Fecal Microbiota Transplant

Policy MP-062

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Disclaimer:
1. Policies are subject to change in accordance with State and Federal notice requirements.
2. Policies outline coverage determinations for U of U Health Plans Commercial, Healthy U (Medicaid) and Advantage U (Medicare) plans. Refer to the “Policy” section for more information.
3. This Medical Policy does not guarantee coverage or payment of the service. The service must be a benefit in the member’s plan and the member must be eligible for coverage at the time of service. Additional payment guidelines may be applied that are not included in this policy.

Description:
Clostridium difficile infection (CDI), is a Gram-positive, spore-forming bacterium usually spread by the fecal-oral route. Patients with recurrent CDI have been observed to have reduced diversity of the intestinal microbiome and diminished numbers of bacteria relative to healthy individuals that can cause disease asymptomatic carriage, mild diarrhea, colitis, or pseudomembranous colitis.

According to the American Journal of Gastroenterology (AJG) Clostridium difficile infection is a leading cause of hospital-associated gastrointestinal illness. Patients typically have extended lengths-of-stay and CDI is a frequent cause of large hospital outbreaks of disease.

Fecal microbiota transplant (FMT) is the term used when stool is taken from a healthy individual and instilled into a sick person with certain conditions, such as recurrent CDI. Studies show that patients with recalcitrant CDI have abnormally proportioned colon microbiota and that reintroduction of normal bacteria via donor feces corrects this imbalance and breaks the cycle of CDI recurrence. The purpose of FMT treatment is based on the premise that an imbalance in the community of microorganisms residing in the gastrointestinal (GI) tract is associated with specific disease states, including susceptibility to infection. In its healthy state, intestinal microbiota performs a variety of useful functions including aiding in the digestion of carbohydrates, repressing the growth of pathogenic microbes, mediating the synthesis of certain vitamins, and stimulating the lymphoid tissue to produce antibodies to pathogens.

FMT may be administered by oral capsules, colonoscopy, retention enema, or through a nasojejunal (NJ)/nasoduodenal (ND) tube in the upper GI tract. The choice of the delivery route
depends in part on patient preferences, individual risk, cost, availability of resources and expertise.

Policy Statement and Criteria

1. Commercial Plans

U of U Health Plans covers fecal microbiota transplantation including capsulized, frozen fecal microbiota transplantation medically necessary for treatment of patients with recurrent Clostridium difficile infection when the following criteria are met:

A. At least 3 episodes of recurrent mild to moderate CDI and failure of a 6-8 week taper with vancomycin with or without an alternative antibiotic (e.g., rifaximin, nitazoxanide, metronidazole, vancomycin); and

B. Persistent positive stool *C. difficile* toxin by testing and at least one of the following:
   i. At least two episodes of recurrent severe CDI resulting in hospitalization and associated significant morbidity, including renal failure.
   ii. Moderate CDI not responding to standard therapy (vancomycin) for at least a 10 days.
   iii. Severe fulminant *C. difficile* colitis with no response to standard therapy after 48 hours.

U of U Health Plans considers fecal microbiota transplantation investigational/experimental in all other circumstances.

2. Medicaid Plans

Coverage is determined by the State of Utah Medicaid program; if Utah State Medicaid has no published coverage position and InterQual criteria are not available, the U of U Health Plans Commercial criteria will apply. For the most up-to-date Medicaid policies and coverage, please visit their website at http://health.utah.gov/medicaid/manuals/directory.php or the Utah Medicaid code Look-Up tool

3. Medicare Plans

Coverage is determined by the Centers for Medicare and Medicaid Services (CMS); if a coverage determination has not been adopted by CMS and InterQual criteria are not available, the U of U Health Plans Commercial criteria will apply. For the most up-to-date Medicare policies and coverage, please visit their search website at: http://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx?from2=search1.asp& or the manual website
**Clinical Rationale**

