megacolon. Patients with fulminant CDI who survive the infection may require a colectomy.

Prompt surgical consultations are recommended for all cases of severe or fulminant CDI. Abdominal imaging is important to evaluate for the presence of toxic megacolon, bowel perforation, or other findings warranting surgical intervention.

### Ribotype 027 (NAP1) Strain

**SAY:**

The NAP1 or ribotype 027 strain is a hypervirulent strain of *C. difficile* that has become increasingly recognized since the early 2000s. This strain is associated with frequent, severe, refractory disease that is more likely to relapse than non-NAP1 strains. This may be because the NAP1 strain generally produces large quantities of toxin.

### Slide 8

**Severe and Fulminant CDI**

- 2018 IDSA/SHEA guidelines state CDI is severe when—
  - Leukocytosis greater than 15,000 cells per milliliter or serum creatinine greater than 1.5 mg/dL.
- Fulminant CDI defined as CDI requiring ICU admission because of—
  - Hypotension
  - Intestinal perforation
  - Toxic megacolon
- Obtain abdominal imaging and prompt surgical consultation

### Slide 9

**Ribotype 027 (NAP1) Strain**

- Associated with frequent, severe, refractory disease that is more likely to relapse compared to non-NAP1 strains
  - Toxin production 16–23-fold greater than wild-type strains
- Fluoroquinolone (FQ) use has been associated with the emergence of the NAP1 strain
  - Reduction in FQ use in the United Kingdom has been associated with dramatic reductions in CDI due to NAP1.
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| Recurrent CDI is defined as a resolution of CDI symptoms, followed by subsequent reappearance of symptoms after treatment has been discontinued. It is helpful to forewarn patients that they may have loose stools and symptoms consistent with irritable bowel syndrome for some time after completing CDI treatment, particularly if still receiving antibiotics. This is usually not indicative of recurrent of CDI. Patients should be retested to confirm the patient indeed has CDI by testing specifically for the presence of toxin and not just simply the presence of the organism, as persistence of the organism may occur in the absence of toxin production. Up to 30 percent of patients experience recurrent CDI within 30 days of treatment. It may not occur immediately after discontinuing antibiotics. CDI recurrences have been shown to present as late as 3 months after the discontinuation of CDI treatment. The risk of another episode increases with each successive recurrence. Recurrent symptoms may be due to a relapse of the initial infecting strain or a reinfection with a new strain, although the former is more common. Among 102 patients with recurrent CDI, isolates obtained 2 to 8 weeks apart were identical in 88 percent of cases. | - Resolution of CDI symptoms followed by reappearance of symptoms after treatment has been discontinued[11]
- ~30% of patients experience recurrent CDI within 30 days of treatment[12]
- Recurrent symptoms may be due to relapse of the initial infecting strain or reinfection with a new strain[12,13] |
| **The Four Moments of Antibiotic Decision Making** | **Slide 11** |
| SAY:                      | The Four Moments of Antibiotic Decision Making |
| Moment 2 reminds us to ensure appropriate testing has occurred once a patient has clinical signs or symptoms suggestive of an infection. | 1. Does my patient have an infection that requires antibiotics?
2. Have I ordered appropriate cultures before starting antibiotics? What empiric therapy should I initiate? |
Several diagnostic tests are available to identify *Clostridioides difficile* infection. The enzyme immunoassay can identify toxin A and B production. It is relatively easy to perform, and results return within 4 hours. The major drawback with this test is that it has limited sensitivity.

There are nucleic acid tests that identify the genes that produce toxin A and B. These are very sensitive, but they cannot distinguish whether the genes are activated and making toxin or present but inactive. Thus, there is concern that they will detect large portions of patients with asymptomatic colonization—likely leading to overdiagnosis and overtreatment.

Glutamate dehydrogenase is an enzyme produced by all *C. difficile* isolates. It is rapid, relatively inexpensive, and easy to perform. It has very good sensitivity for detecting the presence of *C. difficile* but its specificity is poor because it does not distinguish strains that do or do not produce toxin.

