Mining the microbiota to identify gut commensals modulating neuroinflammation in a mouse model of multiple sclerosis

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Abstract

Background: The gut microbiome plays an important role in autoimmunity including multiple sclerosis and its mouse model called experimental autoimmune encephalomyelitis (EAE). Prior studies have demonstrated that the multiple sclerosis gut microbiota can contribute to disease, hence making it a potential therapeutic target. In addition, antibiotic treatment has been shown to ameliorate disease in the EAE mouse model of multiple sclerosis. Yet, to this date, the mechanisms mediating these antibiotic effects are not understood. Furthermore, there is no consensus on the gut-derived bacterial strains that drive neuroinflammation in multiple sclerosis.

Results: Here, we characterized the gut microbiome of untreated and vancomycin-treated EAE mice over time to identify bacteria with neuroimmunomodulatory potential. We observed alterations in the gut microbiota composition following EAE induction. We found that vancomycin treatment ameliorates EAE, and that this protective effect is mediated via the microbiota. Notably, we observed increased abundance of bacteria known to be strong inducers of regulatory T cells, including members of Clostridium clusters XIVa and XVIII in vancomycin-treated mice during the presymptomatic phase of EAE, as well as at disease peak. We identified 50 bacterial taxa that correlate with EAE severity. Interestingly, several of these taxa exist in the human gut, and some of them have been implicated in multiple sclerosis including Anaerotruncus colihominis, a butyrate producer, which had a positive correlation with disease severity. We found that Anaerotruncus colihominis ameliorates EAE, and this is associated with induction of RORγt+ regulatory T cells in the mesenteric lymph nodes.

Conclusions: We identified vancomycin as a potent modulator of the gut-brain axis by promoting the proliferation of bacterial species that induce regulatory T cells. In addition, our findings reveal 50 gut commensals as regulator of the gut-brain axis that can be used to further characterize pathogenic and beneficial host-microbiota interactions in multiple sclerosis patients. Our findings suggest that elevated Anaerotruncus colihominis in multiple sclerosis patients may represent a protective mechanism associated with recovery from the disease.

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The exact autoimmune responses in MS are not fully known, although components of both the adaptive and innate immune systems are involved [5, 6]. Interferon-gamma (IFNγ) producing Th1 and interleukin-17 (IL-17) secreting CD4+ T cells play a central role in the pathogenesis of MS [7, 8].

Genetic and environmental factors contribute to MS development [9]. Among the latter, several studies suggest that intestinal factors modulate MS disease severity [10]. In addition, alterations in the gut microbiota composition [11–16], gut-derived products [17–19], intestinal permeability [20–22], and enteric nervous system functions [23] have been reported in MS patients. The human microbiome encompasses trillions of organisms that inhabit the gut and shape the gut-associated lymphoid tissue. Studies have shown that the gut microbiota shapes the development and function of both innate and adaptive immune cells [24–28]. Different commensals in the gut promote the differentiation of subsets of lymphocytes. Segmented filamentous bacteria (SFB) induce intestinal Th17 [29], Bacteroides fragilis (B. fragilis) colonization of germ-free (GF) mice preferentially induces Th1 cells [30], and polysaccharide A of B. fragilis suppresses Th17 cells in conventional mice [31]. Clostridium clusters IV and XIVa promote T regulatory cells (Tregs) [32], and a subset of Clostridium clusters IV and XIVa which attach to the gut mucosa can also promote Th17 cells [33].

The relationship between the host and its microbiota is generally mutually beneficial [34]. However, perturbations in the composition of the gut microbiota referred to as dysbiosis have been implicated with diseases of various etiologies, including autism, amyotrophic lateral sclerosis (ALS), Parkinson’s disease, and several autoimmune disorders such as inflammatory bowel disease, diabetes, rheumatoid arthritis, and MS [11, 12, 35–42]. Several studies have reported alterations in the gut microbiota composition of MS patients including increases in Akkermansia muciniphila and decreases in butyrate-producing bacteria [11, 14, 15, 40, 41, 43–46]. Whether these gut microbiota alterations in MS contribute to disease pathogenesis or are just a consequence remains unknown. We have previously reported that MS patients and mice at peak EAE have elevated intestinal miR-30d which increases the levels of Akkermansia muciniphila and ameliorates EAE [47]. Furthermore, we recently reported that MS-derived A. muciniphila attenuates EAE clinical scores [40]. These findings suggest that elevated Akkermansia in MS patients may be a consequence of the disease. The finding that transferring the gut microbiota from MS patients into mice exacerbates EAE suggests that the MS gut microbiota can drive neuroinflammation [14, 15]. Consistent with these findings, other studies have reported that long-term antibiotic treatment in MS reduces relapse rates and gadolinium-enhancing lesions as well as improves measures of disability [48, 49]. However, results across gut microbiota studies in MS lack consistency. Sample size, subject heterogeneity, study design, type of controls, geographical location, sequencing platforms, and regions of 16S rRNA gene sequencing may all contribute to lack of reproducibility [50]. As such, the mechanisms through which altered gut-brain axis may contribute to CNS inflammation, demyelination, and axonal loss remain poorly understood. This in turn makes it impossible to determine how the gut microbiome can be targeted for therapeutic purposes in MS.

Studies in animal models demonstrate that the gut microbiota can modulate neuroinflammation. Germ-free mice are protected from EAE, and transferring specific pathogen-free microbiota to these mice restores their susceptibility to EAE [51, 52]. Other studies have reported that oral treatment with broad-spectrum antibiotics significantly altered the gut microbiota and reduced EAE severity in a Treg-dependent manner [53]. Disease amelioration did not occur when antibiotics were given intraperitoneally, thereby bypassing the gut, suggesting that modulation of the gut microbiota produces protective effects. However, these studies did not investigate the gut microbiome composition of the antibiotic-treated mice, and as such was not able to demonstrate that changes in the gut microbiota composition regulate CNS inflammation in EAE mice. Furthermore, SFB, a TH17 inducer, exacerbates EAE [52], whereas Tregs inducers such as polysaccharide A-positive Bacteroides fragilis ameliorate EAE [54]. Prior studies found decreased Prevotella histicola in MS patients, and mice fed human-derived Prevotella histicola are protected from EAE [11, 12, 15, 41, 55–57]. These findings suggest that bacteria depleted in MS can attenuate neuroinflammation. Furthermore, we reported that MS patients receiving a probiotic consisting of a mixture of Lactobacillus-Bifidobacterium-Streptococcus species displayed changes in their gut microbiota composition that were associated with anti-inflammatory immune markers in the periphery [58]. Another study reported that supplementation of MS patients with a mixture of three Lactobacillus species and one Bifidobacterium species was associated with improved EDSS score as well as decreased depression and stress [59]. Taken together, these studies demonstrate that modulating the microbiota has therapeutic potential, and the EAE mouse model is a useful tool to identify human gut-derived bacteria with neuroimmunomodulatory potential.

