Using Precisely Defined in vivo Microbiotas to Understand Microbial Regulation of IgE

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INTRODUCTION

The prevalence of allergies has been increasing over the last 50 years. The hygiene hypothesis postulated a link between decreased microbial exposure and increased type 2 immune responses (1). Later observations revealed that increased hygiene led to changes in both allergic and autoimmune diseases, leading to the counter-regulatory model (2, 3). Growing understanding of the impact of the intestinal microbiome on immune regulation led to formulation of the microflora hypothesis, which suggested that changes in the composition and richness of gut microbial communities underlie allergic diseases (4). Indeed, many studies now support the suggestion that microbial colonization during a critical window in early life is particularly important for development of a regulated immune system in both mice and man (5).

Elevated total serum IgE levels are a hallmark of allergies but are also observed in multiple immunodeficiencies, including some characterized by deficiencies in regulatory T cells (6, 7). We have previously shown that hyper-IgE is associated with an underlying immune dysregulation or deficiency (8). In line with this, germ-free mice display abnormally high serum IgE levels (8–11).
We have previously shown that exposure to an increased diversity of microbes in early life, during a “window of opportunity,” could completely inhibit these high serum IgE levels (11). However, it was unclear whether the inhibition of IgE during this critical window was an additive result of increased diversity or rather reflected the presence of a “keystone” microbial species. The underlying cellular and molecular mechanisms involved in inhibition of IgE were also unclear. Regulatory T cells play a major role in suppressing immune responses to self-antigens, commensal microorganisms, and harmless environmental antigens (12). Defects in regulatory T cell induction or function have been associated with the development of autoimmunity, food allergy and age-dependent unbalanced Th2 responses at mucosal sites (7, 13–15). There are thought to be two populations of Tregs; one derived from the thymus, referred to as tTregs, and another originating from naïve T cells in the periphery (pTregs) (16).

Intestinal bacteria, such as some Clostridia species, have been shown to induce pTreg differentiation via their production of the short chain fatty acid (SCFA) butyrate in the colon (17, 18). Furthermore, a subpopulation of Tregs, defined as CD4+ Foxp3+ RORγt+ Helios−, has been shown to be induced by the microbiota in the small intestine, colon and GALT tissues (19–21). It is unclear, however, whether there is a functional link between microbial colonization, pTreg induction, in particular the RORγt+ Helios− pTregs subpopulation, and IgE regulation. We hypothesized that certain bacterial species and their production of SCFA could induce the immune regulation of IgE by increasing peripherally induced pTregs. Here, we generated an extensive collection of gnotobiotic mouse colonies and found that three bacterial species that colonize the small intestine in early life can together suppress hygiene-mediated hyper-IgE. We identified acetate production, immunogenicity and mucosa association as key characteristics of bacterial consortia that when combined provide the capacity to inhibit the induction of hyper-IgE.

MATERIALS AND METHODS

Mice, Hygiene Status and, Bacterial Colonization

C57BL/6j mice were re-derived to germ-free status via two-cell embryo transfer. Axenic and gnotobiotic mice (germ-free and precisely colonized) were bred and maintained in flexible-film isolators at the Clean Mouse Facility (CMF), University of Bern, Switzerland or at the International Microbiome Centre (IMC), University of Calgary, Canada. The germ-free and gnotobiotic C57BL/6j strain used in Bern and Calgary were identical. For some experiments, mice were housed in individually ventilated isocages (Tecniplast) in the IMC. Germ-free status was routinely monitored by culture-dependent and— independent methods and all germ-free colonies were independently confirmed to be pathogen-free. To start gnotobiotic mouse colonies, germ-free breeding pairs were gavaged with single or mixed bacterial cultures, as indicated. Gram staining, SYTOX™ (Fisher) DNA staining and 16S rRNA gene amplicon sequencing of their intestinal contents were performed to confirm their colonization status. The offspring of these gnotobiotic breeding pairs were analyzed at 10–13 weeks of age, unless stated otherwise. SPF mice were either purchased from Envigo (Switzerland) or bred and maintained at the University of Calgary. The SPF mice at the University of Calgary were fed the same autoclaved diet as their germ-free and gnotobiotic counterparts. Although two different SPF colonies were used, one in Switzerland (Envigo, Netherlands) and another in Canada (in house SPF colony) we did not observe any significant differences in our readouts between these two SPF colonies. The only data derived from Envigo SPF mice are n = 18 of a total of n = 80 mice shown in Figure 1B as well as the data shown in Supplementary Figure 3. All animal experiments were in accordance with the guidelines established by either the Swiss Federal Veterinary Office or by the Canadian Council for Animal Care and were approved by the Commission for Animal Experimentation of the Veterinary Office of Canton Bern or by the University of Calgary Animal Care Committee.

