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# Gut microbiota from multiple sclerosis patients enables spontaneous autoimmune encephalomyelitis in mice

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There is emerging evidence that the commensal microbiota has a role in the pathogenesis of multiple sclerosis (MS), a putative autoimmune disease of the CNS. Here, we compared the gut microbial composition of 34 monozygotic twin pairs discordant for MS. While there were no major differences in the overall microbial profiles, we found a significant increase in some taxa such as *Akkermansia* in untreated MS twins. Furthermore, most notably, when transplanted to a transgenic mouse model of spontaneous brain autoimmunity, MS twin-derived microbiota induced a significantly higher incidence of autoimmunity than the healthy twin-derived microbiota. The microbial profiles of the colonized mice showed a high intraindividual and remarkable temporal stability with several differences, including *Sutterella*, an organism shown to induce a protective immunoregulatory profile in vitro. Immune cells from mouse recipients of MS-twin samples produced less IL-10 than immune cells from mice colonized with healthy-twin samples. IL-10 may have a regulatory role in spontaneous CNS autoimmunity, as neutralization of the cytokine in mice colonized with healthy-twin fecal samples increased disease incidence. These findings provide evidence that MS-derived microbiota contain factors that precipitate an MS-like autoimmune disease in a transgenic mouse model. They hence encourage the detailed search for protective and pathogenic microbial components in human MS.

gut microbiome | multiple sclerosis | experimental autoimmune encephalomyelitis | twin study | germ-free mice

The risk of developing multiple sclerosis (MS) is driven by both genetic factors and environmental exposures (1). Risk genes have been determined by large-scale genome-wide association studies (GWAS), which identified more than 200 different DNA variants associated with disease susceptibility (2). Environmental risk factors include smoking, reduced exposure to sunlight, and infection with Epstein-Barr virus (3). Very recently, the intestinal microbiota emerged as an additional potential triggering factor (4, 5).

The notion that commensal gut bacteria are causally related to brain autoimmunity is supported by a transgenic mouse model of spontaneous experimental autoimmune encephalomyelitis (EAE). In this model, nearly all animals raised in specific pathogen-free (SPF) conditions develop a relapsing-remitting (RR) variant of the disease within months of age (6). Importantly, when kept in a germ-free environment, animals from the same strain remain disease-free. However, spontaneous disease promptly follows exposure of germ-free mice to SPF-derived fecal material (7). Taken together, these observations indicate that the encephalitogenic immune response observed in these mice is mediated by the intestinal microbiota.

Translation of these experimental observations into human MS poses considerable practical challenges. Complicating factors are, in particular, genetic diversity (8) and lifestyle, such as diet (9), both of which profoundly impact the individual gut microbiota. In addition, age, therapy, and neurological condition (10) might

also affect the gut microbial composition. In an attempt to eliminate genetic variance and reduce environmental variance to a minimum, we identified and recruited 34 monozygotic (MZ) twin pairs, discordant for MS, for a microbiome study. All the probands were of Caucasian origin and had grown up together with their healthy twins to adulthood in Germany. We studied their gut microbiota in two tiers: first, intestinal microbial profiles of MS twins and healthy twins were compared by 16S ribosomal RNA (rRNA) amplicon and metagenomic shotgun sequencing. Second, we transplanted fecal samples from selected twin pairs to germ-free mice to assess functional differences in the human intestinal microbiota of MS and healthy twins.

## Results

**MZ Twin Cohorts Discordant for MS.** We assembled a cohort of 34 MZ twin pairs clinically discordant for MS. In each pair, one twin has clinically definite MS according to the current diagnostic criteria (11), whereas the co-twin is unaffected. Our MS twin cohort resembles the general MS population with respect to female

## Significance

Studies using experimental models have indicated that multiple sclerosis (MS)-like disease can be triggered in the gut following interactions of brain autoimmune T lymphocytes with local microbiota. Here we studied the gut microbiota from monozygotic human twin pairs discordant for multiple sclerosis. When we transferred human-derived microbiota into transgenic mice expressing a myelin autoantigen-specific T cell receptor, we found that gut microbiota from multiple sclerosis-affected twins induced CNS-specific autoimmunity at a higher incidence than microbiota from healthy co-twins. Our results offer functional evidence that human microbiome components contribute to CNS-specific autoimmunity.

