Multiple Sclerosis Patients have an Altered Gut Mycobiome and Increased Fungal to Bacterial Richness

Running Title: RRMS Patients have Gut Fungal Dysbiosis

Meeta Yadav¹,²,#, Soham Ali³,#, Rachel L. Shrode⁴, Shailesh K. Shahi¹, Samantha N. Jensen¹,⁵, Jemmie Hoang⁶, Samuel Cassidy⁸, Heena Olalde⁸, Natalya Guseva¹, Mishelle Paullus⁸, Catherine Cherwin⁶, Kai Wang⁷, Tracey Cho⁸, John Kamholz⁸, Ashutosh K. Mangalam¹,⁴,⁵,⁹*

¹Department of Pathology, Carver College of Medicine, University of Iowa, Iowa City, IA, USA, ²University of Iowa College of Dentistry, Iowa City, IA, USA, ³Carver College of Medicine, University of Iowa, Iowa City, IA, USA, ⁴Informatics Graduate Program, University of Iowa, Iowa City, IA, USA, ⁵Interdisciplinary Graduate Program in Immunology, University of Iowa, Iowa City, IA, USA, ⁶College of Nursing University of Iowa, Iowa City, IA, USA, ⁷Department of Biostatistics, College of Public Health, University of Iowa, Iowa City, IA, USA, ⁸Department of Neurology, University of Iowa Hospitals and Clinics, Iowa City, IA, USA, and ⁹Iowa City VA Health System, Iowa City, IA, USA.

* Corresponding author:

Dr. Ashutosh K Mangalam
Department of Pathology,
University of Iowa Carver College of Medicine
25 S. Grand Avenue, 1080 ML
Iowa City, IA 52242, USA
Phone: 1-319-335-8558
Email: Ashutosh-mangalam@uiowa.edu

# Authors contributed equally
Abstract

Trillions of microbes such as bacteria, fungi, and viruses exist in the healthy human gut microbiome. Although gut bacterial dysbiosis has been extensively studied in multiple sclerosis (MS), the significance of the fungal microbiome (mycobiome) is an understudied and neglected part of the intestinal microbiome in MS. The aim of this study was to characterize the gut mycobiome of patients with relapsing-remitting multiple sclerosis (RRMS), compare it to healthy controls, and examine its association with changes in the bacterial microbiome. We characterized and compared the mycobiome of 20 RRMS patients and 33 healthy controls (HC) using Internal Transcribed Spacer 2 (ITS2) and compared mycobiome interactions with the bacterial microbiome using 16S rRNA sequencing. Our results demonstrate an altered mycobiome in RRMS patients compared with HC. RRMS patients showed an increased abundance of Basidiomycota and decreased Ascomycota at the phylum level with an increased abundance of Candida and Epicoccum genera along with a decreased abundance of Saccharomyces compared to HC. We also observed an increased ITS2/16S ratio, altered fungal and bacterial associations, and altered fungal functional profiles in MS patients compared to HC.

This study demonstrates that RRMS patients had a distinct mycobiome with associated changes in the bacterial microbiome compared to HC. There is an increased fungal to bacterial ratio as well as more diverse fungal-bacterial interactions in RRMS patients compared to HC. Our study is the first step towards future studies in
delineating the mechanisms through which the fungal microbiome can influence MS disease.

Introduction

Multiple Sclerosis (MS) is a neuroinflammatory autoimmune disease that affects ~2.5 million people worldwide. Though MS has different clinical subtypes, most of the patients (~85%) present with relapsing-remitting MS (RRMS). The precise etiopathogenesis of MS is unclear, but both genetic and environmental factors have been suggested to play an important role in susceptibility and the pathogenesis of MS. In recent years multiple groups, including ours, have highlighted the role of gut bacteria in the pathobiology of MS\textsuperscript{1-8}, and it has emerged as an important environmental factor. However, the gut microbiome consists of non-bacterial microbes such as fungi, viruses, and archaea\textsuperscript{9, 10}, yet the role of the fungal microbiome (mycobiome) is not well studied in MS.