Bacterial Culture

Bacteria were cultured in Brain Heart Infusion (BHI) medium as previously described (22). Bottles and media were gassed with 10% H2, 10% CO2, and 80% N2 while in a Whitley A95 anaerobic incubator before being sealed and inoculated with specific bacterial species.

Bacterial Flow Cytometry

Bacterial flow cytometry was performed as described previously (23, 24). Briefly, bacteria were cultured for 24–48 h as described above, with the modification that the BHI media was filter-sterilized before inoculation. For growing Akkermansia muciniphila YLA4, 0.25 g/L of autoclaved Type II hog gastric mucin (Sigma) was added to the filter-sterilized BHI media. Bacterial cultures were centrifuged for 10 min at 3,000 g, washed twice and re-suspended in sterile-filtered PBS/2%BSA/0.02% sodium azide and diluted to ~10^7 bacteria/ml (OD_950 0.1 = 10^8 bacteria/ml). Serum was diluted 1:10 in PBS/2%BSA/0.02% sodium azide, heat-inactivated at 56°C for 30 min and centrifuged at 16,000 g for 5 min at 4°C to remove any bacteria sized contaminants. This serum supernatant was used to perform serial dilutions. 2.5 × 10^5 bacteria were added to each well and incubated with the serial dilutions of serum for 1 h at 4°C, centrifuged for 10 min at 3,000 g, washed twice and then re-suspended in monoclonal APC rat anti-mouse IgG1 (1:40, A85-1) (BD) and BV605 rat anti-mouse IgA (1:40, C10-1) (BD) and incubated overnight at 4°C. The bacteria were washed twice in PBS/2%BSA/0.02% sodium azide and then re-suspended in PBS/2%BSA/0.02% sodium azide and acquired on a FACS Canto (BD) using FSC (forward scatter) and SSC (side scatter) in logarithmic mode. Data was analyzed using FlowJo software (Tree Star Inc.).

Small Intestinal Wash Collection

The entire small intestine was removed and washed with 5 ml of ice-cold intestinal wash buffer [10% 0.5 M EDTA pH 8.0, 10% 10x PBS, ddH2O and 0.1 mg ml⁻¹ soybean trypsin inhibitor...
(Sigma)] with 40 µl of 100 µM phenylmethylsulfonylfluoride (PMSF, Sigma) per 5 ml of intestinal wash buffer. The small intestinal wash was centrifuged at 4,000 g for 10 min at 4°C and stored at −80°C.

**DNA Extraction From Intestinal Contents**

Small intestines, ceca and colons were opened longitudinally. Contents were removed and snap-frozen in liquid nitrogen and stored at −80°C. DNA was extracted from the contents using the QIAamp Fast DNA Stool Mini Kit (Qiagen), according to the manufacturer’s instructions. Briefly, samples were homogenized in InhibitEX buffer by bead-beating using differentially sized beads (glass beads, 0.5–0.75 mm; zirconia beads, <100 µM), treated with lysis buffer (1.2% Triton X-100, 2 mM EDTA, 20 mM Tris·HCl) containing lysozyme (20 mg ml⁻¹, Sigma), followed by Proteinase K treatment. Bacterial DNA was ethanol-precipitated on a column membrane and eluted with sterile water. Procedural blank DNA extractions were performed in parallel with each batch of DNA extractions and run as controls during the downstream PCR steps to confirm the absence of contaminating DNA sequences from the extraction process.

**16S rRNA Gene Amplicon Sequencing**

Two platforms were used for 16S rRNA gene amplicon sequencing. For the gnotobiotic models C1 and C2 (Supplementary Table 1) the 16S rRNA gene segments spanning the variable V5 and V6 regions were amplified from DNA from intestinal content samples using a multiplex approach with the barcoded forward fusion primer 5′-CCATCTCA TCCCTGCGTGTCTCCGACTCA G BARCODE ATTAGAT ACCCYGGTAGTCC-3′ in combination with the reverse fusion primer 5′-CTCT CTATGGGCAGTCGGTGATACGAGCTGACGACARCCATG-3′. The sequences in *italics* are Ion torrent PGM-adaptor sequences. The PCR amplified 16S rRNA V5-V6 amplicons were purified and prepared for sequencing on the Ion torrent PGM system according to the manufacturer's instructions (Life Technologies). Samples with over 1,000 reads were used for the analysis. Data analysis was performed using the QIIME pipeline version 1.8.0 (25). Operational taxonomic units were picked using UCLUST (26) with a 97% sequence identity threshold followed by taxonomy assignment using a custom C1 or C2 database for gnotobiotic C1 and C2 intestinal samples. For all other gnotobiotic models, the 16S rRNA gene segment spanning the variable V4 region was amplified from DNA extracted from intestinal content samples using a multiplex approach with the barcoded forward fusion primer 5′-AATGATACGGCGACCACCGAGATCTACAC i5BARCODE TATGGTAATTGTGTGCCAGCMGCGCGGTAA-3′ in combination with the reverse fusion primer 5′-CTCT CTATGGGCAGTCGGTGATACGAGCTGACGACARCCATG-3′. One PCR reaction contained 0.25 mM of each primer, KAPA HiFi Hot Start Ready Mix (Roche) and 20–100 µg of template DNA.
PCR conditions were 98°C for 2 min, followed by 25 cycles of 98°C for 30 s, 55°C for 30 s and 72°C for 20 s and a final extension step at 72°C for 7 min. Amplification of the V4 gene region was verified by electrophoresis on a 1.2% agarose gel. Reactions were purified using NucleoMag NGS clean-up beads (Macherey-Nagel) and were normalized using the SequaPrep Normalization Plate Kit (Illumitrogen). Amplicons were pooled and concentration and quality were determined using the Qubit HS DNA kit (Qubit) and the Tapestation D1000 assay (Agilent), respectively. Amplicon sequencing was done on a MiSeq (Macherey-Nagel) and were normalized using the SequalPrep Reactions were purified using NucleoMag NGS clean-up beads (Promega). The intestines were opened longitudinally, rinsed in Dulbecco's saline (DPBS) lacking CaCl₂ and MgCl₂ (Invitrogen), cut into 1-2 cm long pieces and washed once for 20–30 min (small intestine) or 25–35 min (colon) at 37°C with shaking at 220 rpm. The resulting cell suspension was passed through a cell strainer (100 µM). Cells were washed several times with HBS/5%HS and centrifuged at 400 g for 5 min and re-suspended for quantification.

