Bifidobacteria-mediated immune system imprinting early in life

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SUMMARY

Immune-microbe interactions early in life influence an individual's risk of developing allergies, asthma and some autoimmune disorders. Breastfeeding helps guide the development of healthy immune-microbe relationships, in part by providing nutrients to specialized microbes that in turn benefit the host and its developing immune system. Such bacteria having co-evolved with humans are associated with reduced risks of immune mediated diseases but are increasingly rare in modern societies. Here we map an immunological sequence of events, triggered by microbial colonization that distinguish children with different gut bacterial composition. Lack of bifidobacterial species is associated with elevated markers of intestinal inflammation and immune dysregulation and in a randomized trial of breastfed infants, the infant-adapted *Bifidobacterium infantis* EVC001 silenced intestinal Th2 and Th17 immune responses, while inducing IFNβ, and its metabolites skew T-cell polarization *in vitro*, from Th2 towards Th1, suggesting a healthier immune imprinting during the first critical months of life.

HIGHLIGHTS

An ordered sequence of immune changes after birth, driven by microbial interactions

Low gut *Bifidobacterium* abundance is associated with markers of intestinal inflammation

Feeding B. infantis EVC001 silenced intestinal Th2 and Th17 but upregulates IFNB

B. infantis EVC001 metabolites and/or enteric cytokines skew naïve T-cell polarization towards Th1 and away from Th2

INTRODUCTION

Mounting evidence indicate that the composition of the infant gut microbiome is critical to immunological development, particularly during the first 100 days of life in which aberrations in gut microbial composition are most influential in impacting the developing immune system. Indeed, multiple studies have emphasized how early gut microbiome dysbiosis, described as an overabundance of Proteobacteria (Shin et al., 2015) and loss of ecosystem function (Duar et al., 2020) is associated with both acute and chronic immune dysregulation, leading to common conditions such as colic (Rhoads et al., 2018), atopic wheeze and allergy (Arrieta et al., 2015; Laforest-Lapointe and Arrieta, 2017) and less common, but serious immune mediated disorders such as type-1 diabetes (Vatanen et al., 2016) and Crohn's disease (Hviid et al., 2011). Recent developments in systems immunology now enable us to profile immune system development in humans at the systems level, and map cell-cell regulatory relationships in health and disease (Davis and Brodin, 2018). Further, small blood volumes available from newborn children are no longer prohibitive and as little as 100 µL of whole blood is sufficient for systems-level immunomonitoring (Brodin et al., 2019). Previously, in an analysis of immune development in human newborns, we identified drastic modulation in immune cell and protein abundances that were largely shared among preterm and term children, pointing towards a stereotypic postnatal adaptive response to environmental exposures (Olin et al., 2018). Moreover, gut dysbiosis in a subset of these infants was associated with elevated markers of immune activation in the blood (Olin et al., 2018).

Dysbiosis of the infant gut microbiome is common in modern societies and a likely contributing factor to the increased incidences of immune mediated disorders (Dominguez-Bello et al., 2019; Mohammadkhah et al., 2018; Sonnenburg and Sonnenburg, 2019). Therefore, there is great interest in identifying microbial factors that can support healthier immune system development and potentially prevent some cases of allergy, autoimmunity and possibly other conditions involving the immune system (Renz and Skevaki, 2020). Loss of *Bifidobacterium* early in life is one commonly reported factor in infants more likely to develop autoimmune activity as seen in a birth

cohort in Finland (Vatanen et al., 2016) and atopic wheeze in another cohort in rural Ecuador (Arrieta et al., 2017). Moreover, observational studies have identified a link between the loss of *Bifidobacterium* in infants and enteric inflammation early in life (Henrick et al., 2019; Rhoads et al., 2018).