In this report, we investigated the gut microbiota of untreated and vancomycin-treated EAE mice at multiple time points spanning pre- and postimmunization state with myelin oligodendrocyte glycoprotein (MOG)
to identify bacterial taxa that correlate with EAE severity, and as such has the potential to modulate CNS inflammation. We found that vancomycin treatment ameliorates EAE, and that this protective effect is mediated via the gut microbiota. Notably, we observed increased abundance of Tregs inducing bacteria including members of Clostridium clusters XIVa and XVIII in the gut of vancomycin-treated mice compared to untreated mice. Importantly, many of the bacteria that correlated with EAE severity exist in the human gut microbiome. In addition, several of the bacteria that have been implicated in MS such as A. muciniphila and Anaerotruncus colihominis correlated with EAE severity. We found that human-derived A. colihominis ameliorates EAE, and this is associated with the induction of RORγt+ regulatory T cells in the mesenteric lymph nodes. Our work led to the identification of 50 bacterial species with neuroimmunomodulatory potential. This list of bacteria can serve as a starting point to define pathogenic and beneficial host-microbiota interactions that modulate neuroinflammation in MS. This information will in turn facilitate the development of innovative microbiota-based approaches to target neuroinflammation in MS.

Materials and methods

Mice
C57BL/6J female mice were obtained from the Jackson Laboratory and kept in a specific pathogen-free facility at the Harvard Institute of Medicine or Icahn School of Medicine at Mount Sinai on a 12-h light/dark cycle. C57BL/6J germ-free female mice were obtained from the Massachusetts Host-Microbiome Center at the Brigham & Women’s Hospital and kept in a specific pathogen-free facility at the Harvard Microbiome Facility or the Massachusetts Host-Microbiome Center at the Brigham & Women’s Hospital. Mice were all 8–10 weeks of age and cohoused four mice from the same experimental condition per cage. Mice were assigned randomly to the experimental groups. Mice were fed an ad libitum diet of PicoLab Rodent Diet 5053 and distilled water without added preservatives (provided by animal facility). Animals were housed in a biosafety level 2 facility using autoclaved cages and aseptic handling procedures and kept under a 12-h light/dark cycle. All animal experiments described in this paper were approved by the Institutional Animal Care and Use Committee (IACUC) at Harvard Medical School or Icahn School of Medicine at Mount Sinai and carried out in accordance with those approved animal experiment guidelines.

EAE induction

EAE was induced by injecting 8- to 10-week-old female C57BL/6J mice with 150 μg MOG35–55 peptide (Genemed Synthesis) emulsified in complete Freund’s adjuvant (CFA) (BD Difco) per mouse subcutaneously in the flanks, followed by intraperitoneal administration of 150 ng pertussis toxin (List Biological Laboratories, Inc.) per mouse on days 0 and 2 as previously described [60]. Clinical signs of EAE were assessed according to the following score: 0, no signs of disease; 1, loss of tone in the tail; 2, hind limb paresis; 3, hind limb paralysis; 4, tetraplegia; and 5, moribund. Differences between the groups were determined by Friedman test and Dunn correction for multiple comparisons.

Isolation and identification of Enterococcus faecalis

Adult C57BL6/J female mice received vancomycin (0.5 g/L) in drinking water for 2 weeks. Next, mice were euthanized, cecum was rapidly collected and resuspended in prereduced anaerobically sterilized saline, and 100 μL of 10−4 through 10−7 dilutions was plated on Brucella blood agar (Anaerobe Systems) and incubated in an aerobic incubator. Eight to ten colonies were isolated in pure culture after 48 h incubation. Pure Enterococcus faecalis cultures were identified by biotyping using the Bruker Biotyper LT MALDI-TOF mass spectrometer. The E. faecalis ID was confirmed by amplifying the nearly full-length 16S rRNA gene using the 8F and 1510R primers according to previous methods [61]. After polymerase chain reaction, PCR products were sequenced by Sanger sequencing at GENEWIZ. Identification and percent identity were then performed using batch BLAST, National Centers for Biotechnology Information.

Bacteria strains, growth, and administration

Anaerotruncus colihominis (DSMZ, DSM no.:17241) and Enterococcus faecalis were grown anaerobically at 37 °C in brain-heart infusion (BHI) medium (Cat. no.: DF0037178; Fisher Scientific). Each bacterial culture suspension was subsequently transferred to 3 Brucella blood agar plates, and colonies from all plates were resuspended in 2.5 mL anaerobic sterile PBS yielding a suspension of live bacteria at a density of OD600 = 1.3. An 8-week-old female C57BL/6J mice were colonized with 200 μL of bacteria suspension via oral gavage 3 days per week beginning 3 weeks prior to disease induction, and bacteria treatment was received for the entire duration of the experiment. Control mice were orally gavaged with anaerobic PBS (vehicle).

Histopathology

Mice were euthanized at the termination of experiments and were intracardially perfused with PBS. Lumbosacral spinal cords were fixed with formalin. Tissue was processed and stained as previously described [60]. Paraffin-embedded serial sections were stained with
Luxol fast blue for myelin and Bielschowsky silver for axons. The demyelinated area and axonal/neuronal loss were determined using ImageJ software (National Institutes of Health, USA), and the percentages of demyelinated and axonal/neuronal lost area out of total area were calculated. To detect immune infiltrate, spinal cord sections were stained using Rat Anti-Mouse CD4 antibody (1:100; eBioscience, Cat. no. 14-0042-82) with secondary biotinylated antibodies. Avidin-peroxidase and 3,4-diaminobenzidine were used as the color substrate. CD4+ cells were counted manually using multipoint function on ImageJ and cell density calculated as number of cells per total white matter area.

**Antibiotic treatment**
In order to investigate the effect of vancomycin and neomycin on EAE development, mice were given vancomycin 0.5 mg/mL or neomycin 1 mg/mL (Fisher Scientific) in drinking water for 2 weeks (Fig. 2A). For the cohousing experiment, mice received vancomycin 3 mg in 200 μL nuclease-free water via oral gavage (Fig. 2E). For the fecal transfer experiment using wild-type C57BL/6J mice, to deplete bacteria, mice were given a mixture of antibiotics (ampicillin 1 mg/mL, vancomycin 0.5 mg/mL, neomycin 1 mg/mL, metronidazole 1 mg/mL; Fisher Scientific) in drinking water for 3 consecutive days (Fig. 2G).

**Fecal microbiota transplantation**

**Feces collection, processing, and storage**
Donor mice received vancomycin 0.5 mg/mL in drinking water or normal drinking water for 2 weeks. Two weeks post vancomycin initiation, ceca were removed under aseptic conditions and transferred to an anaerobic chamber (Anaerobe Systems) for processing. The content of 3 ceca from control mice or 1 cecum from vancomycin treated mice were homogenized by vortexing, at room temperature, in 8 mL of pre-reduced and anaerobically sterilized PBS (Anaerobe Systems Cat. no. AS-908). For experiments assessing the effect of control/vancomycin microbiota on germ-free mice (Fig. 2I–L), recipient animals were treated with antibiotics for 3 days as described above. Twenty-four hour post discontinuation of antibiotics, mice (n = 10/group) were administered 200 μL of the control- or vancomycin cecal slurry by oral gavage at −11 and −10 days post immunization with MOG.