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The authors declare no conflict of interest.

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See Commentary on page 10528.

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**Table 1. Demographic and clinical characteristics of MZ twins discordant for MS**

Characteristic	MZ twins
No. of pairs	34
Gender, female/male	26/8
Age in years (range in years)	41.3 ± 10.8 (21–63)
Age in years at disease onset (range in years)	28.0 ± 9.0 (14–47)
Time in years clinically discordant for MS (range in years)	13.2 ± 9.6 (1–33)
Pairs clinically discordant for MS longer than 10 y (%)	18 (53)
Pairs with a positive family history of MS (%)	11 (32)
MS type, <i>n</i> (%)	
CIS	3 (9)
RR-MS	22 (65)
SP-MS	7 (21)
PP-MS	2 (6)
Disease-modifying treatment, <i>n</i> (%)	
Untreated	15 (44)
IFN-β	13 (38)
Natalizumab	4 (12)
Glatiramer acetate	1 (3)
Azathioprine	1 (3)

Continuous data are expressed as mean ± SD. Categorical data are expressed as the number of observations (%). CIS, clinically isolated syndrome; PP-MS, primary progressive MS; RR-MS, relapsing–remitting MS; SP-MS, secondary progressive MS.

preponderance, age distribution, age at onset, and distribution of clinical subtypes [RR MS, secondary progressive MS, primary progressive MS, and clinically isolated syndrome (CIS)] (Table 1) (12). The twin cohort is homogenous in terms of geographic environment and genetic background. Furthermore, immunophenotyping of peripheral blood mononuclear cells (PBMCs) from all twin pairs did not show relevant differences in the frequencies of CD4<sup>+</sup> T cells, Th1 cells, Th17 cells, and Treg cells (Fig. S1).

**Microbial Profiling of MS-Discordant MZ Twins by 16S rRNA and Metagenomic Shotgun Sequencing.** To search for differences between microbiomes of MS-affected versus nonaffected twins, we applied 16S rRNA sequencing and metagenomics shotgun sequencing. 16S rRNA amplicon sequencing revealed no major differences in microbial community structures. Specifically, MS twins and healthy twins exhibited comparable microbial community richness (alpha diversity) (Fig. 1A). A concomitant principal coordinate analysis (PCoA) also did not point to disease-dependent clustering of samples (Fig. 1B). Further, analysis at the level of individual taxa [up to genera or the operational taxonomic units (OTUs) level] did not indicate significant differences between MS and healthy twins. However, when patients were stratified for the use of disease-modifying therapy, several taxa (most notably *Akkermansia muciniphila*) were significantly increased in untreated MS twin siblings (Fig. S2).

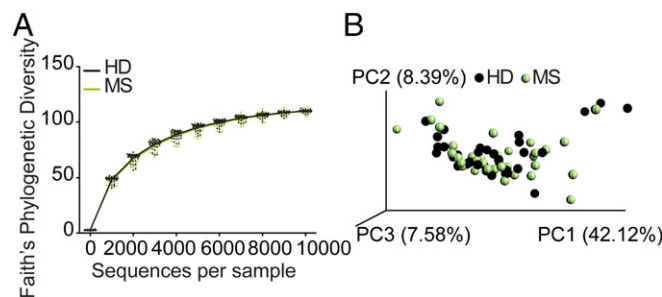
Comparing the microbiome composition of MS and healthy twins is a first, purely descriptive step in exploring the role of the microbiota in MS. Disease-related functional differences in the microbiomes of healthy or MS twins might be revealed by shotgun metagenomic sequencing. Using Human Microbiome Project (HMP) Unified Metabolic Analysis Network (HUMAN2) tool1, we assessed the microbial community, performed functional profiling on the bacterial DNA sequences, and identified 242 unique bacterial species across 98 genera, 1.2 million gene families, and 558 bacterial metabolic pathways. Interestingly, analysis of gut bacterial composition showed higher similarity between discordant