For Figure 3, after digestion and centrifugation the small intestine cell pellet was re-suspended in 40% Percoll solution and layered on top of a 70% Percoll solution (GE Healthcare). Gradient centrifugation was carried out at 700 g for 20 min at room temperature with no break. Cells lying at the 40%/70% interphase were collected and washed with HBS/5%HS and centrifuged at 400 g for 5 min and re-suspended for quantification. Mesenteric lymph nodes (MLNs) and PP were digested with HBS/5%HS containing 1 mg ml⁻¹ collagenase IA (Sigma) and 10 U DNAse I (Roche) for 20 min at 37°C and passed through a cell strainer (100 µM), washed with HBS/5%HS and centrifuged at 400 g for 5 min and re-suspended for quantification. Splenocytes were passed through a cell strainer (100 µM) and centrifuged at 400 g for 5 min. Spleen cell pellets were re-suspended in 1 ml of 0.88% NH₄Cl for 10 min at room temperature to lyse red blood cells. Cells were washed with HBS/5%HS and centrifuged at 400 g for 5 min and re-suspended for quantification.

For intracellular cytokine staining small intestinal single cell suspensions were stimulated for 4 h at 37°C/5% CO₂ in Iscove's Modified Dulbecco's Media (IMDM, Fisher) containing 10% Fetal Bovine Serum (FBS, Fisher) in the presence or absence of 50 ng ml⁻¹ phorbol 12-myristate 13-acetate (PMA, Sigma) and 750 ng ml⁻¹ ionomycin (Invitrogen) with 10 µg ml⁻¹ Breifeldin A (Sigma).

**Flow Cytometry**

Cells were washed twice with DPBS and stained with either fixable viability dye eFluor506 (eBioscience) or fixable viability stain 780 (BD) and anti-CD16/CD32 (2.4G2; BD) in DPBS for 10 min at 4°C. Cells were then washed with FACS buffer containing 2% fetal calf serum (Invitrogen), 2.5 mM EDTA (MP Biomedicals) in PBS. For cell surface staining, cell pellets were re-suspended in antibodies diluted in FACS buffer and stained for 30 min at 4°C. Intra-nuclear staining was performed using the Foxp3 staining kit (eBioscience). Intracellular cytokine staining was performed with the Cytofix/Cytoperm staining kit (BD). The following mouse-specific conjugated antibodies were used: CD45-PerCP-Cy5.5 (30-F11; BD), CD45-BV510 (30-F11; BD), TCRβ-BV786 (H57-597; BD), CD4-Pe-Cy7 (RM4-5; BD), Helios-AlexaFluor488 (22F6; BD), CD3-PB (145-2C11; Biolegend), CD4-BV785 (RM4-5; Biolegend), Helios-FITC (22F6; Biolegend), IL10-Pe-Cy7 (JES5-16E3; Biolegend) Foxp3-AlexaFluor700 (FJK-16s; eBioscience), and RORγt-PE (Q31-378; eBioscience). Cells were acquired on a FACS Fortessa (BD Biosciences) or a FACS Canto (BD) with FACS DIVA software (BD). Data analysis was performed using FlowJo software (Tree Star Inc.).

**IgE ELISA and IgE Electrochemiluminescence Immunoassay (IgE-ECL)**

Blood was collected in serum-separating tubes (Sarstedt and BD) and total serum IgE concentrations were measured.
using either the BD OptEIA Mouse IgE ELISA Set (BD) according to the manufacturer’s instructions, or a multi-array (Meso Scale Discovery) utilizing the BD OptEIA Mouse IgE ELISA Set.