Although fungi make up approximately 0.1% of the gastrointestinal tract\textsuperscript{10, 11}, their impact on human health is significant due to their ability to regulate local and systemic host immune response\textsuperscript{12, 13}. The importance of the fungal microbiome on the immune system was demonstrated in a study where mice that were rewilded had increased intestinal abundance of fungi along with an expansion of granulocytes compared to laboratory mouse controls\textsuperscript{12}. While in human peripheral blood 50-70% cells are neutrophils, only 10-25% cells are neutrophils in mouse peripheral blood\textsuperscript{14}. Additionally, an oral antifungal treatment of mice in colitis model showed increased disease severity and exacerbated allergic airway disease with increased \textit{Aspergillus}, \textit{Wallemia}, and \textit{Epicoccum} and decreased \textit{Candia} spp. Moreover, oral administration of a mixture of these three fungi (\textit{Aspergillus amstelodami}, \textit{Epicoccum nigrum}, and
Wallemia sebi) was sufficient to recapitulate the exacerbating effects of antifungal drugs on allergic airway disease. The importance of the mycobiome in human health and disease is validated by studies showing alterations in mycobiome composition in various autoimmune diseases such as inflammatory bowel disorders (IBD), ankylosing spondylitis (AS), and type 2 diabetes mellitus (DM) compared to healthy controls (HC). These studies have also shown correlations between bacterial and fungal microbiota that are altered in disease states. The relationship between gut bacteria and fungi has been demonstrated in the murine model where bacteria outnumbered fungi by greater than three times prior to antibiotic administration, but after antibiotic administration bacterial abundance dropped three-fold while fungal abundance increased around 40-fold. Thus all these observations suggest that gut mycobiome might play an important role in the health and disease including MS.

Based on the association of gut mycobiota in other autoimmune diseases and on the described relationship between gut bacteria and MS, we hypothesized that the gut fungal microbiota might be altered in MS patients. Therefore, in this study we profiled the fungal microbiota and associated fungal functional characteristics in RRMS patients (MS) (n=20) and compared them with those of HC (n=33). We also analyzed the bacterial microbiome of the same patients to determine the correlation between the fungal and bacterial microbiome. We observed that the MS patients have a distinct fungal microbiome compared to HC, with differential abundances of multiple fungal genera and fungal functions as well as altered fungal-bacterial relationships. A distinct fungal microbiome in RRMS patients could have implications in the pathogenesis of MS, potentially through functional differences and altered interactions with the bacterial microbiome.
Methods

Standard Protocol Approvals, Registrations, and Patient Consents

The study was done in accordance with the guidelines approved by the University of Iowa Institutional Review Board (IRB). A prior written informed consent was obtained from all the subjects to participate in the study.

Human Subjects Enrollment

Relapsing-Remitting Multiple Sclerosis (RRMS) patients (n=22) who fulfill McDonald diagnostic criteria for MS were recruited from the Neuroimmunology Clinic at the University of Iowa Hospital Clinics (UIHC), at the University of Iowa. Healthy controls (HC) (n=34) were also recruited through the University of Iowa (Table 1). Individuals eligible for the study were those 18-63 years of age with a diagnosis of RRMS. All patients were either on MS treatment or not on treatment for at least three months prior to enrollment. Patients and healthy controls were asked to provide samples at least four weeks after their last dose of oral antibiotics or laxatives, and at least three months after their last colonoscopy. Those with active pregnancy or a history of bariatric surgery were excluded from the study.

Table 1. Biometric and MS Patient Treatment Data

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<th></th>
<th>HC</th>
<th>MS</th>
<th>p-value</th>
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<tr>
<td><strong>Age: Mean ± SD</strong></td>
<td>42 ± 14</td>
<td>45 ± 7.8</td>
<td>t-test: 0.33</td>
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<tr>
<td><strong>Sex: Male / Female</strong></td>
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<td>5 / 15</td>
<td>Fisher: 0.48</td>
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<tr>
<td><strong>BMI: Mean ± SD</strong></td>
<td>24 ± 3.7</td>
<td>30 ± 7.9</td>
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</tr>
<tr>
<td>Treatment: Yes / No</td>
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<td>16 / 4</td>
<td>NA</td>
</tr>
<tr>
<td>---------------------</td>
<td>----</td>
<td>--------</td>
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</tr>
<tr>
<td>Treatment</td>
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<tr>
<td>Glatiramer acetate</td>
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<tr>
<td>Dimethyl fumarate</td>
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<td>6</td>
<td></td>
</tr>
<tr>
<td>Not on treatment</td>
<td></td>
<td>4</td>
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</tbody>
</table>