Finally, the cytotoxin assay is considered the “gold standard” for *C. difficile* detection. The test is performed by adding a prepared stool sample to a monolayer of cultured cells. If *C. difficile* toxin is present, it exerts a cytopathic effect. This test has a sensitivity approaching 100 percent, but it is time consuming to perform and results take up to 48 hours. Some laboratories perform “two-step” tests which initially look for the presence of the organism, followed by testing for the toxin if the organism is present. For example, initial testing may look for the production of glutamate dehydrogenase, and if present, then EIA testing for toxins A and B is performed. Or the initial test might be a nucleic acid test followed by the EIA to evaluate for toxin production.

The 2018 IDSA/SHEA guidelines suggest that a nucleic acid test alone is reasonable if there are established criteria for stool testing for *C. difficile* in a facility—for example, only testing unformed stools. But in the absence of these institutional recommendations, the guidelines recommend using a multistep testing algorithm.
Additional Tips Related to Laboratory Testing

SAY:

Keeping in mind the high rates of asymptomatic colonization with *C. difficile*, particularly in patients with recent or active health care exposure, it is important to obtain *C. difficile* testing for patients with a high clinical suspicion for CDI. Testing should be limited to patients with at least three unformed stools within the past 24 hours and without a reasonable alternative explanation, with an exception being fulminant disease with ileus.

At least one study showed that screening and isolating asymptomatic *C. difficile* carriers reduced the incidence of CDI, but there were some limitations with this study. It spanned over a decade, so other changes that occurred during this time period may have impacted outcomes, including the improved institutional hand hygiene compliance over time. Improvements in antibiotic use over time were not evaluated in this study. Similar studies need to be performed to better understand the pros and cons of screening and isolating asymptomatic carriers. As of April 2019, identifying asymptomatic carriers is not considered standard practice. One concern with this approach is that patients with asymptomatic colonization may unnecessarily receive *C. difficile* therapy.

One of the most common reasons people have loose stools in the hospital setting is laxative use. In one study conducted in an academic medical center, 44 percent of patients with positive nucleic acid testing for *C. difficile* received laxatives within the previous 48 hours.

It is important to confirm a patient has not recently received a laxative prior to sending *C. difficile* testing. Also, tests of cure are not necessary. Once a patient has responded to antibiotic therapy, retesting for *C. difficile* is not needed as asymptomatic colonization can persist in the absence of disease.

Because of the high prevalence of *C. difficile* colonization in infants and the very low likelihood of *C. difficile* clinical infection in infants under a year of age, testing for *C. difficile* is discouraged in this age group. Many institutions have expanded this to limit testing for...
### Slide Title and Commentary

children under 2 years of age. Some laboratories require approval by an infectious diseases clinician prior to pursuing *Clostridioides difficile* testing in young children.

Other mechanisms that have been successfully used to limit unnecessary testing of *Clostridioides difficile* include not testing for *Clostridioides difficile* more than once in 7 days, having automatic popups in the electronic health record reminding clinicians to ensure that a patient is not receiving a laxative prior to sending *Clostridioides difficile* testing, or laboratory refusal to test formed stool samples for *Clostridioides difficile*.

### The Four Moments of Antibiotic Decision Making

**SAY:**

The second portion of Moment 2 as well as Moments 3 and 4 focus on antibiotic therapy. Empiric therapy is not recommended for CDI, unless there is a high suspicion for CDI and the presentation is fulminant, particularly if significant delays are anticipated while awaiting laboratory confirmation. As there are generally not opportunities for de-escalation of CDI therapy and the duration of therapy is relatively standardized, Moments 3 and 4 can be combined for CDI management.

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**The Four Moments of Antibiotic Decision Making**

1. Does my patient have an infection that requires antibiotics?
2. Have I ordered appropriate cultures before starting antibiotics? What empiric therapy should I initiate?
3. A day or more has passed. Can I stop antibiotics? Can I narrow therapy or change from IV to oral therapy?
4. What duration of antibiotic therapy is needed for my patient’s diagnosis?
A number of treatment principles should be considered.