**16S rRNA gene sequencing and analysis**
Feces were collected at 11 time points spanning pre- and postimmunization states: before vancomycin treatment (day −22), on vancomycin (days −21, −20, −18, −14), at vancomycin discontinuation (day −6), post vancomycin discontinuation/pre-EAE induction (day 0), post EAE induction-latent period (days 3 and 8), post EAE induction-peak disease (day 15), and post EAE induction-recovery phase (day 29). Fecal samples were collected immediately upon defecation, snap-frozen, and kept at −80 °C until processed. Bacterial DNA was isolated from feces using MoBio PowerLyzer PowerSoil Kit (Qiagen). Amplicons spanning variable region 4 (V4) of the bacterial 16S rRNA gene were generated with primers containing barcodes (515F, 806R) from the Earth Microbiome Project [62] using HotMaster Taq and HotMaster Mix (QuantaBio) and paired-end sequenced on an Illumina MiSeq platform at the Harvard Medical School Biopolymer Facility. Paired-end 16S rRNA gene reads were trimmed for quality (target error rate < 0.5%) and length (minimum 125 bp) using Trimmomatic, merged using FLASH (fast length adjustment of short reads), and quality screened using QIME 1 (quantitative insights into microbial ecology). Spurious hits to the Phix control genome were identified using BLASTN (basic local alignment search tool) and removed. Passing sequences were trimmed of primers, evaluated for chimeras, and screened for mouse-associated contaminant using Bowtie2. True 16S rRNA sequence fragments were confirmed as such by requiring at least one BLAST match against the Greengenes database (default parameters).
Greengenes was only used for this QC step, and not for taxonomic assignment. Chloroplast and mitochondrial contaminants were detected and filtered using the RDP (Ribosomal Database Project) classifier with a confidence
threshold of 80%. High-quality 16S rRNA amplicon sequences were assigned to a high-resolution taxonomic lineage using Resphera Insight (v2.2), for which the performance has been previously benchmarked using high-quality draft genome assemblies of well-defined species from the Human Microbiome Reference Genome Database [http://hmpdacc.org/HMRGD/; ref. 13]. This method utilizes a manually curated 16S rRNA database of 11,000 unique species and a hybrid global-local alignment procedure to assign short next-generation sequencing sequences from any region of the 16S rRNA gene to a high-resolution taxonomic lineage [63]. Operational taxonomic units (OTUs) assignment was performed as follows: sequences that could not be confidently assigned to any known species were clustered de novo, and clusters were given an OTU number identifier and annotated with a closest relative or set of relatives as previously described [64]. Taxonomy summary plots were generated based on relative abundances as implemented in QIIME. Testing for significant differences in alpha diversity was performed by one-way ANOVA followed by Tukey’s post hoc test or a mixed-effects model when comparing matched data with a missing value followed by Dunnett’s corrections (as indicated on the figure legends). Beta diversity was estimated using Bray-Curtis, and distances were then used to perform principal coordinate analysis as implemented in QIIME. Differences in beta diversity were tested using PERMANOVA with FDR correction for multiple comparison testing. Compositional differences were determined using linear discriminant analysis effect size (LEfSe) with alpha set at 0.05 and the effect size set at greater than 2. To identify bacteria linked with EAE severity, Spearman correlations were performed in R using the function cor.test, and FDR adjustment was performed in R using the function p.adjust.

**Fecal microbe quantification by qPCR**

DNA extracted from feces as described above was used for specific bacteria abundance. Quantitative PCR (qPCR) analysis was conducted using a QuantStudio 7 Flex Real-Time PCR System (Applied Biosystems). *A. colihominis*, *E. faecalis*, and *SFB* were quantified by SYBR Green (Applied Biosystems) and primer pairs as follows: all bacteria (universal 16S rRNA gene, reference): forward: ACT CCT ACG GGA GGC AGC AGT, reverse: ATT GCT AAG ACC CCG GAT TGA TGT TAT [65]; *A. colihominis* 16S rRNA gene: forward: GGAGCTTAGGTITGAAGTTTCC, reverse: CTGCTGCTTCCCGTA [66]; *E. faecalis* 16S rRNA gene: forward: TACTGAAACACATTGATG, reverse: AACTTCTGCACCAACCGGCAAC [67]; and *SFB* 16S rRNA gene: forward: GACGCTAGGGC TGAGAGCAT, reverse: GACGGCACGGATTGTTAT TCA [65]. The relative quantity was calculated using the comparative CT method normalizing to the amount of all bacteria in the sample [68].

**Immune cell isolation**

For immune profiling, naive 8-week-old female C57BL/6 mice were orally gavaged with *A. colihominis* or *E. faecalis* or PBS 3 days per week for 3 weeks as described above after which mice were euthanized using carbon dioxide and tissues rapidly dissected. Single-cell suspensions were obtained as previously described [69]. Mesenteric lymph nodes and spleens were dissociated by pressing through 70 μm mesh and red blood cells removed from splenocytes using ALK lysis buffer. Intestinal tissues were cleaned of feces, Peyer’s patches removed, and deepithelialized in 5 mM buffered EDTA before digestion with 0.5 mg/ml collagenase type 4 (Sigma C5138) and 0.5 mg/ml DNase1 (Sigma DN25). Cell suspensions were filtered through 40 μm strainers and washed before use. No further enrichment of lymphocytes was performed.

**Flow cytometry**

For analysis of cytokine production, mononuclear cell suspensions were restimulated with 5 ng/mL phorbol 12-myristate 13-acetate (PMA) and 500 ng/mL ionomycin with monensin (Biolegend) for 3.5 h at 37 °C. All other analysis was performed on unstimulated cells. Intracellular cytokine staining was performed following surface staining and fixation with IC Fixation Buffer (ThermoFisher/eBioscience). Transcription factor staining was performed using FoxP3 fixation/permeabilization buffers (ThermoFisher/eBioscience). Super Bright Complete staining buffer (ThermoFisher) was included when multiple Brilliant Violet-conjugated antibodies were used together. The following anti-mouse antigen fluorochrome-conjugated antibodies were used, obtained from BioLegend unless otherwise stated: CD45 Brilliant Violet 750 (30-F11), CD45 APC-Cy7 (30-F11), CD4 FITC (RM4-5), CD4 PerCp-Cy5.5 (RM4-5), CD3 AlexaFluor 700 (17A2), CD3 PerCP-eFluor710 (17A2, Thermofisher/ eBioscience), CD8-alpha Pacific Orange (5H10, ThermoFisher), CD11b PerCP-Cy5.5 (M1/70), CD11c PE-Cy7 (N418), CD103 Brilliant Violet 510 (2E7), CD64 Brilliant Violet 786 (X54-5/7.1, BD Bioscience), MHC-II I-A/I-E Pacific Blue (M5/114.15.2), CD80 Brilliant Violet 421 (16-10A1), CD86 Brilliant Violet 605 (GL-1), Ly-6G Brilliant Violet 570 (1A8), FoxP3 PE (FJK-16s, Thermofisher/eBioscience), RORγt APC (B2D, BD BioScience), GATA3 Brilliant Violet 421 (16E10A23), IL-17A PE (TC11-18H10.1), IL-10 APC (JES5-16E3), IFNy PE-Cy7 (XMG1.2), and GM-CSF FITC (MP1-22E9). Dead cells were excluded using Zombie Aqua (BioLegend) or eFluor780 Fixable...
Viability Dye (ThermoFisher/eBioscience). Data was acquired on a five-laser Aurora Cytometer (Cytek Biosciences), and raw data was spectrally unmixed using SpectroFlo software (Cytek Biosciences). Unmixed data files were analyzed using FlowJo 10 (BD Biosciences) and statistical analyses performed using RStudio 1.1.463 and Prism 6 (GraphPad).