**Human Sample Collection**

Stool samples were collected by patients and HC in Commode Specimen Collection kits (Fisher PA, USA) provided to them by our laboratory along with the instruction sheet and shipped on frozen gel packs to our laboratory by overnight delivery. Once in the laboratory, the stool was aliquoted and stored at -80 degrees until DNA extraction. The same collection kits, instruction sheets, and processing methods were used for fecal samples from MS patients and HC.

**DNA Extraction and Sequencing**

Microbial DNA was extracted from each fecal sample using Qiagen DNeasy PowerLyser PowerSoil Kit (Qiagen, Germantown, MD) per the manufacturer’s instructions using a bead-beating step (PowerLyzer 24 Homogenizer, Omni International, USA). The fungal microbiome was analyzed using Two-step PCR as described previously for bacterial microbiome analysis except instead of 16S rRNA; we targeted the internal transcribed spacer 2 (ITS2) region of fungal 18S rRNA using
the following primers: forward primer 86 F(5'-3') GTGAATCATCGAATCTTTGAA and reverse primer 4R (5'-3') TCCTCCGCTTATTGATATGC20. Fungal PCR conditions were as follows: 95°C for 5 min, 35 cycles of 95°C for 30s, 59°C for 30s, 72°C for 30s, and 72°C for 10 min. Bacterial 16S rRNA sequencing was performed as per protocol published by our lab19.

**Metagenomic Profiling**

The raw fungal ITS2 (internal transcribed spacer 2) sequence data was processed utilizing DADA221. In brief, the reads were trimmed to remove the primer sequences, then truncated based on a quality score of 25. Reads were then denoised to infer the exact sequence variants by resolving single-nucleotide errors. Paired forward and reverse reads were merged, and any resulting chimeras were removed to produce amplicon sequence variants (ASVs). These ASVs were then assigned fungal taxonomy at the genus level using a naive Bayesian classifier method and the UNITE database (Version 8.2)22. After the removal of three samples (2 MS and one HC) with less than 1000 reads, 20 MS samples and 33 HC samples were available for fungal profiling, with a median number of reads of 37,331.

The raw sequence data of the V3-V4 region of bacterial 16S rRNA was processed utilizing DADA2. These sequences were processed as described above for ITS sequences, and the final ASVs were assigned bacterial taxonomy at the kingdom to species levels using the Silva database (version 138.1)23 with a median number of reads of 49,976.

**Functional Profiling**

The functional profile of the gut mycobiome was generated using FungalTraits
database. Of the 142 identified fungal genera in this study, 86 had functional entries in this database, and the functional profile of each sample was then inferred using its fungal composition and the functional data of each of these 86 fungal genera.

**Statistical Analysis**

Analysis and figure generation of alpha diversity, beta diversity, and differential abundance of all features was performed entirely in R (Version 4.0.3) using the vegan and ggpubr packages with custom R scripts. Prior to statistical analysis, the fungal and bacterial datasets were each constant-sum scaled to one million reads and generalized log-transformed. The fungal and bacterial data were then filtered to remove low prevalence taxa. Alpha diversity and differential abundance analysis between the MS and HC groups were performed using the Wilcoxon test, and p-values were adjusted using the Benjamini-Hochberg algorithm. PERMANOVA was performed to analyze the statistical significance of beta-diversity clustering.

Inter-kingdom correlation analysis was conducted between the bacteria and fungi at the genus level using Spearman's rank correlation, and only correlations showing p-value less than 0.05 are reported. Bootstrapped random forest analysis was conducted using the Boruta package at the suggested significance level of 0.01 with 500 trees for 100 iterations.