First, do not treat asymptomatic patients with a positive *C. difficile* test. Always confirm that the patient has signs and symptoms consistent with CDI before prescribing therapy.

Second, if possible, discontinue any antibiotic therapy the patient is receiving that is not being administered to treat CDI. Continuing additional antibiotic agents may decrease clinical response and increase CDI recurrence rates.

If additional antibiotics are needed because of proven infection, or in a situation where it is not entirely clear if a patient has sepsis from an abdominal source or CDI, select the narrowest agent possible. Avoid agents with a strong association with CDI such as third- and fourth-generation cephalosporins, fluoroquinolones, or clindamycin. As there are some data suggesting the risk of CDI is lower with tetracyclines compared with most other antibiotic classes, consider a tetracycline (such as doxycycline or tigecycline) if appropriate based on the clinical scenario.

Discontinue gastric acid suppression medications whenever possible. Additionally, avoid antimotility agents as some studies suggest that these agents could lead to poor outcomes.

- Do not treat asymptomatic patients with a positive *C. difficile* test
  - Confirm symptoms consistent with CDI exist prior to prescribing therapy
- If possible, discontinue any antibiotic therapy not specifically treating CDI
- If additional antibiotic therapy is necessary:
  - Select the narrowest agent possible
  - Avoid agents with a strong association with CDI
  - Consider a tetracycline if appropriate
- Discontinue gastric acid suppression medications
- Avoid antimotility agents
**Treatment of Initial Episode**

**SAY:**

In the past, metronidazole was advocated for mild to moderate cases of CDI, but the 2018 IDSA/SHEA guidelines recommend enteral vancomycin 125 mg by mouth 4 times a day or fidaxomicin 200 milligrams twice a day for mild, moderate, or severe disease. For children, either oral metronidazole or oral vancomycin are considered reasonable options for nonsevere cases. There are no randomized controlled trials evaluating the comparative efficacy of these agents for CDI in children. For children with an initial episode of severe CDI, oral vancomycin is recommended over oral metronidazole.

Of note, intravenous vancomycin has no effect on CDI because of its minimal excretion into the colon. The liquid formulation of vancomycin compounded from powder which is intended for intravenous administration can be safely and effectively administered as oral vancomycin. This approach is more cost effective than using the standard oral vancomycin formulation, especially while a patient is hospitalized.

**Treatment of Fulminant Disease**

**SAY:**

In cases of fulminant CDI, vancomycin in higher dosages (500 mg instead of 125 mg) is recommended although there is a lack of evidence of a clear benefit with this increased dose. Intravenous metronidazole should also be considered for fulminant CDI, particularly if an ileus prevents the delivery of oral agents to the colon. If an ileus is present, vancomycin can also be administered per rectum as a retention enema, along with intravenous metronidazole. It is anticipated that IV metronidazole is likely to achieve therapeutic concentrations in an inflamed colon. For fulminant CDI, early surgical consultation is essential.
A first recurrence of CDI in adults can be treated with either oral vancomycin as a tapered and pulse regimen or a 10-day course of fidaxomicin. Although the risk of recurrence may be lower with fidaxomicin compared with a tapered and pulse regimen of vancomycin, medication costs should be factored into this decision.

The theoretical benefit of a tapered and pulse vancomycin regimen that extends over several weeks is that it may keep growth of *C. difficile* under control while allowing normal flora to repopulate. A commonly used prolonged tapered and pulse regimen consists of 125 mg of oral vancomycin 4 times per day for 10 to 14 days, followed by twice a day for a week, then once a day for a week, and then every 2 to 3 days for 2 to 8 weeks.

If metronidazole was used for the primary episode, it is recommended that a first recurrence of CDI is treated with a standard 10-day course of vancomycin.

For children, oral vancomycin is recommended for a first recurrence.

