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Multi-Donor Fecal Microbial Transplantation for Critically Ill Patients: Rationale and Standard Operating Procedure

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Abstract: Patients in the intensive care unit often lose a considerable fraction of their gut microbiome due to exposure to broad-spectrum antibiotics and other reasons. Dysbiosis often results in prolonged diarrhea and increase occurrence of multi-drug resistant pathogens in the colon with clinical consequences not yet well understood. Restoring the microbiome by fecal microbial transplantation (FMT) is a plausible therapeutic possibility, so far only documented in case reports and case series using very heterogeneous methodologies. Before FMT with critically ill patients can be tested in randomized controlled trials, there is a burning need to describe a standardized operating procedure (SOP) of the whole process, respecting the specifics of the critically ill population, such as the risk of the disrupted intestinal barrier and time-critical nature of the procedure. We describe the SOP that has been developed for experimental use in critically ill patients by a multidisciplinary team of intensivists, gastroenterologists, and microbiologists based on feedback from regulatory authority (State Institute for Drug Control of the Czech Republic). The hallmarks of these SOPs are multi-donor freshly frozen transplants guaranteed for 2 months consisting of seven aliquots from seven unrelated healthy donors and administered by a rectal tube. In this paper we discuss the rationale for this SOP and the process of its development in detail and release the full proposed SOP is in the form of an online appendix.

Keywords: fecal microbial transplantation; critically ill patients; standard operating procedure; diarrhea

1. Introduction

Although the sayings on the connection between the proper function of the gastrointestinal tract (GIT) and longevity have been known for decades, a full scientific description of such claims based on scientific evidence is yet to be made. Lumen inhabiting microorganisms (termed “microbiota”) together with their metabolic products in the specific host environment (termed “microbiome”) are indispensable for gastrointestinal physiology. The individual specifics of microbiome composition are determined by multiple factors, where genetics, early exposure to microorganisms (mainly transport from mother during birth), environment, and diet are considered the dominant ones [1,2]. Due to the microbiome’s vast complexity, it is being considered as a separate organ by some researchers [3]. The recent progress in 16S rRNA sequencing brought about proof that the microbiome composition of each individual is unique to such an extent, that it may be used in forensic disciplines [4,5]. This enormous microbiome diversity is pivotal to maintaining homeostasis [6,7]. In turn,
a reduction of microbiome diversity (termed “dysbiosis”) results in immune system dysfunc-
tion [8,9]. The association between an altered microbiome and diseases has already
been confirmed for a range of gastrointestinal tract diseases (e.g., inflammatory bowel dis-
ease [10], irritable bowel syndrome—IBS [11], colorectal carcinoma [12]), gastric carcinoma,
and liver disease [13]) as well as for metabolic syndrome [13], gout [14,15], neurodegener-
ative [16], autoimmune (rheumatoid arthritis, ankylosing spondylitis, lupus, sclerosis
multiplex [14,15]) or psychiatric diseases [17], including depression [18].

Although the complex nature of these associations is yet to be understood, it is likely
that two-way communication between microbiota and immune system exists [19] and is me-
diated by microbial-synthesized metabolites regulating host neuroimmune-inflammatory
axes that could physiologically link gut with other organ systems, such as the brain [20] or
liver [21]. Details of the interplay are summarized in a paper by Cibulkova et al. [22].

2. Dysbiosis and Diarrhea in Intensive Care Patients

A recent study in patients admitted to intensive care showed, that 90% of intestinal
microflora is lost during the first 6 h of ICU stay [23]. The most common symptom of
dysbiosis is diarrhea, which may not only complicate ICU stay by threatening skin surfaces,
but also alter dignity and worsen the prognosis of ICU patients [24,25]. The main risk
factors of diarrhea in ICU patients are antibiotics therapy, enteral nutrition [26], proton
pump inhibitors (PPI) therapy [27], the severity of underlying disease [28], and history
of recent surgery [29]. As approximately 70 % of ICU patients worldwide are treated
with antibiotics [30] and 5–25 % of those develop diarrhea [31], antibiotics are indeed the
leading cause of diarrhea in ICU. On the other hand, the risk of diarrhea and its negative
consequences are among the drivers of antibiotic stewardship programs [24,25]. In addition
to this, the adoption of routine ICU care bundles has been suggested to reduce dysbiosis
and diarrhea [32]. Dysbiosis is also linked to changes in intestinal barrier permeability
and possible immune dysregulation and organ failures [33,34] and also increased rate of
colonization with multi-drug resistant pathogens.