**IgA and IgG1 Electrochemiluminescence Immunoassay**

Total concentrations of IgA and IgG1 in serum and small intestinal wash were determined using the Mouse Isotyping
A further increase in microbial diversity suppresses hyper-IgE and demonstrates a dynamic bacterial colonization profile in the small intestine and colon in early life. (A) Relative species abundance in the small intestine (SI) and cecum of a representative adult Oligo-MM$^{12}$ mouse, $n = 4$ mice. (B) Total serum IgE levels in Oligo-MM$^{12}$ ($n = 100$) mice. The blue and orange horizontal lines represent the geometric mean of the serum IgE levels from the GF and SPF cohorts, respectively. (Continued)
Panel 1 kit according to the manufacturer's instructions (Meso Scale Discovery).

**LC/MS/MS Based Short Chain Fatty Acid Metabolite Analysis**

Cecal contents were collected, snap-frozen in liquid nitrogen and stored at −80°C until processed. Native SCFAs were extracted from cecal samples with an ice-cold 50% acetonitrile solution (2:1 (v/w) ratio) containing known amounts of the following 13C-SCFA analytical standards used as internal standards (IS): acetic acid (1,2-13C2, 99%, #CLM-113, Cambridge Isotope Laboratories, Inc, Andover, MA, USA), Propionic acid (IS): acetic acid (1,2-13C2, 99%, #CLM-113, Cambridge Isotope Laboratories, Inc, Andover, MA, USA), Butyric acid (1,2-13C2, 99%, #491993, Sigma-Aldrich) and Propionic acid (13C3, 99%, #589586, Sigma-Aldrich, St. Louis, MO, USA) and Butyric acid (1,2-13C2, 99%, #491993, Sigma-Aldrich) (Supplementary Table 2). Samples were homogenized at 30 Hz for 3 min with a tissue lyzer (Qiagen) and centrifuged at 18,000 g for 10 min at 4°C. Supernatants were collected and centrifuged under the same conditions. Supernatants were then derivatized with EDC (N-(3-Dimethylaminopropyl)-N’-ethylcarbodiimide hydrochloride) and aniline (32, 33) as follows: 100 µL aliquot were incubated with EDC (50 mM, final concentration) and aniline (100 mM, final concentration) for 2 h at 4°C. An aliquot of each sample was further diluted with 50% HPLC-grade methanol/water. All samples were stored at 4°C until analysis on the same day.

LC-MS/MS analysis of cecal contents was performed on a Vanquish™ UHPLC System coupled to a TSQ Quantum™ Access MAX triple quadrupole Mass Spectrometer (Thermo Scientific) equipped with an electrospray ionization (HESI-II) probe. The UHPLC-MS platform was controlled by Xcalibur™ data system (Thermo Scientific).

Chromatographic separation was achieved on a Hypersil GOLD™ C18 column (200 X 2.1 mm, 1.9 µm, Thermo Scientific) using a binary solvent system composed of LC-MS grade H2O with 0.1% (%v/v) formic acid (Solvent A) and LC-MS grade methanol with 0.1% (%v/v) formic acid (solvent B). The following 21 min gradient was used: 0–1 min, 40%B; 1–7 min, 40–98%B; 7–15 min, 98%B; 15–16 min, 98–40%B; 16–21 min, 40%B. LC eluent was diverted to waste for the first 5 min of the run. The flow rate was 200 µL/min and the sample injection volume 2 µL. The auto sampler was kept at 4°C and the column at 30°C.

MS/MS data were acquired in positive ionization mode with the mass spectrometer operating in Selected Reaction Monitoring (SRM) mode. Fragmentation parameters were optimized using the EZ Tune program with direct infusion of the derivatized analytical grade standards. Subsequently, the following transitions, corresponding to the three derivatized native SCFAs and respective derivatized 13C-SCFA standards, were monitored, with a scan time of 0.05 s and a fixed collision energy of 14 eV: [M+H]+ m/z 136.07, 138.08, 150.09, 153.10, 164.10, 166.11 → m/z 94.06. Electrospray ionization source conditions were as follows: spray voltage of 3,000 V, vaporizer temperature of 300°C, sheath gas of 5 (arbitrary units), sweep gas of 1 (arbitrary units), auxiliary gas of 2 (arbitrary units), capillary temperature of 275°C.

Data analyses, on the converted mzXML files, were conducted in MAVEN (34, 35). In short, for each SCFA, the determination of the native SCFA concentration was based on the 13C/13C signal intensity ratio and the respective 13C-IS concentration.