**Data Availability:**

The microbiota 16S and ITS2 sequencing analysis data have been deposited to the Sequence Read Archive (https://www.ncbi.nlm.nih.gov/sra) under PRJNA732670 for free public access. All other data are available from the corresponding author upon request.
Results

Fungal microbiota of RRMS patients is different from healthy controls

To characterize the fungal microbiome diversity, we sequenced fecal samples of 20 RRMS patients and 33 healthy controls (HC) using ITS2 rRNA sequencing. A total of 136 fungal genera were identified, with 31 unique to the HC group, 73 unique to the MS group, and 32 genera were present in both groups. Alpha diversity using the Chao1 index showed greater richness in MS than in HC (Fig 1A), though this was not statistically significant (p > 0.08). Principal coordinate analysis of beta diversity using Bray-Curtis dissimilarity (Fig 1B) demonstrated distinct clustering of the mycobiome of MS compared with HC (p = 0.004).

Overall, the two most prominent phyla by relative abundance were Ascomycota and Basidiomycota (Fig 1C), and the top five fungal genera were Saccharomyces, Candida, Malassezia, Penicillium, and Cladosporium (Fig 1D), comprising 75.5% of all the identified fungal genera in the samples.

Examining the differential abundance between the MS and HC groups at the phylum level showed a lower abundance of Ascomycota (padj = 0.011) and higher abundance of Basidiomycota (padj = 0.011) in the MS group compared to HC (padj = 0.011 and 0.011, respectively). Previous studies in IBD and AS have shown that there is a strong negative correlation between Basidiomycota and Ascomycota characterized by a higher Basidiomycota/Ascomycota ratio in IBD\textsuperscript{17} and AS\textsuperscript{16}. In this study, we also observed that MS patients have an increased Basidiomycota/Ascomycota ratio (p = 0.0053) compared to HC (Fig 1E). Thus, our data demonstrates a phylum-level shift in MS towards Basidiomycota and away from Ascomycota. Interestingly, within the
Ascomycota phylum, the genera *Candida* (padj = 0.021) and *Epicoccum* (padj = 0.021) were more abundant in the MS group (Fig 2A & B), while only *Saccharomyces* (padj = 0.021) was depleted in the MS group (Fig 2B) compared with HC. Species-level analysis showed *Saccharomyces cerevisiae* comprising 88% of the *Saccharomyces* genus and *Candida albicans* comprising 81% of the *Candida* genus across all samples. Thus, our study finds that MS patients have a distinct mycobiome compared to HC with enrichment or depletion of certain fungi.

**Random Forest Analysis to identify the potential discriminating fungi**

We next assessed the ability of the gut mycobiota profile to predict disease phenotype in our samples. A bootstrapped random forest algorithm of 500 trees was used to generate a predictive model based on the gut mycobiota profiles of the samples. The Boruta algorithm was then utilized to identify the fungal genera that were most important in distinguishing between MS and HC samples at a significance level of 0.01. Interestingly, the four aforementioned genera, *Candida*, *Saccharomyces*, *Epicoccum*, and *Malassezia*, were again identified (Fig 2C). The Boruta algorithm additionally identified *Penicillium* which is decreased in MS, and *Malassezia*, which is increased in MS (Fig 2C), as a significantly important feature in distinguishing MS from HC. Thus, the random forest classification analysis suggests that these fungal taxa could be important discriminators between MS and HC.

**Functional Profile of Gut Mycobiome in MS Patients**

Analysis of the mycobiome functional profile determined from the FungalTraits database revealed several enzymes whose enrichment differed significantly between MS and HC samples. Levels of amino acid permease (padj = 0.029), cellobiohydrolase (padj = 0.011), endoglucanase (padj = 0.029), and invertase (padj = 0.0211) were lower in MS patients compared to HC (Fig 3), while amylase (padj = 0.011) was increased in these patients.
These results suggest potential differences in fungal metabolism and functional phenotypes in the gut of MS patients.

**Gut Bacterial Microbiome of RRMS patients is different from healthy controls**

We also characterized the bacterial microbiome of the same samples using 16S rRNA (V3-V4) sequencing. A total of 285 bacterial genera were identified, with 196 genera in the HC group and 185 genera in the MS group. Alpha diversity by Chao1 index demonstrated decreased bacterial richness in the MS group compared with the HC group (p = 0.020; Fig 4A), aligning with our previous study showing less bacterial diversity in the gut of MS patients2. Additionally, the MS and HC groups clustered separately on principal coordinate analysis of beta diversity using Bray-Curtis dissimilarity (p = 0.004; Fig 4B), indicating distinct microbiota profiles.