Fecal microbiota transplantations should be considered for both children and adults with multiple CDI recurrences.
### Fecal Microbiota Transplantation

**SAY:**

The colon harbors a stable community of microorganisms which exist in symbiosis with the host. Antibiotics lead to the selective removal of bacteria that serve as a barrier to colonization with *C. difficile*. Fecal microbiota transplantations or FMTs promote intestinal diversity similar to what was present prior to antibiotic exposure.

There are a number of observational and randomized controlled trials that have evaluated the efficacy of FMTs. Treatment success in these studies range from 50 to 100 percent when used for patients with recurrent CDI. The highest success rates appear to be associated with instillation of feces via colonoscopy rather than the nasogastric route.

### Monoclonal Antibodies

**SAY:**

Monoclonal antibodies against *C. difficile* toxin appear to reduce the recurrence rate of CDI. Bezlotoxumab is a monoclonal antibody directed against toxin B that was approved by the Food and Drug Administration in 2016 as adjunctive therapy for patients with a previous history of CDI who are receiving antibiotic treatment for CDI and who are at high risk for recurrence. Bezlotoxumab together with standard therapy has been associated with lower recurrence rates than standard therapy alone—17 percent versus 28 percent. The role of monoclonal antibodies to prevent and treat CDI recurrences is still being defined as of April 2019.
Select Modifiable Risk Factors

SAY:

Some nonmodifiable risk factors place people at increased risk of CDI. These include elderly age, inflammatory bowel disease, solid organ transplant recipient, or Hirschsprung’s disease in young children.

Some important modifiable risk factors alter a patient’s risk of developing CDI including antibiotic use, gastric acid suppression, probiotic use, and infection control practices. Although the role of lapses in infection control practices is well recognized in the transmission of CDI, this presentation will not review the role of infection prevention practices in limiting the spread of C. difficile.

To observe a demonstrable decrease in CDI rates, a multifaceted approach is generally necessary that starts with judicious testing for C. difficile as well as appropriate antibiotic use, judicious gastric acid suppression use, and infection prevention practices. The reference on this slide by Abbett and colleagues, published in Infection Control and Hospital Epidemiology in 2009, includes a checklist that reviews modifiable risk factors, including enhanced isolation practices and laboratory notification procedures, which successfully reduced the incidence of CDI by 40 percent at a tertiary-care hospital.

Clostridioides difficile and Antibiotics

SAY:

Virtually all antibiotics can increase the propensity for the development of CDI, but the greatest risk exists with third- and fourth-generation cephalosporins, fluoroquinolones, and clindamycin. The risk is highest when patients are receiving antibiotics, but the risk remains elevated up to 12 weeks later. Some data suggest that tetracyclines such as doxycycline or tigecycline may be protective as they have some activity against C. difficile growth and appear to inhibit toxin production. It also appears they have minimal effects on a number of healthy gut flora.
CDI Reductions Associated With Reduction in Cephalosporins and FQ

SAY:

A large quasi-experimental study was undertaken to assess the impact of national antibiotic stewardship and infection control programs on CDI rates in Scotland, evaluating data from over 1 million patients. Stewardship activities included empiric guidelines recommending against the use of agents such as fluoroquinolones, clindamycin, and cephalosporins and finding suitable replacement agents. Approval was needed for use of these agents and susceptibilities of these agents were not routinely provided. In addition, improvements in infection control practices occurred. These interventions led to a 68 percent reduction of CDI in Scottish hospitals.
### How Can We Reduce the Use of High Risk Agents?

SAY:

What are some ways we can reduce the use of high-risk antibiotic agents? Keeping in mind the elevated risk of CDI with antibiotics such as ceftriaxone, cefepime, clindamycin, and fluoroquinolones, use of these agents should be limited as much as possible.