**Statistical Analysis**

Statistical analysis was performed by one-way ANOVA with Tukey’s or Dunnett’s post-test depending on whether the comparison was performed between all means (Tukey’s) or compared to a control mean (Dunnett’s) using GraphPad Prism 8 (GraphPad Software Inc.). The data for serum IgE concentrations was first log transformed before performing one-way ANOVA with Tukey’s post-test. The SCFA in colon vs. cecum (in Supplementary Figure 1) were compared using Student’s t-test. A p < 0.05 was considered statistically significant. P-values are indicated as follows: *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, not significant (ns).

**RESULTS**

**The Presence of a Butyrate-Producing Bacterial Species Has a Minimal Effect on Adult Serum IgE Levels**

We have previously shown that mice colonized with the Altered Schaedler Flora (ASF), harboring eight commensal intestinal bacterial species (36), display a dichotomy in terms of development of high IgE levels (11). High IgE levels in adulthood correlated with a low ASF diversity early in life while a high ASF diversity early in life suppressed induction of IgE. We also noted that the presence of the butyrate-producing bacteria *Pseudoflavonifractor* sp. ASF500 early in life correlated with suppression of IgE (11). We and others have also previously shown that ASF colonization (37) or colonization with microbial consortia containing *Clostridia* species (38, 39) results in the induction of intestinal pTregs and the SCFA butyrate has been demonstrated to support pTreg induction (17, 18). To test whether butyrate produced by *Pseudoflavonifractor* sp. ASF500 could suppress hyper-IgE, we gavaged germ-free C57BL/6 breeding pairs with either microbial...
Community #1 (C1, Supplementary Table 1) consisting of three non-butyrate-producing ASF species we know do not suppress hyper-IgE, or community C2, which had the butyrate-producer *Pseudoflavonifractor* sp. ASF500 added to the community (Supplementary Table 1). This approach was necessary because *Pseudoflavonifractor* sp. ASF500 is unable to colonize a germ-free mouse on its own and requires the presence of other bacterial species (data not shown). Colonization with either C1 or C2 in the small intestine and cecum of the offspring was confirmed by 16S rRNA amplicon sequencing (Figure 1A). *Pseudoflavonifractor* sp. ASF500 was only detected in the C2 group and *Lactobacillus intestinalis* ASF360 (present in the inoculum for C1 and C2) was below the detection limit in both groups.

Total serum IgE levels were measured in large control cohorts of GF and specific pathogen free (SPF) mice (Figure 1B). As previously shown (11), IgE levels in adult GF mice were elevated and significantly increased above SPF levels. We found that IgE levels in adult C1- or C2-colonized offspring remained significantly elevated at the level of GF controls (Figure 1B). We found that colonization with both C1 and C2 led to significant increases in acetate and propionate levels compared to GF levels, whereas only C2 significantly increased butyrate levels in the cecal contents, as expected (Figure 1C). All SCFA levels in the C1 and C2 cohorts remained significantly lower than that found in SPF mice (Figure 1C). These data indicated that low levels of SCFA were insufficient to inhibit hygiene-induced hyper-IgE. Of note, SCFA levels in cecal contents were comparable to the levels observed in colon contents (Supplementary Figure 1).

**A More Diverse Microbial Community Suppresses Hyper-IgE**

We have previously demonstrated that bacterial diversity during early life is important to suppress hyper-IgE (11). Since our sequential increase of ASF diversity (C1 < C2 < C3 < C4) had no inhibitory effect on serum IgE levels, we hypothesized that none of these microbiotas (C1 to C4) reach a high enough diversity early in life to suppress hyper-IgE. Since the full ASF showed only partial suppression of hyper-IgE depending on colonization diversity early in life (11) we hypothesized that further experimentally increasing the microbial diversity will eventually result in suppression of hyper-IgE. We recently established a gnotobiotic microbiota consisting of 12 defined and well-characterized bacterial species. This microbiota is referred to as either stable Defined Moderately Diverse Microbiota from mice (sDMDMm2) (45, 46) or Oligo-mouse microbiota (Oligo-MM12) (22, 47) (Supplementary Table 3). Analysis of the microbiota composition in the small intestine and cecum of our gnotobiotic Oligo-MM12 mouse cohort revealed that 11 of the 12 species were detectable by 16S rRNA amplicon sequencing (Figure 3A) with *Bifidobacterium longum* subsp. *animalis* YL2 undetectable [as has previously been observed (22)]. More importantly, however, colonization with the Oligo-MM12 resulted in suppression of hyper-IgE (Figure 3B), which correlated with increased production of acetate and propionate compared to C4-colonized mice while butyrate levels remained unchanged (Figure 3C). However, the suppression of hyper-IgE did not reach the levels observed in SPF mice (Figure 2B), which indicates that increasing bacterial diversity even further to SPF enhances suppression of hygiene-induced hyper-IgE (11). Notably, colonization with Oligo-MM12 also resulted in significant induction of RORγt+Helios− pTregs (Figures 3D,E and Supplementary Figure 3).