The five most prominent bacterial phyla across all the samples were **Actinobacteria**, **Bacteroidetes**, **Firmicutes**, **Proteobacteria**, and **Verrucomicrobiota**, accounting for 98.8% of all identified bacteria (Fig 4C). Within the **Bacteroidetes** phylum, the **Barnesiellaceae** (padj = 4.4e-3) family and one of its genera, **Barnesiella** (padj = 5.9e-3), as well as the genus **Odoribacter** (**Odoribacteraceae** family; padj = 0.045) were decreased in the MS group compared to HC (Supp 1 and Fig 4D). The other differentially abundant family, **Eggerthellaceae** (**Actinobacteria** phylum; padj = 6.3e-3) and one of its genera, **Eggerthella** (padj = 2.4e-4), were overabundant in MS compared to HC (Supp 1 and Fig 4E). Within the **Firmicutes** phylum, an uncultured genus of the **Oscillospiraceae** family (**Oscillospiraceae** UCG 003) was depleted in the MS group compared to HC (padj = 0.045) while **Blautia** (padj = 0.045) and **Hungatella** (padj = 0.045) were increased in MS (Fig 4E). We have previously also shown **Prevotella** to be significantly decreased in MS patients and beneficial in alleviating symptoms of EAE in mice through modulation of...
the gut microbiome. In this study, the relative abundance of *Prevotella* was decreased in the MS group, though not significantly (padj = 0.70; Supp 2). Similarly, *Akkermansia* was increased in MS but did not reach statistical significance (padj = 0.73; Supp 2). Overall, the shift in the bacterial microbiome in MS patients in this study reveals a number of genera, several of which have already been described in MS microbiome literature.

**Correlation between Gut Mycobiome and Microbiome**

Comparing mycobiome changes with alterations in the bacterial microbiome of the samples show that with decreased bacterial richness comes increased fungal richness. The ITS2/16S ratio is increased in the MS group compared to the HC group (p = 0.040; Fig 5A). Across all samples, bacterial and fungal richness were negatively correlated (R = -0.28, p = 0.042; Fig 5B).

Spearman correlation analysis reveals distinct fungal and bacterial interactions in MS and HC (Fig 5C). The microbial interactions in the HC group are dominated by negative correlations between several bacteria and six fungi. Meanwhile, the MS group exhibited more diverse relationships between the two kingdoms, with numerous positive associations appearing between various fungi and bacteria. MS patients with increased *Blautia* and *Lactococcus* in their stool had decreased *Saccharomyces* and increased *Candida*. *Candida* was also positively associated with *Alistipes* and *Akkermansia*.

Additionally, *Epicoccum* and *Cladosporium* are negatively correlated with *Prevotella*. *Penicillium* is negatively correlated with *Parasutterella* and *Holdemania* and positive correlated with *Terrisporobacter* and *Turicibacter*. *Aspergillus* shares only positive correlations with multiple bacteria: *Paludicola*, an uncultured genus of the *Oscillospiraceae* family, and an uncultured genus of the *Butyricicoccaceae* family.
This demonstrates a shift towards potentially more complex bacterial-fungal relationships in MS patients compared to HC.

**Discussion**

This is the first study to characterize the gut fungal microbiome (mycobiome) profile of RRMS patients compared with healthy individuals and identify connections between the mycobiome and bacterial microbiome. Overall, the results of this study demonstrate that the mycobiota of MS patients differs from that of healthy individuals with enrichment of *Candida*, and *Epicoccum*, lower relative abundance of *Saccharomyces*, and increased ITS2/16S ratio in MS patients. This study highlights the important role of the understudied fungal microbiome in MS.