Human breastmilk contains abundant Human Milk Oligosaccharides (HMOs) and these are not digestible by humans who lack necessary glucosidases (Sela and Mills, 2010). Instead, the maternal energy used to create these complex sugars is justified by the importance of providing a selective nutritional advantage to microbes specialized in metabolizing HMOs and serving an evolutionary important role in the gut of newborn infants. Bifidobacterium longum subspecies infantis (B. infantis) is one example of a bacterial strain adapted to metabolizing HMOs (LoCascio et al., 2010; Sela et al., 2008; Underwood et al., 2015). B. infantis is commonly found in breastfed infants in countries with low incidence of immune mediated disorders, such as Bangladesh (Huda et al., 2014), and Malawi (Grzekowiak et al., 2012), but rare in Europe (Abrahamsson et al., 2014; Avershina et al., 2014; Jost et al., 2012; Roos et al., 2013) and in North America (Azad et al., 2013; Lewis et al., 2015). Strategies for introducing *B. infantis* have been successfully accomplished, such as EVC001 (Evolve BioSystems Inc.), which is able to stably and persistently colonize and achieve a dominant composition in the intestinal microbiome of breastfed infants leading to reduced markers of intestinal inflammation such as fecal calprotectin (Henrick et al., 2019).

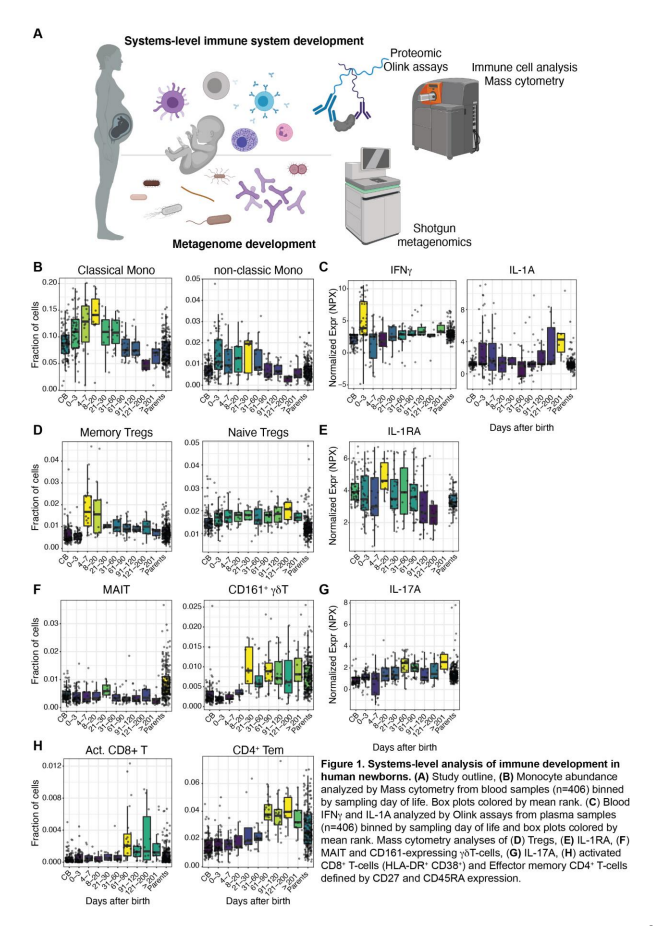
Here, we extend previous findings by combining systems-level analyses of immune development and microbiome colonization in 155 Swedish infants and find that low abundance of gut bifidobacteria is associated with intestinal inflammation and immune dysregulation at three months of life. We also demonstrate a silencing of intestinal inflammation in infants supplemented with *B. infantis* EVC001 (Evolve BioSystems Inc.) in an intervention study in California. Fecal water from infants colonized by *B. infantis* EVC001 skew naïve T-cell polarization *in vitro* from Th2 towards Th1, and modulate Th17 cell states, indicative of a healthier immune system imprinting during the first critical months of life.