twin siblings than among unrelated twin pairs (Fig. S3), thus confirming the influence of the host's genetics in the composition of the gut microbiome. To determine possible associations of specific bacterial species, gene families, or metabolic pathways with MS, we performed logistic regression analysis. In harmony with 16S sequencing findings, after adjusting for twin pair, number of genome equivalents sequenced, and multiple comparisons, no specific bacterial species, gene families, or metabolic pathways were significantly associated with the MS disease phenotype.

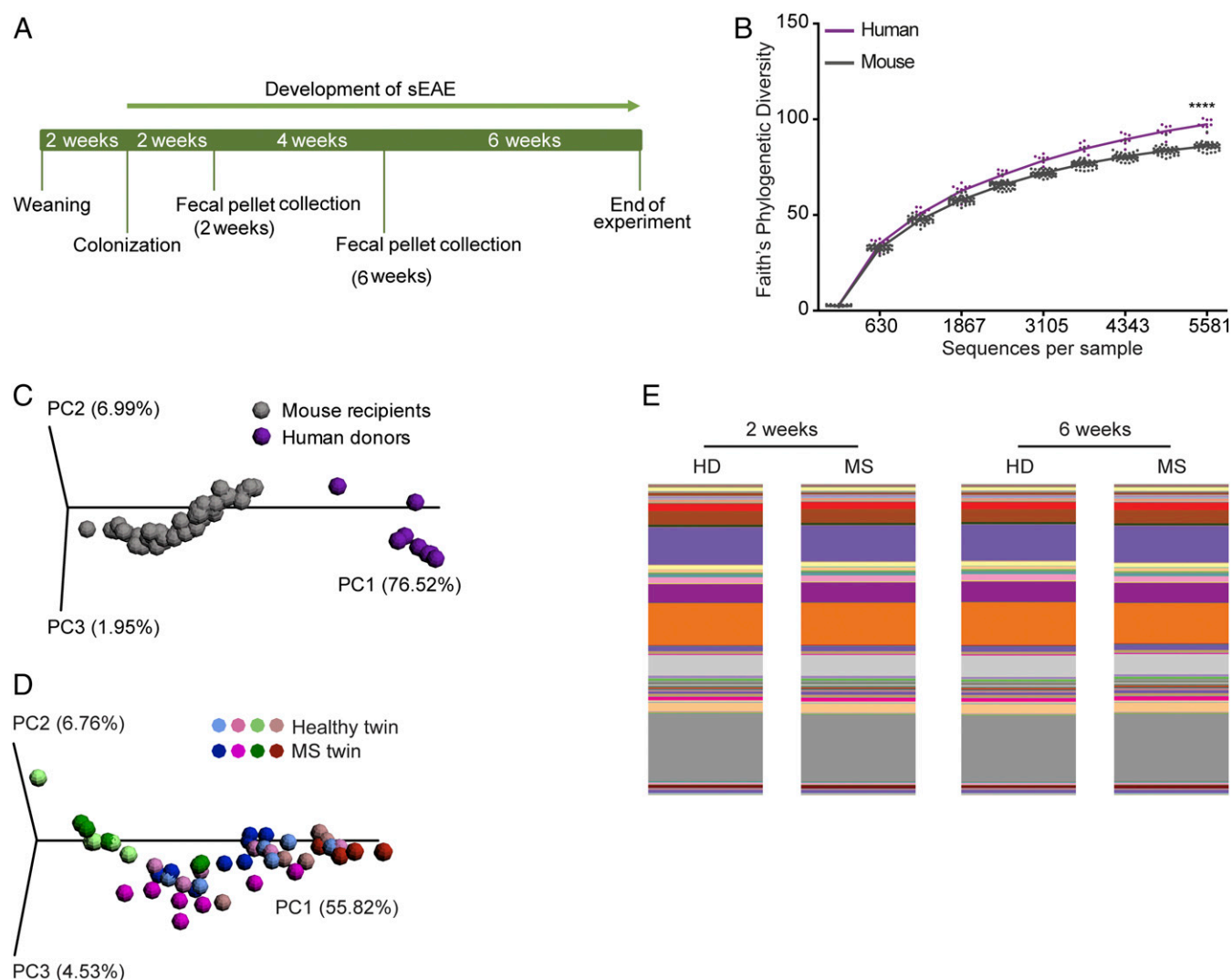
**RR Mice Colonized with Human MS Twin-Derived Microbiota.** To explore the functional role of human microbiota in CNS inflammation, we used our previously described spontaneous RR mouse model (6, 7). Several qualities render these mice a particularly promising reporter system. RR mice express a transgenic myelin oligodendrocyte glycoprotein (MOG)-specific T cell receptor in more than 70% of their CD4<sup>+</sup> T cells. Beginning at 2 mo of age, more than 80% of the mice spontaneously develop an inflammatory demyelinating disease that recapitulates features of early human MS: The disease course is often relapsing–remitting, affects varying parts of the brain and spinal cord with lesions featuring round-cell infiltrates and large confluent areas of demyelination with axonal degeneration, and responds to B cell-depleting therapy (6). Most importantly, spontaneous disease critically depends on an intact commensal microbiome, as germ-free RR mice remain healthy (7).