We next followed the Oligo-MM12 colonization dynamics in early life in the small intestine and colon starting from day 8 until day 43 after birth (Figure 3F). Up until about 2 weeks of age there was a dominance of *Enterococcus faecalis* KB1 and *Akkermansia muciniphila* YL44 in both the small intestine and colon. The level of *E. faecalis* KB1 then declined drastically concomitantly with an increase in *Bacteroides caccae* IRS148 (Figure 3F). These changes coincided with weaning of the pups, suggesting that the change in diet from milk to solid food contributes to the increase of certain bacterial species. In light of our previous findings (11), these data suggest that the two species (*E. faecalis* KB1 and A. propria (cLP) (Figures 2D,E). C4 colonization also led to an increase of RORγt+Helios− pTregs in the spleen (Figures 2D,E). The transcription factor Helios has been suggested to identify tTregs whereas pTregs are characterized as Helios negative (40, 41). However, it remains controversial whether Helios, or other markers such as Neuropilin-1 (42, 43), can unequivocally identify tTregs vs. pTregs (44). We have therefore included expression of RORγt to better identify microbially-induced pTReg (19–21).

These results indicate that although the microbial communities of C3 and C4 produced SCFA and induced RORγt+Helios− pTregs, this was not sufficient to inhibit hyper-IgE.
muciniphila YL44) most prominent in early life may contribute to regulation of IgE levels. In addition, since Clostridia species are known to be dominant producers of SCFA, and we found that C. clostridiiforme YL32 and Clostridiales species B. coccoides YL58, were also present in early life, albeit at lower levels, we further dissected the protective potential of these four bacterial species.

**Two Dominant Early Life Clostridiales Are Unable to Regulate Serum IgE Levels**

Since they are often obligate anaerobes, many Clostridiales species are unable to monocolonize a germ-free gut. We therefore added B. coccoides YL58 to C2 to generate community C5 and added C. clostridiiforme YL32 to C5 to generate C6 (Supplementary Table 1). Successful addition of these species was confirmed by 16S rRNA amplicon sequencing (Figure 4A). We found that total IgE levels were not altered from germ-free levels in either C5- or C6-colonized mice (Figure 4B). Addition of the Clostridiales species also did not significantly alter SCFA levels compared to C4-colonized mice (Figure 4C), despite induction of RORγt+Helios− pTregs in this cohort (Figures 4D,E and Supplementary Figure 4). These results indicated that other Oligo-MM12 members are likely contributing to suppression of hyper-IgE.

**Acetate Produced by B. coccoides YL58 Contributes to Suppression of Serum IgE Levels**

Since regulation of IgE is dependent on early-life microbial colonization (11), we next investigated whether E. faecalis KB1 and A. muciniphila YL44, the two most dominant members of the Oligo-MM12 consortia in early life (Figure 3F), were able to individually regulate serum IgE levels. We monocolonized breeding pairs with either E. faecalis KB1 or A. muciniphila YL44 and determined the serum IgE levels and cecal content SCFA concentrations in their offspring (Figure 5). Successful colonization with these individual bacteria was confirmed by 16S rRNA full length sequencing of DNA extracted from the feces and by gram staining of intestinal contents (data not shown). We found that total serum IgE levels in E. faecalis KB1 and A. muciniphila YL44 monocolonized mice were not significantly altered from germ-free levels (Figure 5A).

As neither E. faecalis KB1 nor A. muciniphila YL44 by themselves were able to suppress IgE we next tested these two in combination. Unfortunately, 16S rRNA amplicon sequencing of our double-colonized colony revealed that it got contaminated with the additional Oligo-MM12 species B. coccoides YL58 (YL44 + KB1 + YL58) (Figure 5B). We did, however, observe a reduction in serum IgE levels in this YL44 + KB1 + YL58 colony (Figure 5A). To investigate whether this effect relied on the presence of B. coccoides YL58 we generated B. coccoides YL58 monocolonized as well as E. faecalis KB1/B. coccoides YL58 (KB1 + YL58) and A. muciniphila YL44/B. coccoides YL58 (YL44 + YL58) double colonized cohorts. We found that while B. coccoides YL58 alone was not able to reduce serum IgE levels, we observed reduced IgE levels when B. coccoides YL58 was present in combination with E. faecalis KB1 or A. muciniphila YL44 (Figure 5A). This indicated that functional cooperation between bacterial species with different characteristics was required to induce an IgE-suppressing effect. Of note, pTreg proportions in these different cohorts did not correlate with reduced serum IgE levels indicating that inhibition of IgE was not mediated solely by induction of pTregs (Supplementary Figures 5, 6). In addition, the absence of butyrate in all cohorts (Figure 5C), and the absence of propionate in the KB1 + YL58 model suggests that propionate and butyrate are not required for suppressing hygiene-induced hyper-IgE. In contrast, acetate production correlated with reduced serum IgE levels with the exception of the B. coccoides YL58 monocolonized group (Figures 5B,C). Therefore, since B. coccoides YL58 alone was not sufficient to suppress IgE, we concluded that acetate provided by B. coccoides YL58 (Figure 5C) is required but not sufficient for reducing IgE levels. Since E. faecalis KB1 and A. muciniphila YL44 do not produce high levels of acetate (Figure 5C), we next investigated what could be the particular characteristics of E. faecalis KB1 and A. muciniphila YL44 that are required to suppress IgE when present in combination with an acetate-producer.