Quantification and statistical analysis
All graphs, calculations, and statistical analyses were performed using the GraphPad Prism software for Mac (GraphPad Software, San Diego, CA, USA). Comparisons of three or more groups following a normal distribution were performed using one-way ANOVA followed by Dunnett’s test or Tukey’s test as specified in figures. For datasets of non-normal distributions, the Mann-Whitney test was used for comparisons between two groups. EAE clinical scores were analyzed over time with the non-parametric Friedman test for repeated measurements as specified in figures [70, 71]. Exact statistical instruments, sample sizes, and P-values are indicated in each figure. We did not use statistical methods to determine sample size; we used sample sizes that were similar to those in our previous publications and those of others [60, 72–75].

Results
Characterization of the gut microbiome during EAE
To investigate changes in the gut microbiota composition during EAE, we collected fecal pellets at 11 time points spanning pre- and post-MOG immunization state from an 8-week-old C57BL/6 (B6) mice (Fig. 1A and B). Next, we analyzed the microbiota from these mice by sequencing the V4 region of the microbial 16S rRNA gene. We first assessed alpha diversity and found no significant changes at any time point (Figs. 1C, S1A). We then examined beta diversity to assess whether the composition of the microbiota changes during EAE and found no significant differences in community structure between pre- and postimmunization states (Fig. 1D). We next investigated whether the relative abundances of bacteria differed between pre- and postimmunization states at the species level. We observed a change in the abundance of species belonging to several genera including Akkermania, Turvibacter, Lactobacillus, and Clostridium (Figs. 1E, S1B). We observed that most changes in taxa abundance occurred within the first 2 weeks post EAE induction (Figs. 1F, S2). In particular, we found increased levels of Clostridium cocleatum, Ruminococcus flavefaciens, Clostridium ruminantium, and Clostridium chauvoei at 3 days post immunization (DPI) compared to 0 DPI (Figs. 1F, S2). We observed increased abundance of 8 taxa including Clostridium scindens, a member of Clostridium cluster XIVa as well as three bacteria belonging to Clostridium sensu stricto: C. chauvoei/quinii operational taxonomic unit (OTU) 8778, C. chauvoei, and C. celatum at 8 DPI compared to 0 DPI (Figs. 1F–G, S2). Family Verrucomicrobiaceae which contains species Akkermanisia muciniphila were enriched in the gut of mice at 15 DPI compared to 0 DPI. These findings are consistent with prior studies reporting increased A. muciniphila in EAE mice [47, 76, 77]. In addition, several studies have reported increased level of A. muciniphila in MS compare to healthy control [11, 14, 15, 40, 46]. C. scindens was also increased at 15 DPI compared to 0 DPI (Figs. 1F–G, S2). Interestingly, we have recently reported that C. scindens is increased in MS patients compare to healthy control [40]. Two Clostridia species were decreased at 3 DPI compared to 0 DPI: Clostridium viride (OTU 5938), a member of Clostridium cluster IV, and Clostridium indolis (OTU 1977), a member of Clostridium cluster XIVa (Figs. 1F–G, S2). Furthermore, several species belonging to Clostridium cluster XIVa were decreased at 15 DPI compared to 0 DPI: C. boltea/clostridiform/oroticum (OTU 8033), C. polysaccharolyticum (OTU 7948), C. indolis (OTU 1977), C. hylemonae, and C. aldenense/indolis (Figs. 1F–G, S2). These results are consistent with findings from one study reporting decreased abundance of species belonging to Clostridium clusters IV and XIVa in MS patients [41]. We also found decreased Dorea formicigenerans (OTU 2094) and Lactobacillus gasseri/hominis/johnsonii/taiwanensis at 3 DPI compared to 0 DPI (Figs. 1F–G, S2). These findings are consistent with results from prior studies reporting decreased genera Dorea and lactobacilli as well as decreased L. johnsonii/
Fig. 1  (See legend on previous page.)
Amelioration of EAE after vancomycin is mediated via the microbiota

To this date, it remains unclear if the neuroprotective effects of antibiotics in EAE mice are driven by changes in the gut microbiota composition. To investigate this, 8-week-old B6 mice were treated with vancomycin or neomycin, two poorly absorbable antibiotics, for 2 weeks, after which antibiotics were discontinued (Fig. 2A). Next, mice were immunized with MOG for EAE induction. We found that, compared to control mice, vancomycin-treated mice had significantly less severe disease, whereas neomycin-treated mice behaved like the control group (Fig. 2B–D). Hence, these results are consistent with our prior report that vancomycin treatment ameliorates EAE [78]. To determine if the protective effect of vancomycin is mediated via the microbiota, we conducted three follow-up experiments. First, we did a cohousing experiment in which 8-week-old B6 mice receiving normal drinking water or vancomycin once a day via oral gavage were either housed with the same treatment group (single treatment) or cohoused between treatment groups. Two weeks later, vancomycin was discontinued, and mice were immunized with MOG to induce EAE (Fig. 2E). As expected, single-treatment housed control mice developed more severe disease than the single-treatment housed vancomycin mice (Fig. 2F). We observed that vancomycin-treated mice that were cohoused with control mice developed more severe disease than single-treatment housed vancomycin mice (Fig. 2F). We also found that control mice that were cohoused with vancomycin-treated mice had delayed disease onset compared to single-treatment housed control mice (Fig. 2F).

Second, we performed a cecal transfer experiment where 8-week-old conventionally raised B6 mice were treated with an antibiotic mixture containing neomycin, vancomycin, ampicillin, and metronidazole for 3 days. Twenty-four hours after antibiotics discontinuation, half of these mice were fed feces from control mice, and the remaining half received feces from vancomycin-treated mice. Ten days post gavage, EAE was induced, and EAE scores were monitored overtime (Fig. 2G). We found that mice that received feces from vancomycin-treated mice had less severe disease than mice who were fed feces from control mice (Fig. 2H). Third, we conducted a second cecal transfer experiment, this time using 4-week-old B6 germ-free (GF) mice that were either gavaged with feces from control mice or with feces from vancomycin-treated mice. At 4-week post gavage, EAE was induced, and EAE scores were monitored overtime (Fig. 2I). We observed that mice that were fed feces from vancomycin-treated mice had less severe EAE compared to those that received feces from control mice (Fig. 2J–L). Taken together, these results suggest that the vancomycin-protective effect is mediated via the microbiota.
Fig. 2 (See legend on previous page.)
mediated via the microbiota. Segmented filamentous bacteria (SFB) exacerbates EAE via induction of Th17 cells, and given that SFB is sensitive to vancomycin, we repeated the fecal transfer experiment in GF mice using wild-type B6 mice microbiota from the Jackson Laboratory which has been previously shown to be free of SFB [29]. We confirmed that the microbiota from our B6 mice lacks SFB by quantitative real-time PCR using SFB-specific primers (Fig. S3A–B). Next, 4-week-old GF mice were gavaged with SFB-free microbiota, and 3 weeks later, half of the mice were treated with vancomycin via oral gavage once daily for 2 weeks (Fig. S3C). We found that vancomycin treatment protects mice from EAE even in the absence of SFB (Fig. S3D–F). Hence, the protective effect of vancomycin is not modulated by SFB.