We reasoned that if the gut microbiota is a determinant factor contributing to the severity and course of human MS, transplantation of human fecal microbial communities from MS twins should result in a higher frequency of spontaneous EAE than the transfer of fecal material from healthy twins. We selected a subgroup of five twin pairs who were all discordant for RR MS and in which the affected twins had different disease durations and were either without therapy or were treated with IFN-β (for further details refer to Table S1). Each donor fecal sample was diluted in prereduced PBS and introduced into a group of 6-wk-old, germ-free RR mice by oral gavage. Mouse fecal pellets were collected 2 and 6 wk after transplantation to assess the stability as well as composition of the intestinal microbiota by 16S rRNA amplicon sequencing (Fig. 2A).

Unsurprisingly, we detected significantly reduced alpha diversity (~10%) in colonized mice compared with the donor humans (Fig. 2B). PCoA confirmed the divergent gut microbial community structures in human and mouse samples (Fig. 2C), indicating that only a subset of microbial species from human donors was able to colonize mouse recipients. The microbiota transfer rate was consistent across samples and independent of the disease state of the donor. Since the composition of the intestinal microbiota is profoundly imprinted by host genetics, diet,



**Fig. 1.** No overt differences in alpha or beta diversity were detected by comparing the fecal microbial profiles of healthy twins with those of MS twins. (A) Phylogenetic (alpha) diversity of fecal microbiota in healthy twins (HD, black; *n* = 34) and MS twins (MS, green; *n* = 34). (B) PCoA of weighted UniFrac community distances for fecal microbiota of healthy twins (black; *n* = 34) and MS twins (green; *n* = 34).