Under normal conditions, nutrients introduced into the digestive system stimulate the
renewal of the mucosa in the digestive tract by acting on it directly, as well as by releasing
gastrointestinal hormones, such as gastrin [35], cholecystokinin [36] and ghrelin [37]. These
protective mechanisms of enteral feeding might be disrupted in some critically ill patients,
due to a decrease in mucosal blood flow due to splanchnic ischemia [38,39]. In addition,
small bowel hypoxia leads to gut barrier failure associated with bacterial translocation,
 systemic inflammation and the development of multiple organ dysfunction syndromes [40].

3. Fecal Microbial Transplantation: Defining the Needs of Standard Operating
Procedure

Fecal microbial transplantation (FMT sometimes termed “bacteriotherapy”) is the
transfer of stool preparation from a healthy donor (or donors) into the gastrointestinal tract
of the recipient to treat or prevent disease [41]. The procedure aims to restore a healthy and
diverse microbiome. Animal studies indicate that enteral nutrition or probiotics alone are
not sufficient to restore commensal colonization of intestinal crypts [42,43]. Apart from
induction the healthy and diverse bacteria [44], FMT also introduces dead bacteria, viruses,
fungi, microbial peptides, and metabolites, as well as intact desquamated donor colonocytes,
which all contribute to FMT effects [45–47]. FMT triggers the release of antimicrobial factors,
modulates the proliferation of immune cells, stimulates mucus formation and immune
globulin A synthesis, inhibits the activation of nuclear factor Kappa B in inflammatory
epithelial cells, and increases mucin secretion, which in turn may restore the epithelial
barrier and intestinal function.

FMT is not a new method, its first use was documented in China in the 4th-century AD,
where Ge Hong used a human stool to treat food poisoning or severe diarrhea [43]. In World
War II, German soldiers were eating camel excrements as a remedy for bacterial dysentery
during the Battle of Africa. In standard Western medicine, FMT was first used in 1958 as a
therapy for pseudomembranous colitis. The method gradually became a well-established treatment for recurrent and refractory *Clostridioides difficile* infection (CDI) [45]. Along with the buildup of epidemiological evidence of an association between dysbiosis and a range of diseases, FMT has been considered a potential treatment beyond recurrent CDI [46]. Clinical trials have been published for the use of FMT in inflammatory bowel disease [47,48], and primary infection with *Clostridioides difficile* [49], and research is underway for Parkinson’s disease [50], amyotrophic lateral sclerosis (NCT03766321, [51]), and Alzheimer disease (NCT03998423). Given the rapid onset of dysbiosis in the ICU patients and probable links with diarrhea, immune dysregulation, intestinal barrier disruption, multi-organ failure and MDR infections, early restoration of the microbiome through FMT may bring substantial benefit to ICU patients [52]. On the other hand, introducing a surge of living microbes into the intestine of a very sick patient with dysregulated immunity and altered intestinal epithelial barrier function can bring about significant risks, mainly of bacterial translocation, sepsis, or worsening systemic inflammation—in addition to other known risks of FMT in noncritically ill population, including the introduction of multi-drug resistant organisms (MDRO) from donor to the recipient and death [53]. To date, only a case report and small case series have been published [52,54–56], and albeit their unequivocally positive results in treating diarrhea and improving overall patients’ conditions are encouraging, these studies may be subjected to publication and attribution biases. There is a burning need for robust prospective, randomized controlled trials to evaluate the FMT safety and efficiency in this vulnerable patient’s population.

Indeed, performing high-quality studies compliant with good clinical practice principles requires the approval of the respective regulatory authorities. These may feel uneasy to approve and oversee trials on FMT, particularly in vulnerable populations, such as critically ill patients, e.g., due to the difficulties of standardization of the investigational product. In addition, logistical challenges, and the feasibility of FMT in a specific environment of ICU must be considered, too. For example, the State Institute for Drug Control (Statní ústav pro kontrolu léčiv—SUKL) of the Czech Republic decided in 2019 to consider FMT transplant a drug and decided to apply the same principles of regulation and oversight as is in place for drug trials. Consequently, standard operating procedures (SOPs) are required for donor selection, transplant preparation, and recipient monitoring, ensuring the safety and reproducibility of FMT in clinical trials. In collaboration with the regulatory authority of the Czech Republic, SUKL, we have developed such an SOP, which we describe and explain below. Although directly applicable in only one EU country and only in the context of a clinical trial, we believe that it can be useful in its current form or after minor modification in other countries too and help to standardize FMT procedure in clinical trials in critically ill patients.