Think about infectious processes where these agents are commonly used in your facility and consider alternative agents. For example, ceftriaxone is commonly used for the treatment of community-acquired pneumonia. Consider replacing it in your local community-acquired pneumonia guidelines with ampicillin or ampicillin/sulbactam. Ceftriaxone is also commonly used for cystitis. The reality is most cystitis in the hospital is really asymptomatic bacteriuria. Before prescribing treatment, confirm that the patient has urinary symptoms. Further, intravenous therapy generally is not needed for cystitis. If treatment appears necessary, consider 5 days of nitrofurantoin, 3 days of trimethoprim/sulfamethoxazole, or 7 days of cephalexin for cystitis.

Cefepime is commonly used for neutropenic fever. Although all antipseudomonal beta-lactam antibiotics have the potential for adverse events, if you are having higher rates of CDI in your oncology wards than expected, consider replacing your standard neutropenic fever beta-lactam with piperacillin/tazobactam. Data for both children and adults suggest the risk of CDI is lower in units where piperacillin-tazobactam is used as the standard neutropenic fever agent compared with other anti-pseudomonal beta-lactams.
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**How Can We Reduce the Use of High-Risk Agents?**

- **Clindamycin**
  - Consider cephalexin for nonpurulent cellulitis
  - Consider trimethoprim/sulfamethoxazole or doxycycline for mild purulent cellulitis when methicillin-resistant *Staphylococcus aureus* is a consideration. Rates of resistance to clindamycin for both methicillin-susceptible and -resistant *S. aureus* have increased in many parts of the county, making clindamycin a poor choice for purulent cellulitis in these areas.

- **Fluoroquinolones**
  - Avoid for cystitis
  - Limit use for community-acquired pneumonia to severe penicillin allergies

- **Overall**
  - De-escalate therapy whenever possible
  - Limit durations of therapy

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**How Can We Reduce the Use of High-Risk Agents?**

**SAY:**

Clindamycin is frequently used for both nonpurulent and purulent cellulitis. For nonpurulent cellulitis, consider the use of cephalexin. Consider trimethoprim/sulfamethoxazole or doxycycline for mild purulent cellulitis when methicillin-resistant *Staphylococcus aureus* is a consideration. Rates of resistance to clindamycin for both methicillin-susceptible and -resistant *S. aureus* have increased in many parts of the county, making clindamycin a poor choice for purulent cellulitis in these areas.

Fluoroquinolones should be avoided for cystitis and intra-abdominal infections whenever possible. Additionally, consider limiting their use for community-acquired pneumonia to patients with severe penicillin allergies.

And, in general, always remember to stop or de-escalate antibiotic therapy whenever possible when reviewing Moment 3, and limit the duration of antibiotic therapy to what is appropriate for the condition you are treating when considering Moment 4.
### Gastric Acid Suppression

**SAY:**

Proton-pump inhibitors and histamine receptor antagonists have been associated with an increased risk of CDI in both pediatric and adult studies.

In an observational study including 650 children with CDI, children with CDI had 22 times the odds of having recently receiving PPIs within the previous 90 days than children without CDI. Based on the results of three meta-analyses including over 45 observational studies of adults, adults with CDI had about twice the odds of having recently received PPIs than adults without CDI.

Evaluating Centers for Disease Control and Prevention data from 2009 to 2011, which included a relatively large percentage of community-associated CDI cases, 36 percent did not report antibiotic exposure in the preceding 12 weeks. However, among patients without reported antibiotic exposure, 31 percent received proton pump inhibitors. The increased risk of CDI is likely due to a decrease in the protective effect of stomach acid. This decreased protective effect allows the entry and survival of *C. difficile* in the upper gastrointestinal tract.

The relationship between the route and duration of gastric acid suppressive agents and CDI is unknown. Although some debate exists regarding the true impact of gastric acid suppressive agents on CDI, as with all medications, their use should be limited to when they are clearly necessary.