A 2017 systematic review (Quraishi et al) investigated the effect of FMT in patients with recurrent or refractory CDI. In the pooled analysis, 92% of patients had a resolution of CDI (95% CI, 89% to 94%); heterogeneity was classified as likely moderate (I²=59%). Additionally, in the 7 trials that evaluated FMT, the intervention overall was associated with an increase in the resolution of recurrent and refractory CDI (relative risk, 0.23; 95% CI, 0.07 to 0.80). The 30 case series reported resolution rates for CDI ranged from 68% to 100%. The reviewers considered the RCTs as having a low risk of bias for adequate randomization with allocation concealment and intention-to-treat analysis. Nor did they report an assessment of bias in terms of blinding, sample size adequacy, or possible differences in baseline characteristics. However, they disputed that none of the trials demonstrated the efficacy of FMT as being truly placebo-controlled, and the case series followed patients until resolution of CDI (10 weeks to 8 years), though some had an insufficient follow-up.

In another 2018 meta-analysis (Khan et al.) researched the literature of pooled data on the use of FMT as a treatment option for recurrent CDI. Reviewers only selected randomized controlled trials that compared FMT (fresh or frozen) with medical treatment. Among the selected studies, there was a non-significant trend toward the resolution of diarrhea following a single fresh FMT infusion compared with frozen FMT or medical treatment (odds ratio, 2.45; 95% CI, 0.78 to 7.71; p=0.12, I²=69%), but different forms and routes of FMT administration were shown to be equally efficacious. In conclusion, FMT is a promising treatment modality for recurrent CDI. However, the authors found limited data for the variability of FMT dose usages, small trial populations and time frames to assess the success or failure of treatment.

In 2019, a third meta-analysis (Tariq et al) evaluated the efficacy of FMT as a treatment option for recurrent CDI on the basis of results from open-label studies and placebo controlled clinical trials. The authors wanted to investigate their observations on FMT cure rates for CDI being high in observational studies (e.g., >90%) but then appear to be consistently lower in open-label studies and clinical trials. Thirteen studies were included for evaluation, including six placebo-controlled RCTs and seven open-label studies. Out of 610 patients receiving FMT, 439 patients achieved clinical cure (76.1%; 95% confidence interval [CI]: 66.4% to 85.7%); study heterogeneity was significant (I² =91.35%). Cure rates were found to be lower in randomized trials (139/216, 67.7%; 95% CI: 54.2% to 81.3%) vs open-label studies (300/394, 82.7%; 95% CI: 71.1% to 94.3%; p < 0.001). Subgroup meta-analysis by FMT route of administration indicated lower cure rates with enema than colonoscopy (66.3% vs 87.4%; p < 0.001). However, no differences between colonoscopy and oral delivery were detected (87.4% to 81.4%; p= 0.17). Lower cure rates were observed for studies that included both recurrent and refractory CDI than those that only included patients with recurrent CDI (63.9% vs 79%; p < 0.001). The authors concluded that colonoscopy and oral route are more effective than enema for stool delivery and the efficacy seems to be higher for recurrent than for refractory CDI.

A recent UpToDate (2020) review summarizes FMT as an effective treatment for recurrent (≥3 recurrences) CDI. The authors concluded that transplantation of stool microbiota from healthy individuals to patients with recurrent CDI can break the cycle of CDI recurrence. In addition, the initial results from the FMT National Registry published in Gastroenterology 2021 showed a 1 month cure rate of over 90% in 200 patients who received just 1 FMT. At a 6 month follow-up, only 4% of patient had a recurrence of CDI and less than 1% had a complication of treatment includes IBS or newly diagnosed IBD. However, to minimize risk of infection, rigorous screening of potential healthy stool donors for occult pathogens must be done. The optimal approach for administration is uncertain. If feasible oral capsules should be tried first, then colonoscopy followed by enema retention and finally NJ or ND tube for patients who cannot undergo FMT via the alternate routes.
In 2016, the U.S. Food and Drug Administration (FDA) issued an updated draft guidance on enforcement policy regarding investigational new drug requirements for use of FMT to treat CDI not responsive to standard therapies. The draft guidance is similar to the 2013 guidance and states that the FDA is continuing to consider how to regulate FMT and that, during this interim period, the agency will use enforcement discretion regarding the use of fecal transplant to treat treatment-resistant CDI. The FDA requires that physicians obtain adequate informed consent from patients or their legal representative before performing the intervention. The document also stated that selective enforcement does not apply to the use of fecal transplant for treating conditions other than treatment-resistant CDI.