RESULTS

Sequential waves of immune cell expansions during the first months of life

We analyzed longitudinal blood samples (n=406) postnatally from infants (n=155) born at the Karolinska University Hospital between April 2014 to December 2019 using a mass cytometry panel with 44 antibodies targeting activation and differentiation markers across all immune cell populations. In total, we profiled 110 immune cell populations from 270 blood samples and quantified 267 unique plasma proteins using using Olink assays (Lundberg et al., 2011) (Olink, Uppsala, Sweden), which elucidated immune responses and developmental changes (Olin et al., 2018) as well as infectious disease susceptibility postnatally (Pou et al., 2019). We also collected paired fecal samples (n=168) performed shotgun metagenomic sequencing from these same children and timepoints (Figure 1A).

When ordering immune cell frequency measurements by day of sampling, we observed an early expansion of classical and non-classical monocytes during the first few days of life (Figure 1B), and a concomitant surge in circulating IFN γ , and a less pronounced increase in IL-1 α (Figure 1C). Starting a few days after this monocyte expansion, a significant increase in memory Tregs was observed uniformly across children, while naïve Tregs abundance remained stable (Figure 1D). This expansion of antigen experienced Tregs indicate a tolerizing response to antigen following the initial innate reaction mediated by monocytes. This memory Treg expansion is also associated with a transient increase in circulating IL-1RA at day 21-30 (Figure 1E), further emphasizing a balanced response to initial innate reactions starting immediately after birth. Gut associated cells such as MAIT cell populations were found at a consistently lower frequency as compared to parents, but with a small transient increase around 1 month of life (Figure 1F).



At this same timepoint a more robust increase in circulating CD161⁺ $\gamma\delta T$ cells was observed and reached cell frequencies similar to adult levels (Figure 1F). Such cells are important producers of IL-17A (Maggi et al., 2010), which was corroborated by increased levels of IL-17A in plasma during the same initial 3-month time frame (Figure 1G). A final activation event involve the expansion of CD4⁺ and activated CD8⁺ $\alpha\beta T$ -cells from 2 months onwards (Figure 1H), and collectively these findings indicate transient immune activation events in early and mid-infancy, likely triggered by colonizing microbes like the weaning reaction described in mice (Nabhani et al., 2019), but noticeably different with respect to timing and immune cell populations involved.

Expansion of mucosal-specific CD4⁺ T-cells in the blood of newborn children

Work in murine model systems and human subjects have revealed that immune cells primed by antigens at mucosal surfaces circulate and are detectable in the peripheral blood. Specifically, memory CD4⁺ T-cells identified by the expression of CD38 and lacking the lymphoid tissue homing marker CD62L are examples of such mucosalspecific T-cells that are abundant in the intestinal tissue but also identifiable in the blood of humans at a frequency of ~4-8% of total CD4+ T-cells (Pré et al., 2011). Here, we identified the same population of CD4⁺ T-cells in the blood of newborn children; however, at a greater abundance of 33.6% and expanding in the blood during the first weeks of life to reach more than 60% of memory CD4⁺ T-cells within the first 100 days of life (Figure 2A). Since these cells have downregulated CD45RA we believe these cells have responded to their native antigen, presumably at the mucosal surfaces and undergone memory phenotype transition. To understand the functional aspects of these T-cells, we sorted these by flow cytometry and performed bulk mRNA-sequencing. We also sorted out total memory CD4⁺ T-cells from the same samples and compared bloodtranscriptional modules (BTMs)(Li et al., 2013) between mucosal-specific memory CD4+ T-cells and total memory CD4⁺ T-cells. The most enriched hallmark pathways in mucosal-specific memory CD4⁺ T-cells were type-I and type-II IFN responses (Figure 2B) and the most upregulated genes in mucosal specific memory CD4⁺ T-cells included the Complement regulatory Factor H (CFH), the cytokine IL-15 important for NK cell

homeostasis as well as the monocyte growth factor CSF1 (M-CSF), suggesting that the mucosal-specific memory CD4⁺ T-cells serving a regulatory function vis-à-vis NK-cells and monocytes in the newborn intestinal mucosa (Figure 2C-D).