**Immunogenicity Is an Additional Requirement for IgE Suppression**

A recent study identified A. muciniphila as a strong inducer of systemic IgA and IgG1 (48). We therefore measured serum IgA and IgG1 reactivity against A. muciniphila YL44 in the YL44 monocolonized, YL44 + YL58 bicolonized and YL44 + KB1 + YL58 tricolononized cohorts (Supplementary Figure 7). All A. muciniphila YL44 colonized cohorts displayed IgA and IgG1 reactivity against A. muciniphila YL44. While E. faecalis KB1 monocolonized mice also displayed specific serum IgA reactivity, B. coccoides YL58 monocolonized mice did not (Supplementary Figure 7). In order to extend this immunogenicity screen, we measured total IgA (as opposed to bacteria-specific IgA) in small intestinal washes and serum from various gnotobiotic cohorts (Supplementary Figure 8). We also included segmented filamentous bacteria (SFB) monocolonized mice as a control as SFB is known to be a potent inducer of IgA (49). We found that A. muciniphila YL44, E. faecalis KB1 and SFB were able to promote induction of IgA (Supplementary Figure 8). Importantly, although SFB induced a strong IgA response (Supplementary Figure 8) as well as a RORγt+Helios− pTreg response in the colon, SFB alone was not sufficient to suppress hygiene-induced hyper-IgE, possibly due to the lack of acetate production (Supplementary Figure 9). We therefore conclude that IgA immunogenicity in combination with acetate production is required to drive an IgE suppressive phenotype.

**IL10 Production Does Not Correlate With Protection From Hyper IgE**

Lastly, we investigated whether IL10 production in the small intestine, which is required to maintain intestinal immune homeostasis (50), was required to inhibit hygiene-induced hyper-IgE. We found similar proportions of IL10+CD45+TCRβ− cells in the small intestine of cohorts with high vs. low serum
Two early-life Oligo-MM\textsuperscript{12} Clostridiales bacteria are unable to regulate serum IgE levels. (A) Relative species abundance in the small intestine (SI) and cecum of a representative C5- and C6-colonized mouse, \( n = 2–4 \) mice per group. (B) Total serum IgE levels in C5 (\( n = 30 \)) and C6 (\( n = 49 \)) mice. The blue and orange horizontal lines represent the geometric mean of the serum IgE levels from the GF and SPF cohorts, respectively (from Figure 1B). (C) SCFA levels in the cecal contents.

(Continued)
IgE levels indicating that small intestinal IL10 production alone is not sufficient to suppress hygiene-induced hyper-IgE (Supplementary Figure 10). In addition, SFB monocolonized mice, which had the highest proportion of IL10⁺ cells (Supplementary Figure 10) retained hygiene-induced hyper-IgE (Supplementary Figure 9).

In summary, we generated an extensive range of clearly defined and quality controlled gnotobiotic microbiotas in vivo in combination with read-outs for serum IgE levels, intestinal SCFA levels, Treg induction, IgA and IL10 production to identify characteristics of individual or communities of commensal bacterial species with the capacity to suppress the hyper-IgE syndrome observed in germ-free mice. We found that the IgA-inducing A. muciniphila YL44 and E. faecalis KB1 bacterial species, both dominant in the small intestine in early life, can cooperate with acetate-producing B. coccoides YL58 to inhibit hyper-IgE. Furthermore, this inhibitory effect was independent of propionate and butyrate production, frequencies of IL10⁺CD45⁺TCRβ⁻ cells in the small intestinal lamina propria and frequencies of RORγt⁺Helios⁻ pTregs in the intestinal tissues and GALT.

**DISCUSSION**

Elevated serum IgE levels in germ-free, but also in colonized immuno-deficient animals, has been a long-standing observation (8). We have also previously demonstrated that there is a critical period in early life at around the time of weaning whereby exposure to a diverse group of microbes is essential to prevent isotype switching to IgE (11). However, the mechanism by which this aberrant IgE induction is regulated following intestinal colonization and which commensal species are potent...
mediators of protection remains unknown. Here we show that colonization with A. muciniphila or E. faecalis, that are both predominantly present before weaning and induce IgA, in combination with the acetate-producing bacterial species B. coccoides YL58, results in partial protection from serum hyper-IgE. Surprisingly, this protection did not correlate with microbial induction of RORγt+Helios− pTregs in the mucosal tissues, induction of IL10+CD45+TCRβ− cells, or production of propionate or butyrate.