In 2019, the FDA issued a safety regarding the use of FMT due to the potential risk of serious adverse reactions or life-threatening infections caused by due to the transmission of multi-drug resistant organisms (MDROs). Two immunocompromised individuals received investigational FMT and developed invasive infections caused by the transmission of extended-spectrum beta-lactamase-producing Escherichia coli. One of the affected individuals died. The donor stool used in each patient's FMT procedures had not been tested for extended-spectrum beta-lactamase (ESBL)-producing gram-negative organisms prior to use. Follow-up testing verified donor stool was positive for MDROs identical to the organisms isolated from the two patients. Due to these events, the FDA has determined that the following additional protections are required for any investigational use of FMT:

1) Donor screening that specifically addresses risk factors for colonization with MDROs and exclusion of individuals at higher risk of colonization with MDROs such as health care workers, persons who have recently been hospitalized or discharged from long-term care facilities, persons who regularly attend outpatient medical or surgical clinics, and persons who have recently engaged in medical tourism.

2) MDRO testing of donor stool and exclusion of stool testing positive for MDROs. At the minimum, tests should include: extended-spectrum beta-lactamase-producing enterobacteriaceae, vancomycin-resistant enterococci, carbapenem-resistant enterobacteriaceae and methicillin-resistant Staphylococcus aureus.

3) All FMT products currently in storage for future use must be quarantined until donor MDRO carriage risk can be assessed and FMT products are tested and found negative for MDROs.

4) The informed consent process for FMT treatment subjects should describe the risk of MDRO transmission and infection and the measures being implemented for donor screening and stool testing.

Lastly, the American College of Gastroenterology (2021) published guidelines on diagnosis, treatment, and prevention of CDI. The guidelines addressed fecal microbiota transplant for treatment of three or more CDI recurrences, as follows: "For the treatment of one to two CDI recurrences, the guidelines recommend: tapering/pulsed-dose vancomycin for patients experiencing a first recurrence after an initial course of fidaxomicin, vancomycin, or metronidazole (strong recommendation, very low quality of evidence). In addition, Fidaxomicin for patients experiencing a first recurrence after an initial course of vancomycin or metronidazole. FMT be considered for patients with severe and fulminant CDI refractory to antibiotic therapy, in particular, when patients are deemed poor surgical candidates."

The Infectious Diseases Society of America and Society for Healthcare Epidemiology of America updated their clinical practice guidelines in 2018 for the diagnosis and treatment of CDI in children and adults. For pediatric patients fecal microbiota transplantation may be used after multiple recurrences of CDI following standard antibiotic treatments, this is considered a weak recommendation with very low quality of evidence. In adult patients, fecal microbiota transplantation is strongly recommended for patients with multiple recurrences of CDI who have failed appropriate antibiotic treatments as there is moderate quality of evidence.
No recommendations were found from the U.S. Preventive Services Task Force.

**Applicable Coding**

**CPT Codes**

**44705** Preparation of fecal microbiota for instillation, including assessment of donor specimen

**HCPCS Codes**

**G0455** Preparation with instillation of fecal microbiota by any method, including assessment of donor specimen

**ICD-10 Codes**

**A04.7** Enterocolitis due to Clostridium difficile

**A04.71** Enterocolitis due to Clostridium difficile, recurrent

**A04.72** Enterocolitis due to Clostridium difficile, not specified as recurrent

**References:**

3. Costello SP, Hughes PA, Waters O et al. Effect of Fecal Microbiota Transplantation on 8-Week Remission in Patients With Ulcerative Colitis: A Randomized Clinical Trial. JAMA, 2019 Jan 16;321(2). PMID 30644982


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