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- Proton-pump inhibitors (PPIs) and histamine 2 receptor antagonists
  - Associated with an increased risk of CDI in studies of children and adults
- Likely due to breaches in the protective effect of stomach acid facilitating the entry and survival of *C. difficile* in the upper gastrointestinal tract
- Relationship between the risk of CDI and route and duration of gastric acid suppressive agents is unknown
- Limit gastric acid suppressive agent use whenever possible
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**Probiotics**

**SAY:**

Probiotics include microorganisms that may reduce the risk of colonization by pathogenic bacteria. They are becoming increasingly available as capsules and dairy-based food supplements. Several meta-analyses have indicated probiotics may be effective at preventing CDI when given to patients receiving antibiotics. However, randomized trial data have not shown the same benefit.

There have also been several meta-analyses that have included randomized trials that suggest a benefit of probiotics when administered to patients with mild to moderate CDI. Similar data are not available for patients with severe CDI.

**Bottom Line With Probiotics**

**SAY:**

The role of probiotics in preventing and treating CDI is evolving. They are not routinely recommended as of April 2019.

There continue to be gaps in knowledge regarding the right strains, dosages, frequency, and duration of probiotics for both the prevention and treatment of CDI. There are no current requirements to demonstrate safety or manufacturing consistencies of probiotics to consumers. Probiotics generally seem associated with little harm and may have a benefit for immunocompetent patients. But they may cause harm if administered to immunocompromised patients. There have been cases of probiotic-associated bacteremia or fungemia in immunocompromised patients as well as premature neonates. Although the role of probiotics in the prevention or treatment of CDI is still not clear, it is important to remember that encouraging probiotic use is not a standalone intervention if you are trying to make changes in your facility to reduce CDI rates.

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**Probiotics**

- Include microorganisms that may reduce the risk of colonization by pathogenic bacteria
- Becoming increasingly available as capsules and dairy-based food supplements
- Prevention of CDI
  - Several meta-analyses indicate probiotics may be effective at preventing CDI when given to patients receiving antibiotics
  - No randomized trials have shown a benefit with probiotics in preventing CDI
- Treatment of CDI
  - Meta-analysis of RCTs suggest a benefit of probiotics for mild-moderate CDI therapy
  - Data not available for severe CDI

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**Bottom Line With Probiotics**

- Role of probiotics in preventing and treating CDI is evolving and currently not routinely recommended
- Right strain(s), dosage, duration unknown
- No requirement to demonstrate safety or manufacturing consistencies of probiotics to consumers
- Likely little harm and may have a benefit for generally immunocompetent patients
  - Cases of probiotic-associated bacteremia or fungemia have been reported, particularly in immunocompromised patients and premature neonates
- Encouraging probiotic use is not a standalone intervention to reduce CDI.
**Take-Home Messages**

**SAY:**

A few take-home points are worth reemphasizing.

First, judicious *C. difficile* laboratory testing is critical. Only test stool for *C. difficile* when clinical criteria are met!

Make sure you understand the *C. difficile* testing approach used in your institution and work with your microbiology laboratory and other important stakeholders to determine if measures can be undertaken to reduce unnecessary testing.

It is critical to discontinue additional unnecessary antibiotic agents when a diagnosis of CDI is made. If additional antibiotic therapy is necessary, make sure to select the narrowest antibiotic agent(s) possible, avoid using agents with a strong association with CDI (such as ceftriaxone, cefepime, clindamycin, and fluoroquinolones), and limit the duration of antibiotic therapy to the lowest effective duration.

Also, think about where in your institutional guidelines the use of high-risk antibiotic agents can be replaced with agents that pose a lower risk for CDI.

**Disclaimer**

**SAY:**

The findings and recommendations in this presentation are those of the authors, who are responsible for its content, and do not necessarily represent the views of AHRQ. No statement in this presentation should be construed as an official position of AHRQ or of the U.S. Department of Health and Human Services.

Any practice described in this presentation must be applied by health care practitioners in accordance with professional judgment and standards of care in regard to the unique circumstances that may apply in each situation they encounter. These practices are offered as helpful options for consideration by health care practitioners, not as guidelines.
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