We identified induction of IgA and mucosa association as key characteristics of the bacteria that were involved in inhibition of IgE. Indeed, both A. muciniphila and E. faecalis are present near the epithelial surface of the small and large intestines (51, 52). Although A. muciniphila is known to have barrier-protective functions (53, 54), it is also a potent inducer of bacteria-specific systemic IgA and IgG1 responses (48). Furthermore, decreases in A. muciniphila have been associated with childhood atopy (55). This indicates that A. muciniphila is an immunogenic commensal species and this characteristic contributes to inhibition of hygiene-induced IgE. In humans, a strong serum IgG response against a common set of intestinal microbial antigens has been associated with protection against allergy development during childhood (56). It is also known that mucosa-association promotes an IgA response (57). Nevertheless, these characteristics are not sufficient to inhibit IgE because monoclonization with A. muciniphila, E. faecalis or SFB failed to reduce IgE levels.

Interestingly, we identified acetate production as a critical characteristic required for IgE inhibition. B cell class switch to IgE occurs in the PP in germ-free mice (11) and we found that acetate (1.9 ± 0.31 μmol/gram), but not butyrate or propionate, and the acetate-producing bacteria B. coccoides YL58 was present in the small intestine. In addition, ffar2/gpr43, an acetate receptor, is also expressed in the small intestine, predominantly on leukocytes in the lamina propria (58). In humans, acetate was found to be significantly reduced in infants at 3 months of age who subsequently developed atopy and wheeze (59) and acetate has also been identified to be the most abundant SCFA in the feces of 3–5-month-old infants (60). Therefore, acetate may be an important bacterial-derived metabolite that regulates early life immunity, including regulation of IgE.

We have previously shown that there is a critical window in early life where microbial colonization is required to inhibit hygiene-induced IgE (11). We now show that the bacteria that are capable of IgE inhibition are dominant in early life prior to weaning.

Al Nabhani et al. has recently characterized a vigorous immune response at weaning, termed the weaning reaction, as it is associated with changes in the microbiota at weaning (61). The weaning reaction consists of a time-dependent increase in tumor necrosis factor alpha (TNF-α) and interferon-γ (IFNγ), and is driven by bacteria, SCFA and vitamin A (61). Thus, it is tempting to speculate that a strong weaning reaction induced by bacteria such as A. muciniphila YL44 and E. faecalis KB1 in the presence of an acetate-producing bacteria (such as B. coccoides YL58) during this critical window is sufficient to suppress IgE. This colonization status could potentially favor IgA and IgG1 isotype class switching events while still inducing the important RORγt+Helios− pTregs population (61). Although induction of RORγt+Helios− pTregs at weaning was critical for protection from inflammation later in life (61), our data indicates that induction of RORγt+Helios− pTregs alone is insufficient to inhibit IgE.

Taken together, we show that both bacterial richness and the presence of key bacterial species in early life are important for regulating serum IgE. A recent publication has demonstrated that the increased IgE levels in germ-free mice are induced by food antigens (62). In context of the weaning reaction (61) and our findings, this suggests that appropriate bacterial signals early in life are required to prevent sensitization to food antigens. Therefore, in addition to the Clostridia species previously identified (63), our data provides additional bacterial candidates and avenues for microbial-based prevention strategies in food allergy.

It is important to note that our bacterial combinations only provided partial suppression of serum IgE levels and other additional microbial characteristics are needed to fully inhibit IgE to the levels found in SPF mice. Therefore, it will be interesting to further investigate the role of other bacterial characteristics and metabolites on IgE regulation in order to develop microbial-based therapeutics for prevention of allergic diseases in humans. Furthermore, it would be interesting to screen for commensal species that are dominant in early life, induce IgA and produce acetate, to see whether these species alone can suppress hyper-IgE. Unfortunately, thus far we did not come across species with these capabilities in our extensive in vivo gnotobiotic screen. Alternatively, once appropriate genetic tools are available, A. muciniphila YL44 and E. faecalis KB1 can be genetically modified to produce acetate and tested in isolation.

DATA AVAILABILITY STATEMENT

The 16S rRNA amplicon sequences generated for this study can be found on the figshare repository at https://doi.org/10.6084/m9.figshare.c.4763306.

ETHICS STATEMENT

The animal study was reviewed and approved by The Commission for Animal Experimentation of the Veterinary Office of Canton Bern and the Health Sciences Animal Care Committee of the University of Calgary.

AUTHOR CONTRIBUTIONS

MW contributed to the experimental design, performed and analyzed all the experiments, and contributed to writing the manuscript. KB contributed to the bioinformatics analysis. FT, KB, CT, and FR contributed to experiments assessing pTreg induction. MK and VF performed and analyzed the bacterial FACs. DB and IL established the targeted metabolomic protocol for the SCFAs. MG and KM analyzed and interpreted the data and wrote the manuscript. KM conceived and designed the project.
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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2019.03107/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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