**Vancomycin effect on the gut microbiota during EAE**

Given that vancomycin-treated mice had less severe EAE and we found that this protective effect is mediated via the microbiota, we next profiled the 16S rRNA gene of these mice. Examining alpha diversity, as expected, we found that vancomycin-treated mice had decreased Shannon diversity, Observed species, and Faith’s diversity compared to control mice (Fig. 3A). However, starting at 3 DPI, we found no change in alpha diversity between vancomycin-treated mice that were cohoused with control mice and single-treatment housed control mice (Fig. 3A). In addition, starting at 0 DPI, cohoused vancomycin-treated mice had increased Shannon diversity compared to single-treatment housed vancomycin mice (Fig. 3A). We also observed increased Observed species and Faith’s diversity in cohoused vancomycin-treated mice compared to single-treatment housed vancomycin mice starting at 0 DPI (Fig. 3A). Examining beta diversity, the gut microbiota composition was not significantly different among the 4 mice groups at −22 DPI (baseline) prior to vancomycin initiation (q > 0.29). We observed that overall microbiota community structure of single-treatment housed vancomycin mice differed from that of single-treatment housed control mice (q < 0.003; Fig. 3B) as well as that of cohoused control mice (q < 0.002; Fig. 3B) starting at −21 DPI (1 day post vancomycin initiation). At −21 DPI, single housed vancomycin-treated mice clustered with cohoused vancomycin mice (q = 0.192; Fig. 3B). However, starting at −6 DPI, single-treatment housed vancomycin mice clustered differently from cohoused vancomycin mice (q < 0.002; Fig. 3B). Hence, despite receiving the same vancomycin treatment, single-treatment housed vancomycin mice and cohoused vancomycin treated mice have different alpha diversity and microbial composition. In addition, the overall microbiota community structure differed between cohoused vancomycin-treated mice and single-treatment housed control mice at all time points (q < 0.002; Fig. 3B) and between cohoused vancomycin-treated mice and cohoused control mice at −21, −6, and 0 DPI (q < 0.005; Fig. 3B). However, starting at 3 DPI, cohoused vancomycin-treated mice clustered with cohoused control mice (q > 0.05; Fig. 3B).

To identify taxa that account for the difference in beta diversity in these mice, we generated a taxa plot. As expected, we observed a change in the abundance of several bacteria between vancomycin-treated and untreated mice including *A. muciniphila*, *T. sanguinis*, *Lactobacillus*, *Enterococcus*, and *Clostridium* species (Fig. 3C). We found several taxa that are increased in single-treatment housed vancomycin mice compared to single-treatment housed control mice or cohoused vancomycin-treated mice including the following: *A. muciniphila*, *T. sanguinis*, *Lactobacillus* gasseri/hominis/johnsonii/taiwanensis, and several *Clostridium* species belonging to Clostridium cluster XIVa and *Clostridium* sensu stricto (Figs. 4A–B, S4A–B). These findings are consistent with our previous report that *A. muciniphila* ameliorates EAE by inducing Tregs [47] as well as several studies observing that administering various lactobacilli strains to mice leads to an attenuation of EAE clinical course by either inducing Tregs or suppressing Th17 cells induction [79–81]. We also found that single-treatment housed vancomycin mice had decreased level of most species belonging to Clostridium cluster XIVa that we were able to detect during the pre-symptomatic stages of EAE (0, (See figure on next page.)

**Fig. 3** Effect of vancomycin on alpha and beta diversity during EAE. A Alpha diversity metrics for Shannon diversity, Observed species, and Faith’s diversity were calculated at an average sampling depth of 1500 reads per sample in single-treatment housed untreated mice (control), single-treatment housed vancomycin mice (vancomycin), untreated mice cohoused with vancomycin-treated mice (ControlCoho), and vancomycin-treated mice cohoused with untreated mice (VancoCoho). Results are expressed as mean ± SEM (n = 11–12 mice/group). Statistical comparisons are made using one-way ANOVA followed by Tukey test. *Significant difference with control group (p < 0.05), +significant difference with vancomycin group (p < 0.05), #significant difference with control-cohoused group (p < 0.05). B Principal coordinate analysis of intestinal microbiota samples based on Bray-Curtis shows significantly different clustering between control and vancomycin (q < 0.003), ControlCoho and vancomycin (q < 0.002), VancoCoho and control (q < 0.002) at all time points, vancomycin and VancoCoho (q < 0.002), control and ControlCoho (q < 0.002) starting at −6 DPI, and VancoCoho and ControlCoho (q < 0.005) at −21, −6, and 0 DPI but not at 3, 8, 15, and 29 DPI. q = PERMANOVA p-values adjusted for false discovery rate. Each dot represents the microbiota from one mouse. C Taxa plots showing compositional differences in fecal microbiota at the indicated time points in control, vancomycin, ControlCoho, and VancoCoho mice. EC1, *Enterococcus canintestini/canis/dispar/durans/faecalis/faecium/hiraec/factis/mundhi/ratti/rivorum/villorum*
**Fig. 3** (See legend on previous page.)
3, and 8 DPI) (Fig. S4A). However, at peak disease (15 DPI), we observed an increase of four species belonging to Clostridium cluster XIVa in single-treatment housed vancomycin mice compared to single-treatment housed control mice including \textit{Clostridium scindens} (OTU 2213), \textit{Clostridium asparagiforme/lavalense} (OTUs 1808 and 2825), and \textit{Clostridium indolis/jejuense} (OTU 6292) (Figs. 4A, S4A). In addition, \textit{Clostridium cocleatum}, a member of Clostridium cluster XVIII, was increased in single-treatment housed vancomycin mice compared to single-treatment housed control mice or cohoused vancomycin mice (Figs. 4A–B, S4A–B). Interestingly, species
that fall within clusters XIVa and XVIII of *Clostridia* are strong Tregs inducers [32, 82, 83]. When comparing the microbiota composition of vancomycin-treated mice and control mice that were cohoused, *A. muciniphila* was increased in cohoused vancomycin-treated mice compared to cohoused control mice (Figs. 4C, 5A). We also observed that *C. indolis/jejuense* (OTU 6292) and *C. cocleatum* (OTU 2322), members of Clostridium cluster XIVa, as well as *C. coeleatum* were increased in cohoused vancomycin-treated mice compared to cohoused control mice (Figs. 4C, 5A). Furthermore, *Caprobacillus cateniformis* (OTU 4628), a butyrate producer [84], was increased in cohoused vancomycin-treated mice compared to cohoused control mice (Figs. 4C, 5A). Interestingly, butyrate is a Tregs inducer that has been shown to ameliorate EAE [85, 86]. Taken together, these findings suggest that a potential mechanism by which vancomycin treatment attenuates EAE severity is via promoting the proliferation of bacterial species that are Tregs inducers. These results also suggest that single-treatment housed vancomycin mice had less severe disease than cohoused vancomycin-treated mice because they had more Tregs inducing bacterial species. Several bacteria were decreased in single-treatment housed vancomycin mice compared to single-treatment housed control mice or cohoused vancomycin-treated mice. Notably, *Anaerotruncus colihominis* (OTU 2694) was depleted in single-treatment housed vancomycin mice (Fig. 4A–B, S4 A–B). Interestingly, we and others have previously reported increased *A. colihominis* in MS patients compared to healthy subjects [19, 40].