We observed that *Ascomycota* and *Basidiomycota* were the main phyla present in MS and HC. Ascomycota and Basidiomycota were negatively correlated with each other which had been previously shown in other autoimmune diseases such as IBD and AS. MS patients had increased *Basidiomycota* compared to *Ascomycota* and an increase in *Basidiomycota/Ascomycota* ratio, similar to the previously reported ratio in IBD. While in AS patients, *Ascomycota* abundance was increased, and *Basidiomycota* abundance was decreased. These related findings suggest that shifts in *Ascomycota* and *Basidiomycota* proportions might play a role in the pathogenesis of inflammatory diseases including MS. We detected two fungal genera, *Candida* and *Epicoccum*, with higher relative abundance, and one genus, *Saccharomyces*, with lower abundance in MS compared to HC. Although fungi belonging to *Candida* species are considered commensal microbes, this genus has also been linked with pathogenicity, especially in individuals with compromised immunity due to antibiotic usage, immunodeficiency, or barrier breach. Interestingly, *Candida* species overgrowth had been reported in other autoimmune diseases such as IBD. Thus, it
is possible that bacterial dysbiosis and inflammatory conditions in the gut create favorable environments for expansion of Candida species which can then exacerbate inflammatory diseases such as MS and IBD through induction of pro-inflammatory response. In contrast to Candida, Saccharomyces, which is considered a beneficial fungus, has been shown to be decreased in IBD\textsuperscript{17, 31}. Our findings showing a decreased abundance of Saccharomyces and increased abundance of Candida in MS is in concordance with previous findings in IBD\textsuperscript{17}. Additionally, in colitis it has been observed that S. cerevisiae may exhibit regulatory effects on the host, notably by inducing higher IL-10 production from dendritic cells compared to Candida\textsuperscript{17}. Moreover, colonic tissue of mice receiving S. cerevisiae\textsuperscript{31} showed increased expression of IL-10, suggesting either S. cerevisiae has anti-inflammatory potential or is poorly adapted to an inflammatory environment\textsuperscript{17}. As MS is an inflammatory disease, gut mucosal inflammation might explain the decrease in Saccharomyces and increase in Candida.

Gut bacterial profiling in this study revealed an increased relative abundance of Hungatella, Eggerthella, and Blautia as well as decreased relative abundance of Odoribacter, Barnesiella, and an unidentified genus of Oscillospiraceae. These findings align with previous MS microbiome studies showing increased relative abundance of Blautia and Eggerthella in MS\textsuperscript{2, 3} and decreased relative abundance of Barnesiella \textsuperscript{1}. Additionally, one study using a mouse model had shown increased Odoribacter in mice with relapsing-remitting EAE\textsuperscript{32}. Previous studies have shown depletion of Prevotella and higher abundance of Akkermansia in MS patients\textsuperscript{1, 2, 33}, and while we also observed similar trends, these did not reach statistical significance. A number of factors could have contributed to differences in the microbiome profiles of our study, including different 16S rRNA variable region-specific primers, a more recently developed analysis pipeline (ASV vs OTU-based taxonomic mapping), as well as the geographical origin of
the samples\textsuperscript{34-36}. Additionally, our female to male ratio of 5:1 in this study is higher than 3:1\textsuperscript{1,2} and 2:1\textsuperscript{6} in previous studies, which could contribute to differences.

In a healthy state, there is a symbiosis between bacteria, fungi, viruses, and the host. Similar to bacteria, commensal fungi have been shown to play a significant role in maintaining immune homeostasis, and perturbation of the healthy mycobiome can influence the local and peripheral immune systems directly or indirectly through modulation of bacterial populations\textsuperscript{37-39}. Fungi may directly interact with the immune system in the gut and influence innate and systemic immune response. Intestinal fungi and their metabolic products may also leak out of the enteric luminal surfaces and activate immune cells\textsuperscript{13, 40-42}. Besides the direct effect of fungi on the host, relationships between fungi and bacteria might play a critical role in host immunity. Fungi and bacteria can compete for nutrients and adhesion sites and modulate the environment\textsuperscript{43, 44}. Antibiotics have been linked to increasing growth of fungi in the human intestinal tract\textsuperscript{45}. In the mouse model, an increased fungal abundance was observed after antibiotic treatment which reduced after antibiotic cessation, pointing towards a balancing mechanism between microbiota and mycobiota in the gut\textsuperscript{17, 18}. Decreased bacterial diversity favors the growth of certain fungal species, and certain fungi can switch from commensal to pathogenic phenotypes\textsuperscript{43}. In this study, we observed decreased bacterial alpha diversity and increased fungal to bacterial richness in MS compared to HC, as evident from the ITS2/16S ratio, suggesting that the MS gut might favor fungi at the expense of bacteria.