Variable colonization of the infant gut after birth

To better understand the expansion of mucosal-specific memory CD4⁺ T-cells and the microbial antigens potentially driving the observed response early after birth, we performed shotgun metagenomic sequencing of longitudinal fecal samples from the same children. We focused on bacterial components and found a highly variable alphadiversity at birth but also a trend towards increasing alpha-diversity over time after birth (Figure 3A). The most variable genera over time were identified partial least square regression analysis vs. days after birth and a latent vector explaining changes over time extracted. The top contributing genera are shown as fraction of reads vs. time after birth and colored by mode of delivery (Figure 3B). Several notable changes are seen, and largely in line with previous literature such as high abundance of *Bacteroides* in vaginally delivered children while staphylococci and streptococci are abundant in cesarean-delivered infants (Shao et al., 2019) (Figure 3B).

There is a drastic expansion of Bifidobacteria occurring after birth in the majority of newborns, and in some infants, all identified reads are mapping to the *Bifidobacterium* genus (Figure 3B), indicating their potential to dominate the gut microbiome early in life, particularly in breastfed infants. We found that multiple species of bifidobacteria expanded early in life in both preterm and term children, in particular *B. longum*, *B. breve*, and *B. animalis* (Figure 3C). Given these findings, the changes in bifidobacteria can be used as a surrogate markers of gut microbiome evolution in the newborn children with three principal patterns seen in our cohort, 1) rare cases of abundant bifidobacteria that are lost over time, 2) the most common pattern of increased abundance of bifidobacteria over time, and 3) children with very rare, or even undetectable levels of bifidobacteria over time during the first three months of life (Figure 3D). Given the known associations between Bifidobacterial abundance and the risk of developing immune mediated diseases in the future (Arrieta et al., 2015, 2017;

Vatanen et al., 2016) we were interested to explore immune system development in children with these different surrogate patterns of microbial colonization.

Immune dysregulation in infants not expanding gut Bifidobacteria

To capture imprinting effects, we focused on blood immune measurements around three months of life involving 268 plasma proteins and all major immune cell populations in blood. We compared these immune system features between infants with (pattern #2) and without bifidobacterial expansion (Pattern #3). We find that 3-month immune cell composition in infants with low gut bifidobacterial abundance was characterized by expanded populations of effector cells and proinflammatory immune cell populations such as proinflammatory monocytes and MAIT-cells, important T-cells in the intestinal immune system compartment and known to respond to bacterial Vitamin B-metabolites in the mucosa (Ioannidis et al., 2020) (Figure 4A).

In contrast, children with abundant gut Bifidobacteria have elevated plasmablast frequencies and also higher frequencies of antigen-experienced regulatory T-cells which are both essential for healthy immune-microbe relationships in the intestine (Smith et al., 2013). The plasma protein analyses revealed similarly upregulated markers of inflammation such as IL-6 and the Oncostatin M (Figure 4A). The latter is notable given results from West et al, implicating the latter protein as a mediator of intestinal inflammation in patients with inflammatory bowel disease (West et al., 2017). To better understand the regulatory relationships in the newborn immune system in these groups of infants, we assessed the cell-cell dependencies. To this end, we calculated Spearman correlation matrices in infants with high levels of *Bifidobacterium* versus low Bifidobacterium abundance, respectively (Figure 4C). Such cell-cell correlation matrices can be informative by revealing coregulated cell populations, and in particular, context-dependent differences to this co-regulation network (Rodriguez et al., 2020). We found that selected cell-cell relationships are shared among high levels of Bifidobacterium versus low Bifidobacterium abundance including strong positive correlation among naïve CD4⁺, naïve CD8⁺ and naïve regulatory T-cells, likely reflecting

the overall thymic output of an individual child (Figure 4C). Other modules of coregulated cell populations were markedly different between the groups of children.