**Correlation between bacterial abundance and EAE severity**

To identify gut commensals that are associated with EAE severity, we performed Spearman correlation analysis. We identified 36 bacteria that positively correlate with EAE severity, we performed Spearman correlation analysis. We identified 36 bacteria that positively correlate with EAE severity (Fig. 5A–B). These findings are consistent with prior studies reporting that some *Lactobacillus* species/strains ameliorate EAE while others exacerbate EAE [79–81, 88]. The majority of bacteria found to have a positive correlation with EAE severity have not been studied in EAE yet including *Dorea formicigenerans*, *Anaerospirabacter*, *Robinsoniella*, *Flavonifractor*, *Papillibacter*, *Sporobacter*, and *Anaerotruncus colihominis* (Fig. 5A). We identified 13 bacteria that negatively correlated with EAE severity and were enriched in vancomycin-treated mice (Fig. 5A). We found that *A. muciniphila* negatively correlated with EAE severity (Fig. 5A–B). Consistent with these findings, we previously reported that mice colonized with MS-derived or commercially obtained *A. muciniphila* are protected from EAE [40, 47]. Three species belonging to Clostridium sensu stricto: *C. chauvoei*, *C. celatum*, and *C. chauvoei/quinii* (OTU 8778) negatively correlated with EAE severity (Fig. 5A). These findings are consistent with prior studies reporting a negative correlation between OTUs belonging to Clostridium sensu stricto and EAE cumulative disease score [87]. Several genera that negatively correlated with EAE severity have not yet been investigated in the EAE mice model including *Turicibacter sanguinis*, *Clostridium cocoletum*, *Olsenella profusa*, and *Enterococcus* (Fig. 5A).

*A. colihominis* ameliorates EAE and induces RORyt+ regulatory T cells in the mesenteric lymph nodes

To this date, no human gut-derived bacteria have been shown to exacerbate EAE. Given prior report that *A. colihominis* is increased in the gut of MS patients and given that it is depleted in mice protected from EAE, we hypothesized that *A. colihominis* might be an important driver of the pro-inflammatory response in EAE/MS. However, *A. colihominis* is a butyrate-producing bacteria [89, 90] which could potentially play a beneficial role.
Fig. 5 (See legend on previous page.)
given prior reports that butyrate ameliorates EAE via induction of Tregs [85, 86]. Thus, whether Anaerotrun\-cupus contributes to disease or could protect against CNS autoimmunity is needed to be tested experimentally. To investigate this, B6 mice were fed A. colihominis, Enterococcus faecalis, a commensal gut microbe not altered in EAE mice, or phosphate-buffered saline (PBS), 3 days per week for the entire duration of the experiment to maintain steady level of the bacteria (Fig. 6A). To confirm that mice were successfully colonized with A. colihominis and E. faecalis, feces were collected from mice 3 weeks post initiation of gavage, fecal DNA was extracted, and bacterial abundance was assessed via quantitative PCR (Fig. 6B). Three weeks post initiation of gavage, EAE was induced by immunizing mice with MOG. We found that mice fed A. colihominis had less severe disease than mice fed E. faecalis or PBS (Fig. 6C). Hence, A. colihominis ameliorates EAE, whereas mice fed E. faecalis developed slightly more severe disease than PBS-treated mice. These findings support a beneficial role for A. colihominis in EAE/MS.

To examine the effect of A. colihominis and E. faecalis on gut and peripheral immunity, naive B6 mice were fed A. colihominis, E. faecalis, or PBS three times per week for 3 weeks. Mice were then sacrificed, and lamina propria, mesenteric lymph nodes, and spleen were collected for immune profiling. We observed increased frequency of Th17 cells (CD3+CD4+RORγt+) in the colon of mice fed E. faecalis compared to mice who received PBS or A. colihominis (Figs. S5A, 6D–E). No difference in the frequency of Th17 cells (CD3+CD4+RORγt+) was seen in the ileum or mLN of these mice (Fig. 6D–E). We also observed no difference in level of IL17A, IFNγ, or IL10 in the gut or spleen of these mice (Fig. S5B–D). GMCSF was increased in the colon of A. colihominis fed mice compared to E. faecalis or PBS-treated mice, but no difference in GMCSF level was observed in the ileum, mLN, or spleen of these mice (Fig. S5E). These findings suggest that E. faecalis may exacerbate EAE via induction of Th17 cells in the colon. Consistent with this notion, one study reported that blocking encephalitogenic Th17 cell entry into the colon ameliorates EAE [91]. Furthermore, another study found a positive correlation between frequency of intestinal Th17 cells and disease activity in MS [57].

Next, we looked at the frequency of T regulatory cells (CD4+Foxp3+) in the gut of these mice and found no difference in the frequency of these cells in the ileum and mLN (Fig. 6F–G). There was a trend toward increase frequency of T regulatory cells (CD4+Foxp3+) in the colon of mice fed A. colihominis compared to mice fed E. fae- calis or PBS (Fig. 6F–G). We observed that the Treg:Th17 ratio was trending up in the colon and ileum of B6 mice fed A. colihominis compared to mice fed E. faecalis or PBS (Fig. 6H). We found increased Treg:Th17 ratio in the mLN of PBS fed mice compare to mice fed E. fae- calis and A. colihominis (Fig. 6H). A study reported that peripheral RORγt+ regulatory T cells suppress myelin-specific Th17 cell-mediated CNS auto-inflammation in a passive EAE model [92]. Hence, we investigated the effect of A. colihominis and E. faecalis on the frequency of peripheral RORγt+ regulatory T cells. We found no difference in the frequency of peripheral RORγt+ regulatory T cells in the spleen of these mice (Fig. S5F). Studies have demonstrated that intestinal RORγt+ Treg cells are microbiota dependent, enriched in the gut, and have a strongly suppressive capacity during intestinal inflammation [93, 94]. Hence, we next examined the frequency of intestinal RORγt+ Treg in these mice. The frequency of RORγt+ Treg was comparable in all 3 groups in the colon and ileum (Fig. 6I–J). We observed increased frequency of RORγt+ regulatory T cells in the mLN of mice fed A. colihominis compared to mice fed E. faecalis or PBS (Fig. 6I–J). These findings are consistent with other studies reporting that a selected mixture of Clostridia strains from the human microbiota including A. colihominis
Fig. 6 (See legend on previous page.)
induces T regulatory cells in mouse colonic lamina propria [32, 82]. Taken together, our findings suggest that elevated *A. colihominis* in MS patients may be a host-triggered response to suppress neuroinflammation. Dendritic cells from the mesenteric lymph node (mLN) are known to play a crucial role in Treg induction [95–97]. Hence, we next examined the frequency of tolerogenic dendritic cells in the mLN of these mice. We found no difference in the frequency of tolerogenic dendritic cells (CD11c+CD11b+CD103+) in the mLN of these mice (Fig. S6A–C). Hence, *A. colihominis* does not induce tolerogenic dendritic cells. Furthermore, the frequency of activated dendritic cells (CD11c+CD103−) was comparable in all 3 groups in the mLN and spleen (Fig. S6D–E).