4. Standard Operating Procedure of Fecal Microbial Transplantation in Intensive Care Unit Patients

The full text of SOPs can be found in Supplementary Materials. There we describe the development of a standard operating procedure for multi-donor fecal microbial transplantation in critically ill patients, starting from donor selection and finishing with post-procedure recipient care. In summary, our SOP recommends the use of a multi-donor transplant consisting of seven 50 mL aliquots from seven donors. The aliquots are prepared from healthy unrelated volunteers, who regularly donate. In addition to standard guidelines-driven [57–59] precautions and tests (See Supplementary Digital Content) in donors to prevent transmissible infection, the deep-frozen transplant mixed with glycerol is quarantined for at least 2 months before use. The transplantation itself is performed by a semi-rigid rectal irrigation tube in a patient positioned to the left semi lateral Trendelenburg’s position for 15 min and then in the right semi lateral Trendelenburg’s position for another 15 min. This ensures the distribution of the transplant well throughout the length of the colon (Figure 1). This recommendation has been developed as a modification of existing guidelines for the use
of FMT to eradicate *Clostridioides difficile* in other than intensive care settings [60,61]. All modifications and their justifications are described below.

![Abdominal X-ray showing the spread of 350 mL transplant mixed with 5 mL of contrast agent 15 min after administration to a sedated ventilated patient via semirigid rectal tube. Fecal derivation system (Flexiseal©) has been clamped immediately after administration with the rectal balloon inflated being clearly visible on the image. Note: The addition of a contrast agent is normally not a part of FMT and it should be noticed that there are no robust data on how FMT without contrast agent spreads after rectal administration.](image-url)

**Figure 1.** Abdominal X-ray showing the spread of 350 mL transplant mixed with 5 mL of contrast agent 15 min after administration to a sedated ventilated patient via semirigid rectal tube. Fecal derivation system (Flexiseal©) has been clamped immediately after administration with the rectal balloon inflated being clearly visible on the image. Note: The addition of a contrast agent is normally not a part of FMT and it should be noticed that there are no robust data on how FMT without contrast agent spreads after rectal administration.
4.1. Donor Selection and Rationale for Multi-Donor Strategy

Appropriate donor selection is the key condition determining FMT success [57,62]. The success rate is generally greater in FMT from a family member compared to an unrelated donor (93% vs. 84%) [63] or from so-called “superdonors” with known high microbiome diversity [64]. It seems that the success of the FMT procedure depends on the interaction (or rather compatibility) between the recipient’s and donor’s microbiome [62,65]. In theory, an ideal donor should have a microbiome as similar as possible to the premorbid microbiome of the recipient [66], or at least diverse enough that the mix certainly contains bacteria that the recipient will accept [60,62]. This idea led to concepts of fecal autotransplantation [67], a superdonor [64] or a multi-donor transplantation [68,69]. Multi-donor means mixing frozen-thawed small aliquots of stool from multiple donors into one graft for one recipient. We have adopted a multi-donor strategy for the use in the critically ill for the following reasons:

- Frozen samples are readily available and allow the best timing for the patient and staff regardless of donors’ defecation habits. No need to exclude a large proportion of critically ill patients without relatives eligible for and willing to donate stool.
- Increase microbiome diversity as compared to a single donor.
- Possibility of some degree of standardization, because aliquots from the same donors yield grafts of very similar composition.
- Cost-effectiveness and affordability as compared to commercial preparations from super donons.

As a downside, the multi-donor strategy increases the risk of transmission of infectious diseases that have escaped thorough screening [57]. It should be stressed that the ultimate responsibility for donor acceptance or rejection is with the physician examining them. The battery of tests represents the mandatory minimum, which can be extended if needed. Yet there always will be infections that are potentially transferable and are not investigated and the risk of those should be mentioned in the informed consent. The 2-months graft quarantine with a repeated examination of the donor aims to minimize this risk. The number of aliquots from seven donors has been determined by the convenience of practical 50 mL size aliquots and the required amount of transplant (350 mL, see below).