**Fig. 2.** Human microbiota can be efficiently transferred to mouse recipients. (A) Design of humanized gnotobiotic mouse experiment: 6-wk-old, germ-free RR SJL/J mice were gavaged with fecal samples from the healthy twin or the MS twin from a MZ twin pair. Fecal samples for 16S rRNA sequencing analysis were harvested at 2 and 6 wk after colonization. Humanized gnotobiotic RR mice were observed for the development of clinical signs of EAE for 12 wk. (B) Phylogenetic (alpha) diversity of fecal microbiota in human twin pairs (purple;  $n = 8$ ) and mouse recipients (gray;  $n = 47$ ). \*\*\*\* $P < 0.0001$  (Mann–Whitney  $u$  test). (C) PCoA of weighted UniFrac community distances comparing fecal microbiota of human donors (purple;  $n = 8$ ) and mouse recipients (gray;  $n = 47$ ). (D) PCoA of weighted UniFrac community distances for fecal microbiota of mice transplanted with microbiota from healthy ( $n = 23$ ) or MS ( $n = 24$ ) twins. (E) Mean relative abundances of bacterial genera in fecal samples of mice that received microbiota of healthy donors (HD;  $n = 20$ –26) or MS patients (MS;  $n = 22$ –26) at 2 and 6 wk after transplantation.

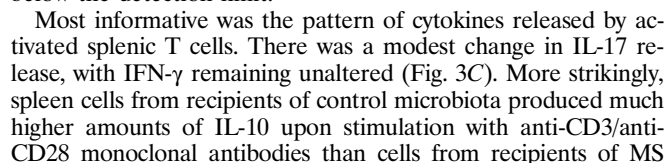
and other environmental factors (13), it was not surprising that the composition of gnotobiotic mouse fecal samples reflected incomplete colonization by the human donor microbiota. Similar changes were noted in previous human-to-mouse transplant experiments (14, 15). PCoA of mouse samples demonstrated clustering by donor and twin pair but not by EAE disease state (Fig. 2D). In addition, comparison of the percentage of microbial genera that was transferred from the human donor to recipient mice did not indicate any disease-based trends and highlighted a comparatively high mean transfer rate of 90.3% (Table S2). Finally, the recipient mouse microbiome remained highly stable between 2 and 6 wk after colonization, as indicated by representative bar plots of mean relative abundance of bacterial genera and corresponding analysis (Fig. 2E).

**Increased Incidence of Spontaneous EAE in RR Mice Colonized with Microbiota from MS-Affected Twins.** Remarkably, RR mice transplanted with MS patient-derived microbiota developed spontaneous EAE at a higher frequency than animals colonized with

intestinal bacteria from healthy twins (Fig. 3A). The triggering of EAE in germ-free RR mice by human microbial transfer per se is notable, considering that so far this has been achieved only with some mouse-associated bacteria (7). However, monocolonization with experimental bacterial consortia (such as altered Schaedler flora) and segmented filamentous bacteria did not induce spontaneous EAE (7).

We next set out to determine the differences that made MS-derived microbiota more pathogenic than healthy-derived samples. Analysis of the mouse microbiome at the level of individual taxa revealed significant differences between MS- and healthy twin-colonized mice, both at the genus and OTU taxonomical levels. While the genera *Adlercreutzia* and *Tannerella* were more abundant in the feces of mice transplanted with the healthy twin fecal material, fewer bacteria of the genera *Ruminococcus* were present in these mice. The most significant difference, however, was a reduced abundance of the genus *Sutterella* in mice colonized with microbiota from the MS twins compared with recipients of



