

Microbiome-based companion diagnostics: no longer science fiction?

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In the last decade, our knowledge on the role of the gut microbiome in health and disease has greatly increased, accompanied by an unseen hype around both its diagnostic and therapeutic potential. Yet, one application area of the microbiome has thus far remained understudied: its role as guidance for therapeutic decisions, treatment monitoring and prediction of response.

In this issue of *Gut*, Mondot *et al*¹ investigate to what extent the microbiome can both inform us on as well as contribute to treatment outcome in resection surgery for Crohn's disease (CD). Although often the only resort next to dilatation,² this intervention is not curative, with first-year endoscopic recurrence rates at 28–93% and frequent necessity of reoperation.³ As such, there is an important need for effective postoperative maintenance strategies and predictions for postoperative outcome.

In a longitudinal study of both the faecal and mucosa-adherent microbiota in 20 patients undergoing resection surgery, Mondot *et al* describe the microbial groups recolonising the intestinal lining around the anastomosis and show different trajectories in recurring patients *versus* those in remission; the latter were found to exhibit a stable, cohesive community structure, which is spatially homogeneous before and after the anastomosis. The species associated with recurrence could form companion drug targets for this procedure, although their causal role still needs to be established. A particularly intriguing aspect of this study is the identification of multiple operational taxonomic units whose presence before surgery seems predictive of early recurrence. Signatures like these open up the exciting prospect of a quick companion diagnostic that would allow a preoperative evaluation whether the patient is likely to remain in long-term remission after treatment. Of course, the current study is

underpowered; larger scale validation in multiple centres will be needed before such a companion diagnostic becomes reality, but conceptually this is a very attractive model.

Thus far, similar studies are scarce, but equally promising. For example, Rajca *et al*⁴ showed that low rates of *Faecalibacterium prausnitzii* and *Bacteroides* are predictors of relapse after infliximab withdrawal in CD. Machiels *et al*⁵ found the presence of *Ruminococcus gnavus*, *Bacteroides vulgatus* and *Clostridium perfringens* and the absence of *Blautia* and *Roseburia* in faecal samples of patients with UC before surgery to be associated with a higher risk of pouchitis after ileal pouch-anal anastomosis. Kaddurah-Daouk *et al*⁶ identified two secondary, bacterial-derived bile acids in a metabolomics screen that contributed to predicting the magnitude of statin-induced low-density lipoprotein cholesterol lowering in responders. Vétizou *et al*⁷ show that the success of ipilimumab anticancer treatment is even causally dependent on the presence of *Bacteroidales*. Together, these studies suggest that the role of the microbiome in clinical guidance is relevant in a wide range of conditions and interventions.

Perhaps, the most likely therapeutic area where microbiome-based treatment guidance is that where the microbiota is part of the treatment itself: faecal microbiota therapy (FMT), also known as faecal transfer. FMT has been extremely effective in *Clostridium difficile* infections, yet for other pathologies success rates are more limited or unclear, and microbiome-informed treatment steering might be a solution. One such area is that of UC. Studies seem to agree upon a ± 25 –30%

success rate.^{8–10} Yet, as the placebo efficacy was of a similar order of magnitude in some studies,⁹ the initial enthusiasm was somewhat curbed. To improve upon these numbers, microbiome-based treatment guidance could be an option. As a point in case, we recently found that donor richness and the number of transferred phylotypes were associated with sustained remission and found indications that FMT success was associated with the successful transfer of *Roseburia* and *Oscillibacter*.⁸ Although still early days, this does suggest that microbiome-based patient and donor selection, with the latter ultimately replaced by construction of personalised probiotic cocktails, is likely to benefit FMT outcome in UC, and possibly in other pathologies.

Overall, these studies indicate that the microbiome field is slowly but surely approaching the clinic. I believe the future role of microbiome monitoring in daily medical practice can be found at four different levels (figure 1). First, microbiome markers can be used for diagnosis (and potentially prognosis) of disease. Second, analysis of patient microbiota could predict the outcome of treatment options. Third, based on the patient's microbiome, a personalised treatment strategy can be devised, be it based on the administrations of specific microbial cocktails ('precision probiotics'), targeted microbial nutrients ('precision prebiotics'), personalised dietary interventions or targeted antibiotics and phages. Finally, treatment success and establishment of normobiosis can be monitored using microbiome time-series analysis.¹¹ The thrilling fact that multiple aspects of this microbiome-based therapeutic model are nearing clinical implementation reflects how the field is shaking off its 'growing pains' and is increasingly becoming a true translational discipline.

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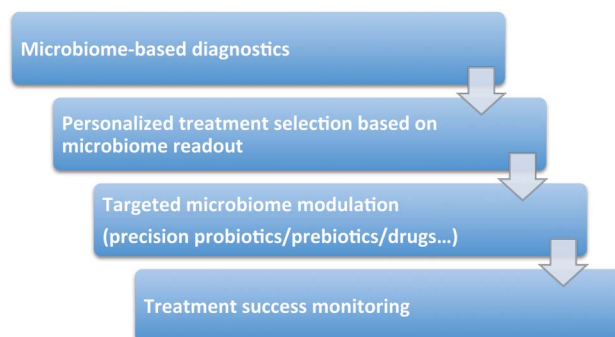


Figure 1 Microbiome-based therapeutic model.

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