The microbial interactions in the HC group are dominated by negative correlations between several bacteria and fungi. However, our MS group exhibited more diverse relationships between the bacteria and fungi. Specifically, MS patients with increased \textit{Candida} also had increased \textit{Alistipes}, \textit{Akkermansia}, and \textit{Lachnospiraceae GCA}, and
those with increased *Saccharomyces* also had increased *Lactococcus, Blautia,* and *Romboutsia*. It has been shown that in healthy mucosa, yeast (e.g., *Candida albicans*) is kept at low levels or excluded by the indigenous bacterial microbiota by the mechanism of colonization resistance which is not yet defined\(^4^6\). In fact, a previous study showed that germ-free mice are highly susceptible to *Candida* infection and antibiotics treatment leads to an expansion of fungal population\(^1^8, 4^7\). Additionally, IBD patients with decreased abundance of *Saccharomyces* had decreased abundance of several bacteria that are depleted in IBD, such as *Bifidobacterium, Blautia, Roseburia,* and *Ruminococcus*, while unidentified *Malasseziale* followed an opposing pattern\(^1^7\). In AS patients, there is a positive corelation between abundance of *Saccharomyces* to *Clostridium sensu stricto, Escherichia/Shigella,* and *Veillonella* and negative correlation between abundance of *Saccharomyces* to *Roseburia* and *Faecalibacterium*\(^1^6\). Thus, our study along with previous studies suggests a role of fungal mycobiome in MS as well as other autoimmune diseases.

Functional profiling of the fungal communities in this study suggests modulation of fungal carbohydrate and amino acid pathways. More specifically, while the functional profile of the MS group mycobiome suggested decreased cellulose metabolism by the gut mycobiome (decreased cellobiohydrolase and endoglucanase), starch metabolism by amylase was suggested to be increased. Invertase, which was decreased in MS, can be produced by fungi, mainly *Saccharomyces cerevisiae*. It hydrolyzes sucrose into glucose and fructose and also has antibacterial and antioxidant properties. Sucrose provides a primary substrate for the generation of starch and cellulose\(^4^8\). Both cellulose and starch are carbohydrates that can be metabolized into short-chain fatty acids (SCFAs) in the human gut. However, while the metabolism of starches has been shown to increase fecal butyrate levels\(^4^9\), metabolism of cellulose-related structures\(^5^0\) leads to
increased gut propionate levels\textsuperscript{51}.

Currently, there are limited tools for functional analysis of the mycobiome, and this study utilized a still growing database of fungal functional profiles. Moreover, functional profiling of fungi is difficult due to the difficulty of culturing fungi and highly varied, environment-dependent fungal trophic modes\textsuperscript{24}. The quality of these profiles derived from ITS1/2 sequencing data will continue to improve as the field progresses and these databases grow. Alternatively, while more cost and time-intensive, whole genome sequencing of the mycobiome would provide more detailed data for more reliable functional analysis of the mycobiome.

We acknowledge that our small sample size is a limitation of the study. However, the subject recruitment was impacted due to the COVID-19 pandemic. We did not want to recruit new subjects as the study may have been confounded by COVID-19 infections. Due to small sample size, we were also not able to analyze the impact of different drug treatments on bacterial and fungal composition, therefore we cannot rule out the impact of treatment on the microbiome. Another limitation of this study was the different BMI distributions in the HC and MS groups. Interestingly, BMI was not significantly correlated to most of the differentially abundant bacteria or fungi except for *Eggerthella*. Nonetheless, future studies would benefit from larger sample sizes in order to better isolate the microbiota differences related to MS from those related to BMI and different treatment groups.