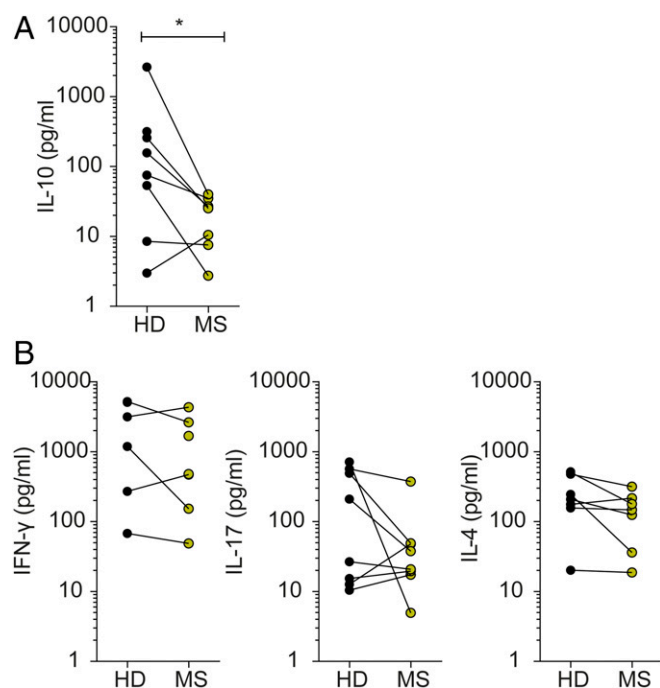


microbiota (Fig. 3C). This difference was of functional relevance. We colonized germ-free RR mice with healthy donor fecal material and subsequently treated them with anti-IL-10 neutralizing or isotype control antibodies. Blocking IL-10 increased disease incidence compared with isotype control-treated mice (from 30 to 55%;  $P = 0.0006$ , Fisher's exact test) (Fig. 3D).

The immunological profiles observed in the colonized RR mice were consistent with the patterns observed in PBMCs from a subgroup of eight twin pairs which included several donors used for the mouse colonization experiments. There, anti-CD3/anti-CD28 stimulation induced stronger production of anti-inflammatory IL-10 in T cells from healthy than from MS donors (Fig. 4A). In contrast, we found no differences in the frequencies of CD4<sup>+</sup> T cells, Th1 cells, Th17 cells, and Treg cells (Fig. S1). Also, production of IFN- $\gamma$ , IL-17, and IL-4 cytokines in response to in vitro stimulation with phytohemagglutinin (PHA) or anti-CD3/anti-CD28 was largely indistinguishable between T cells from MS-affected and healthy twins (Fig. 4B).

## Discussion

Chronic inflammatory diseases have been recently associated with altered profiles of the intestinal bacterial flora (21). These changes were revealed by studies making use of new sequencing technologies, which, however, cannot per se resolve whether the microbial changes were primary or secondary to cognate diseases. Causal relations were established in metabolic conditions such as obesity by transplanting human-derived microbiota to rodents; these



**Fig. 4.** Gut bacteria from healthy twins trigger an antiinflammatory T cell response. (A) IL-10 production of CD4<sup>+</sup> T cells isolated from PBMCs of selected twin pairs. T cells were stimulated for 96 h with 1  $\mu$ M anti-CD3 and anti-CD28 antibodies. Levels of IL-10 (seven pairs, two singletons) in the supernatants were measured by ELISA. \* $P < 0.05$  (Wilcoxon test). (B) Cytokine profiles of CD4<sup>+</sup> T cells isolated from PBMCs of selected twin pairs. Depending on PBMC availability and quality, eight twin pairs were selected for further in vitro stimulation assays, and data are depicted for each cytokine where stimulation-dependent production above the individual detection level could be documented. T cells were stimulated for 48 h with 5  $\mu$ M PHA. Levels of IFN- $\gamma$  (five pairs, three singletons), IL-17 (eight pairs), and IL-4 (seven pairs, one singleton) in the supernatants were measured by the Luminex Bead-based Multiplex Assay.

recipients of disease-derived fecal samples developed an equivalent metabolic condition (22). Applying similar strategies to studies of chronic autoimmune or degenerative diseases (23–25) has remained more challenging. This definitely applies to exploring links between MS and changes of intestinal microbiota.