4.2. Donor Stool Processing and Sample Preparation

Notably, among numerous guidelines and protocols for FMT, there is large heterogeneity in the process of sample preparation and often these important details are not reported at all. We use 1:3 \( w/w \) dilution with normal saline before homogenization and double layer gauze filtration as this provides optimal viscosity and the use of normal saline provides a better outcome compared to sterile water [63]. Similarly, we have chosen a 350 mL volume as a compromise between 500 mL that demonstrated a 97% success rate, and 200 mL, which was successful in 80% of cases [57,63]. Inline, using 200 mL could compromise the efficacy of the procedure and larger volumes than 350 mL would require either using more donors or aliquots larger than 50 mL, which is the maximum volume that can be filled into a standard Jeanette syringe. The use of frozen transplants has been chosen for the practical and safety reasons discussed above. It seems that <12 months of freezing does not significantly alter FMT efficacy [53], but further durability studies were requested by the SUKL and are underway. In addition, we are now developing a completely closed system of sample preparation in a paddle blender ("Stomacher") using a sealable plastic bag with an infusion port and customized nylon filter.

4.3. Southern Way of Administration

With regards to the way of administration, various methods were considered. The lower GI or “Southern” route (via colonoscopy or retention enema) has been found more efficient in microbiome restoration compared to the upper GI “Northern” routes (esophagogastroduodenoscopy, nasogastric, nasojejunal tube, poop pills) [70]. Additionally, we considered upper GI administration of fecal material into the small intestine with a po-
tentially disrupted epithelial barrier too risky. Of particular concern, was aspiration, a recognized complication of the upper GI route [71]. Moreover, in a survey, the majority of ICU nurses considered the administration of fecal material into the colon more acceptable for them as opposed to nasogastric administration (Iva Havirova thesis, unpublished). Given the logistic issues with the availability of endoscopists and considerable risks of endoscopy [72], we have chosen administration by an enema, despite its lower efficiency, compared to colonoscopy. With appropriate patient positioning, we found graft distribution throughout the colon acceptable, reaching the ascendent colon (Figure 1). Indeed, this case-based anecdotal data are far from conclusive with regards to FMT spread and effect size. To maximize engraftment, we recommend performing FMT no sooner than 48 h after the last dose of antibiotics. An important objective of the upcoming trials should not only be to provide more data on the distribution of the graft in the recipient’s colon, but also to test the efficacy of donor microbiome engraftment.

4.4. Safety Considerations and Measures to Maximise Safety for Recipients

Critically ill patients represent a very specific and vulnerable patient population, in which a normally very safe procedure [40] with only mild and self-limiting safe effects [73] can have additional serious risks that might outweigh benefits. In theory, the bacterial load could lead to sepsis and worsen organ failure. This is why we recommend thorough patient investigation and monitoring before, during, and after the procedure, on top of the safety precautions described above. So far published case reports and case series do not report any serious adverse and only one death in 31 patients (See Table 3 of Cibulkova et al. [22]), but this can be heavily biased as only successful procedures may have been selected for publication. Until more data is gained from randomized controlled trials, we only recommend lower GI administration of FMT to critically ill patients either for established indications in non-critically ill populations (such as in C. dif. infection) or limiting the study population for new indications (e.g., antibiotic-associated diarrhea) to hemodynamically stable patients, that do not have perforated viscus, recent colorectal surgery (less than 3 months) or severe neutropenia (less than 500 per microliter). In addition to continuous monitoring of vital functions, which is mandatory in the ICU, we recommend routine blood cultures be taken 3 h post-FMT plus whenever there are signs of an inflammatory response (such as rigors or fever). Only when initial clinical trials yield more safety data, do we recommend expanding research further. This SOP aims to make FMT in critically ill patients as safe as possible, but it should be stressed that good quality data on FMT safety in critically ill patients are still not available.

5. Conclusions

Critically ill patients often suffer from rapid-onset intestinal dysbiosis resulting in prolonged diarrhea and potentially other adverse consequences. Restoration of the physiological microbiome through FMT has been proven safe and effective in a range of conditions outside the intensive care unit. Despite biological plausibility and strong collateral evidence, there is no data on the safety and efficacy of FMT to treat dysbiosis-related conditions in critically ill patients. We have developed a standard operating procedure for the use of FMT in critically ill patients. The hallmark of the methodology is the use of frozen multi-donor transplant quarantined for 2 months and administered by an enema to carefully monitored recipients. It should be stressed that FMT in critically ill patients should only be used in the context of randomized controlled trials. A multi-disciplinary approach is warranted to the design and conduct of clinical trials in this exciting field.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/futurepharmacol2010005/s1.

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