**Conclusion**

We observed that RRMS patients exhibit altered gut mycobiome along with altered gut microbiome. RRMS patients also showed variations in bacterial and fungal relationships with increased ratios of fungal/bacterial abundance and richness. Shifts in fungal functional profiles were also observed in RRMS patients compared to HC. Thus, our
data suggest that the mycobiome may play an important role in the pathobiology of MS. Whether changes in the mycobiome play a role in initiating MS pathophysiology or modulating its presentation, or whether the pathology of MS leads to changes in the fungal mycobiome remains an important question for future studies.

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Disclosure

AM is one of the inventors of a technology claiming the use of *Prevotella histicola* to treat autoimmune diseases. AM received royalties from Mayo Clinic (paid by Evelo Biosciences). MY, SA, RS, SS, SJ, JH, SC, HO, NG, MP, CC, KW, TC, JK declare no commercial or financial relationships that could be a potential conflict of interest.
Figure 1. Fungal microbiota of RRMS patients is different from healthy controls

(A) Fungal ASV richness estimated by Chao1 index in MS and HC. (B) Principal coordinate analysis of Bray-Curtis dissimilarity of HC and MS shows that the mycobiome of HC and MS are distinct (PERMANOVA: p=0.004). Ellipses correspond to 95% confidence intervals around the centroids for each group. (C) Bar plot showing the relative abundances of fungal phyla. *Basidiomycota* was increased and *Ascomycota* was decreased in MS compared to HC. (D) Bar plot showing the top 10 fungal genera in HC and MS (determined by average relative abundance across all samples). The top 10 genera account for 85.5% of all identified fungal genera. (E) *Basidiomycota/Ascomycota* ratio is significantly increased in MS (p=0.0053).
Figure 2. Differentially abundant fungal genera in RRMS versus healthy controls.
(A) Bar plot showing relative abundances of differentially abundant taxa (p < 0.05) at the phylum, family, and genus level. (B) Differentially abundant fungal genera in MS and HC using Wilcoxon signed rank test and adjusted for multiple comparisons with the Benjamini-Hochberg method at a significance level of 0.05. *Candida*, *Epicoccum*, and *Malassezia* are increased in MS compared to HC. *Saccharomyces* is decreased in MS compared to HC. *Penicillium* was identified in random forest analysis as a significant feature and was decreased in MS, though the Wilcoxon test did not reach statistical significance after adjusting for multiple comparisons. Abundance values are sum-scaled to 1 million and generalized log-transformed. The * symbol indicates p-value <0.05. (C) Importance of features determined by random forest and tested for significance with the Boruta algorithm at a significance level of 0.01.

**Figure 3. Functional Profile of Gut Mycobiome in MS Patients** Differentially enriched fungal functions using Wilcoxon signed rank test and adjusted for multiple comparisons with the Benjamini-Hochberg method at a significance level of 0.05. Amino acid permease, cellobiohydrolase, endoglucanase, and invertase are decreased in MS compared to HC. Amylase is increased in MS compared to HC. Abundance values are sum-scaled to 1 million and generalized log-transformed.
Figure 4. Gut Bacterial Microbiome of RRMS patients is different from healthy (A) Bacterial ASV richness estimated by Chao1 index showing decreased bacterial richness in MS (p = 0.020). (B) Principal coordinate analysis of Bray-Curtis dissimilarity of HC and MS showing that the microbiome of HC and MS are distinct (PERMANOVA: p=0.004). Ellipses are visual and do not correspond to any statistical analysis. (C) Abundance of bacteria at phylum level. (D) Differentially abundant bacteria between HC and MS using Wilcoxon signed rank test and adjusted for multiple comparisons with the Benjamini-Hochberg method at a significance level of 0.05. Abundance values are sum-scaled to 1 million and generalized log-transformed. The *,**, and *** symbol indicates p-values of <0.05, <0.01, <0.001, respectively.
Figure 5. Correlation between Gut Mycobiome and Microbiome  
(A) Ratio of ITS2 to 16S compared between groups. (B) Linear regression of fungal and bacterial richness shows a negative correlation (Spearman’s $R = -0.28$, $p = 0.042$). (C) Correlation matrix between bacteria and fungi using Spearman correlation. Values range from $-1$ to $1$ with positive values as orange and red and negative values as purple and blue ($-1 \leq R \leq 1$). Only statistically significant correlations ($p < 0.05$) are shown.