### Discussion

Several studies have identified alterations in the gut microbiome of MS patients, and it is now accepted that the microbiome plays an important role in the pathophysiology of MS. Nevertheless, given that the mechanisms by which the microbiome affects MS have not been well defined and many confounding factors exist, findings from human microbiome studies must be validated well defined and many confounding factors exist, findings from human microbiome studies must be validated. Perturbations of the gut microbiota composition of mice have been shown to influence EAE development. Yet, few studies have characterized the microbiome during EAE and link changes in the EAE microbiome to disease severity. Hence, here we investigated changes in the gut microbiota of untreated and vancomycin-treated EAE mice at multiple time points before and after EAE induction to identify gut commensals with neuro-immunomodulatory potential.

We found that vancomycin treatment ameliorates EAE, and this protective effect is mediated via the microbiota. We observed an enrichment of Tregs inducing bacterial species in the gut of vancomycin-treated mice including members of Clostridium cluster XIVa. We next investigated the microbiome of untreated and vancomycin-treated EAE mice to identify taxa that regulate neuroinflammation. We observed a negative correlation between *A. muciniphila* and EAE severity. Interestingly, we and others have reported increased *A. muciniphila* in MS patient compared to healthy control (HC) [11, 14, 15, 40]. In a recent study, we observed that *Akkermansia* negatively correlated with disability and T2 lesion volume and positively correlated with brain volume. These findings are consistent with our results showing that MS-derived *Akkermansia* ameliorates EAE, and this protective effect is associated with decreased in RORγt+ and IL-17+ producing γδ T cells [40]. Consistent with our findings in the B6 model, another study found that mice with higher levels of *Akkermansia* had less progression in the nonobese diabetic (NOD) progressive EAE model [98]. Taken together, our findings suggest that increased *Akkermansia* could be a beneficial compensatory microbiome response in EAE and MS.

We also found a negative correlation between level of *Lactobacillus* and EAE severity at 15 DPI. These findings are consistent with prior studies reporting that daily administration of a mixture of *Lactobacillus* strains in EAE was effective both at preventing disease development and reversing established disease, an outcome that was IL-10 dependent and correlated with Tregs induction in mesenteric lymph nodes and the CNS [79]. Similarly, treatment with a mixture of *Lactobacillus plantarum* of human origin and *Bifidobacterium animalis* attenuated EAE clinically and induced Tregs in lymph nodes and spleen [99]. Oral administration of human-derived *L. reuteri* after immunization ameliorated EAE with decreased in Th1/Th17 subsets and related cytokines [80]. In humans, one study report decreased *Lactobacillus* in MS patients [12]. We have previously reported that administering a probiotic mixture containing four lactobacilli species to MS patients twice daily for 2 months leads to reduce pro-inflammatory markers in peripheral monocytes [58]. Consistent with our findings, two small double-blinded randomized controlled trials in MS patients receiving a mix of *Lactobacillus* and *Bifidobacterium* daily for 12 weeks showed significant improvements in disability score, depression, anxiety, and inflammatory markers including reduced IL-8 and TNF-alpha expression in peripheral blood mononuclear cells (PBMCs) [59, 100]. Hence, these results suggest that decreased *Lactobacillus* levels in the gut of EAE mice and MS patients could promote CNS inflammation.

We observed a negative correlation between EAE severity and abundance of *Turicibacter sanguinis*, *Olsenella*, and 3 species belonging to *Clostridium* sensu stricto: *C. chauvoei/quinii* (OTU 8778), *C. chauvoei*, and *C. celatum*. Interestingly, we have previously reported elevated *T. sanguinis* and *Olsenella* in MS [40]. Another study reports increased *Clostridium* sensu stricto in MS patients, but they did not identify these taxa at the species level [101]. Little is known about the functions of these bacteria in neurologic diseases. Hence, more studies are needed to investigate the role of these bacteria in EAE/MS. Several other genera that positively correlated with EAE severity have neither been implicated in MS nor have they been investigated in EAE mice including *Flavonifractor, Anaeroспорobacter*, and *Papillibacter*.

We found a positive correlation between EAE severity and 6 taxa belonging to Clostridium cluster XIVa including *C. indolis* (OTU 1977) and *C. jejuna* (OTU 7388). Interestingly, we have reported elevated *C. scindens* in...
MS patients [40]. Another study reported decreased species belonging to Clostridium cluster XIVa in MS patients, but they did not identify those bacteria at the species level [41]. Several members of Clostridium cluster XIVa are butyrate producers, and butyrate has been shown to induce T regulatory cells [32, 82, 83, 102]. Furthermore, several studies have reported decreased butyrate producers in MS patients [11, 41, 43, 45]. Hence, it is conceivable that loss of butyrate producers is implicated in the pathophysiology of MS, and more studies are needed to identify species belonging to Clostridium cluster XIVa that play a role in MS pathophysiology.

One of the most striking differences between the microbiota of vancomycin-treated mice and untreated EAE mice was depletion of Anaerotruncus colihominis (OTU 2694) in the single-treatment housed vancomycin group. We found a positive correlation between A. colihominis and EAE severity. Interestingly, we and others have previously reported increased abundance of A. colihominis in MS patients [19, 40]. To this date, no human-derived bacteria has been shown to exacerbate EAE. Given that A. colihominis is depleted from the gut of mice protected from EAE, positively associated with EAE severity, and is increased in MS patients, we hypothesized that it would exacerbate EAE. However, A. colihominis is a butyrate producer [90], and butyrate has been shown to ameliorate EAE via induction of Tregs [85, 86]. Thus, whether Anaerotruncus contributes to disease or could protect against CNS autoimmunity is needed to be tested experimentally. We found that A. colihominis dampened EAE severity, and this was associated with increased frequency of RORγt+T regulatory cells in the mesenteric lymph nodes. RORγt+Tregs, secrete IL-10, is a cytokine critical for immunoregulation in EAE, suggesting a possible mechanism by which these cells attenuate disease [103–105] (see also supplementary material: Additional file 7). Taken together, our findings suggest that similar to Akkermansia muciniphila, increased Anaerotruncus colihominis in MS patients represents a protective mechanism associated with recovery from the disease and is not part of the pathogenic mechanism that induces or maintains the disease. Studies have demonstrated that intestinal RORγt+Treg cells are microbiota dependent, enriched in the gut, and have a strongly suppressive capacity during intestinal inflammation [93, 94]. Hence, these findings suggest that suppressing intestinal inflammation may result in decreased CNS inflammation. To this date, intestinal RORγt+Treg cells have not been implicated in CNS autoimmune diseases; hence, future studies should investigate the role of these Tregs in EAE mice and MS.