Here we took a two-tiered approach to address this question. First, we compared the intestinal microbial profiles of MS twins and healthy twins by microbial 16S rRNA amplicon and metagenomic shotgun sequencing. Second, we transplanted fecal samples from MS-affected and healthy twins into germ-free mice. We did not detect significant differences in MS-related microbial profiles between MS-affected and healthy twins. In this respect, our results differ from findings reported by others. A recent study using pyrosequencing of 16S rRNA gene in 20 Japanese patients with RR MS detected associations with *Clostridium* and *Bacteroidetes* species (26). These signatures were not seen in our cohort, nor were they reported in another independent screening (27). However, it should be noted that these studies (including our own) are relatively limited in size. Second, it is likely that microbial associations vary according to geographical region, human genetic background, and many other factors, including treatment. However, it is worth noting that, after adjusting for treatment, we found *Akkermansia* species were increased in untreated MS twins, which was also found to be increased in another independent cohort of MS patients [see ref. 27 and the companion paper by Cekanaviciute et al. (28)]. Third, fecal material may be suboptimal for studying processes possibly restricted to circumscribed intestinal segments. Gut microbiota differ radically in their functional potential, density, and composition along different segments of the small and large intestine (29). Thus, in the feces, pathogenic organisms populating the small intestine at low density could be eclipsed by high-density colonic communities. Obviously, more detailed studies of much larger cohorts are needed to further explore the complexities of MS-related microbial associations.

The second tier of our strategy, transfer of human microbiota into germ-free RR mice, led to a particularly interesting observation. We found that not only could human microbiota trigger classic RR EAE, but, even more significantly, transplants from MS-affected donors triggered EAE at higher rates than did transplants from their healthy control-derived counterparts. This difference was not large in magnitude but was significant and reproducible. A detailed analysis of 16S rRNA sequences revealed significant differences between MS- and healthy twin-colonized mice at the level of individual taxa (genera and OTUs). The most prominent MS-dependent change was a reduction in the bacterial genus *Sutterella*.

The increased incidence of spontaneous EAE in MS sample-recipient gnotobiotic RR mice could be due to either increased activation of autoimmune effector T cells or a weakened regulatory mechanism. Our results seem to favor the latter. Indeed, functional screening of peripheral blood lymphocytes from a subgroup of monozygotic twins indicated decreased IL-10 production in MS-derived samples. This is in accord with previous work, which described decreases of Tr1-like, IL-10-producing T cells (30–32). The differential IL-10 response noted in human PBMCs was mirrored in gnotobiotic RR mice. In germfree RR mice recolonized with MS-derived fecal samples, spleen T cells produce less IL-10 than their equivalents from control sample recipients. This recalls previous reports describing enhanced IL-10 production in the intestinal immune system driven by bacterial polysaccharide A (33) and protection from actively induced CNS autoimmune disease (34). A relative lack of protective factors rather than an increase in disease-promoting factors is also supported by our observation that in vivo treatment with anti-IL-10 antibody resulted in an increased incidence of spontaneous EAE in our RR model. Although there is currently no

conclusive evidence of a genetic association of MS with the IL-10 gene or IL-10-related pathways, this by no means excludes the possibility that this crucially important pathway is modulated by external factors (e.g., microbiota).

In conclusion, we here provide evidence that human MS-derived microbiota contain factors that precipitate an MS-like autoimmune disease in a humanized transgenic mouse model. This observation provides possibilities for characterizing the precise role and functional mechanisms by which the human intestinal microbiota contributes to the pathogenesis of neuro-inflammatory diseases. The findings may eventually have important implications not only for the pathogenesis but also for the therapy and potentially even prevention of human MS.

## Methods

Details of cohort recruitment, sample acquisition, and information collection are provided in [SI Methods](#). The study was approved by the local Ethics Committee of the Ludwig-Maximilians University Munich, and all participants gave written informed consent. All animal procedures were in accordance with the guidelines of the Committee on Animals of the Max Planck Institute of Neurobiology and the Max Planck Institute of Immunobiology

and Epigenetics with a license from the Regierung von Oberbayern as well as the Regierungspräsidium Freiburg. Procedures for 16S and metagenomic sequencing, data analysis, mouse colonization, flow cytometry, ELISA, cytokine profiling, and real-time qPCR are described in detail in [SI Methods](#).

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