Conclusions
In summary, we identified 50 gut commensals that correlate with EAE severity. Interestingly, most of the bacteria that showed a correlation with EAE severity exist in the human gut. As stated above, some of these bacteria have been implicated in MS. We also identified several bacteria that correlate with EAE severity that have not yet been implicated in MS. Some of our correlation findings have already been validated in the EAE mice model. In addition, we experimentally validated our finding of a correlation between A. colihominis and EAE severity. This approach can serve as a framework to test additional candidates that we identified in our study. Furthermore, future studies could perform shotgun metagenomics and metabolomics to identify species/strain-specific factors in the bacteria we have identified as well as metabolites that modulate neuroinflammation in the context of EAE/MS. The findings from these studies will ultimately lead to the identification of bacteria communities that are either pathogenic or neuroprotective in MS. This in turn will facilitate the development of probiotics and other gut bacteria-derived products for the prevention and treatment of MS.

Abbreviations
ALS: Amyotrophic lateral sclerosis; B6: C57BL/6J mice strain; BHI: Brain-heart infusion; CPA: Complete Freund’s adjuvant; CNS: Central nervous system; DPI: Days post immunization; EAE: Experimental autoimmune encephalomyelitis; EDSS: Expanded Disability Status Scale; FDR: False discovery rate; GF: Germ-free; GMSCF: Granulocyte-macrophage colony-stimulating factor; HC: Healthy control; IFNy: Interferon gamma; IL-17: Interleukin-17; IL-10: Interleukin-10; LEfSe: Linear discriminant analysis effect size; mLN: Mesenteric lymph node; MOG: Myelin oligodendrocyte glycoprotein; MS: Multiple sclerosis; OTU: Operational taxonomic unit; PBS: Phosphate-buffered saline; PBMCs: Peripheral blood mononuclear cells; RORγt: RAR-related orphan receptor gamma T; SFB: Segmented filamentous bacteria; Th: T-helper cell; Treg: T regulatory cell.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s40168-022-01364-2.

Additional file 1: Figure S1. Changes in alpha diversity and microbiota composition during EAE. A. α-Diversity metrics for Shannon diversity, observed species and Faith’s Diversity were calculated at an average sampling depth of 1500 reads per sample. No significant differences were observed for any of the diversity estimators analyzed (Mixed-effect model followed by Dunnett’s test) at the indicated time points. B. Taxa plots showing compositional differences in fecal microbiota in mice at the indicated time points. EC1: Enterococcus canisterstini/canis/dispar/durans/faecalis/faecium/hiraer/actic/munditii/ratti/nivorum/villorum.

Additional file 2: Figure S2. Compositional differences in the microbiota during EAE. Bar plots showing changes in the relative abundance of all significantly altered species identified by Linear Discriminant Analysis effect size. Results are presented as mean ± SEM (n = 11-12 mice). Asterisks represent significant difference at the time point indicated compare to day 0 (prior to immunization). *p < 0.05, **p < 0.01, ***p < 0.001. Numbers in parenthesis represent operational taxonomic unit.

Additional file 3: Figure S3. Effect of Vancomycin on EAE development in germ free mice colonized with SBF-free microbiota. DNA agarose
Additional file 4: Figure S4. Compositional differences in the microbiota of untreated and vancomycin treated mice using EAE. Linear Discriminant Analysis (LDA) effect size of significantly altered bacteria at the species level at the indicated time points in: A. single-treatment housed untreated mice (Control) and single-treatment housed vancomycin mice (Vancomycin). Positive LDA effect size is increased in Vancomycin. B. single-treatment housed vancomycin mice (Vancomycin) and vancomycin treated mice cohoused with untreated mice (VancoCoho). Positive LDA score is increased in the underlined group. C. untreated mice cohoused with vancomycin treated mice (ControlCoho) and vancomycin treated mice cohoused with untreated mice (VancoCoho). Positive LDA score is increased in the underlined group. OTU = Operational taxonomic unit; Numbers in parenthesis represent OTU. EC1: Enterococcus canintensis/cani/dispar/durans/faecalis/faecium/hirae/lactis/mundtii/ratti/rivorum/silver街区/5 mice per group are shown. Error bars denote mean ± SEM; Mann Whitney test was performed. *p < 0.05

Additional file 5: Figure S5. Effect of Anaerotruncus colihominis and Enterococcus faecalis on intestinal and peripheral T cell populations. (A) Gating strategies used to identify and quantify Th17 cells (RORγt+ of CD3+FoxP3+ cells), regulatory T cells (RORyt+ of CD3+FoxP3+ cells) and RORyt+ regulatory T cells (RORyt+ of CD3+FoxP3+ cells). FSC – forward scatter, SSC – side scatter. (B-E) The proportion of IL-17A+ IFNγ+, IL-10+ and GM-CSF+ CD4 T cells (cytokine+ of CD4+CD45+ cells) in colon and ileum lamina propria, mesenteric lymph nodes (mLN) and spleen of naive conventionally raised mice orally gavaged with Anaerotruncus colihominis (Anaero), Enterococcus faecalis (Entero) or vehicle (PBS). (E) The proportion of RORyt+ regulatory T cells (RORyt+ of CD3+FoxP3+ cells) in the spleen of naive conventionally raised mice orally gavaged with Anaero, Entero or PBS. Boxplots show the median and interquartile range (IQR) with error bars showing the range. Each point shows data from one mouse. Statistical comparisons are made using ANOVA with Tukey correction. *p < 0.01, ns – not significant (p > 0.05).

Additional file 6: Figure S6. Effect of Anaerotruncus colihominis and Enterococcus faecalis on dendritic cell populations. (A) Gating strategies used to identify and characterize tolerogenic CD103+ dendritic cells (CD103+ of CD11c+MHCII+ cells) and CD80+ dendritic cells (CD80+ of CD11c+MHCII+ cells). FSC – forward scatter, SSC – side scatter. (B and C) The proportion of tolerogenic CD103+ dendritic cells (CD103+ of CD11c+MHCII+ cells) in the mesenteric lymph nodes (mLN) of naïve conventionally raised mice orally gavaged with Anaerotruncus colihominis (Anaero), Enterococcus faecalis (Entero) or vehicle (PBS). (D and E) The proportion of CD80+ dendritic cells (CD80+ of CD11c+MHCII+ cells) in the mLN and spleen of naïve conventionally raised mice orally gavaged with Anaero, Entero or PBS. Boxplots show the median and interquartile range (IQR) with error bars showing the range. Each point shows data from one mouse. Statistical comparisons are made using ANOVA with Tukey correction. ns – not significant (p > 0.05).

Additional file 7.

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Authors’ contributions
SKT, PB, GJB, EMS, and KI performed experiments; LMC, PB, DSW, and JCC analyzed the microbiome data; SL assisted with animal experimentation; GJB and SKT analyzed the FACS data; SKT, PB, GJB, and DSW generated the figures; and SKT and PB wrote the manuscript with input from all authors. SKT conceived the study, designed the analysis, and finalized the manuscript. The authors read and approved the final manuscript.

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Availability of data and materials
The microbiota 16S rRNA sequence data will be submitted to the National Center for Biotechnology Information Short Read Archives and made available to the public once the paper is published. Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Stephanie Tankou (stephanie.tankou@msm.edu).

Declarations
Ethics approval and consent to participate
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Consent for publication
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Competing interests
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