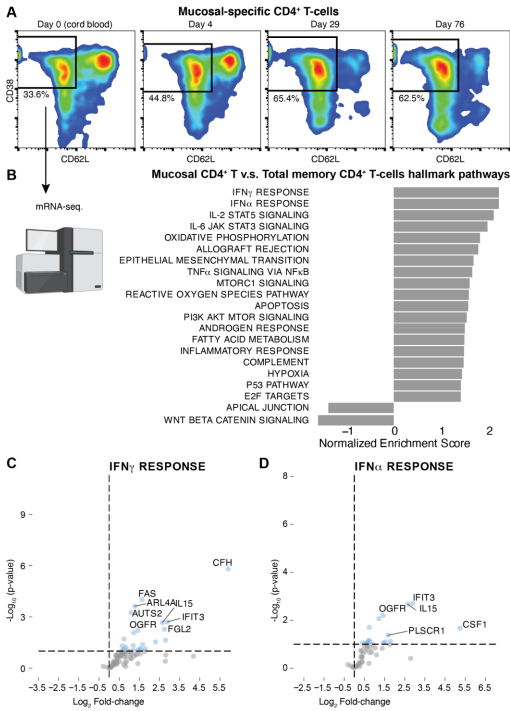


Figure 2. Mucosal specific CD4* T-cells expand and perform type-I and II IFN-responses. (A) Representative FACS plots of CD38* CD62L* CD4* T-cells collected at postnatal day 0, 4, 29 and 76. **(B)** Bulk mRNA-seq. and gene set enrichment analysis showing top enriched hallmark pathways in mucosal-specific v.s. total memory CD4* T-cells. Volcano plots of differentially regulated mRNA transcripts in mucosa-specific CD4* T-cells vs total memory CD4* T-cells within the hallmark pathways **(C)** IFN_Y responses and, **(D)** IFN_α responses.

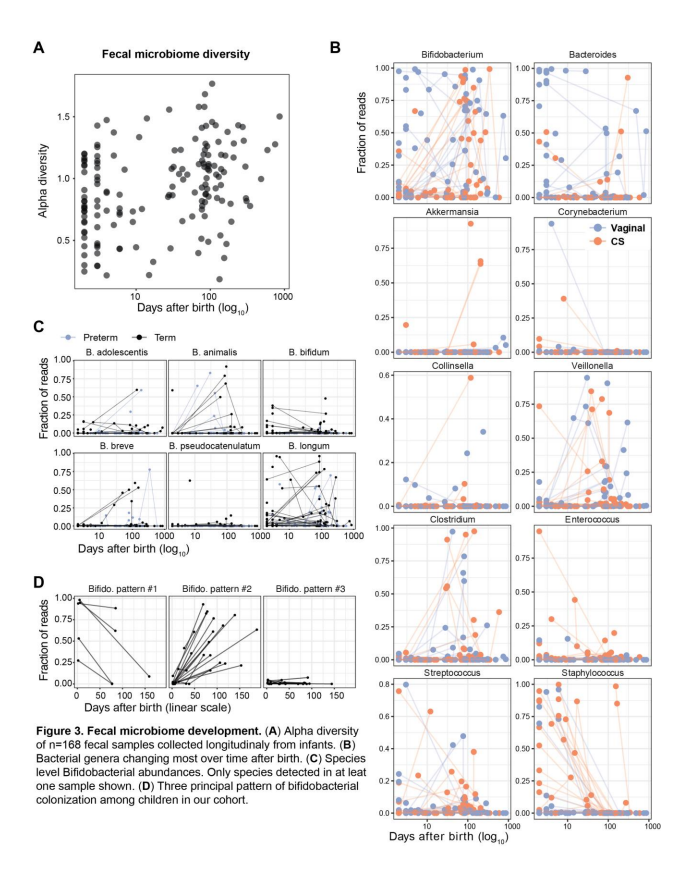
In particular, in infants with higher *Bifidobacterium* abundance, we found that memory-phenotype regulatory T-cells are inversely correlated with the levels of activated CD8⁺ T-cells, proinflammatory monocytes and few additional immune cell populations, whereas this relationship is completely abolished in infants at three months of life, who have low abundance of *Bifidobacterium* (Figure 4C). In contrast, these children have strong positively correlated levels of memory Tregs and activated CD8⁺ T-cells, suggesting ongoing immune activation, rather a homeostatic balance between tolerance and inflammation (Figure 4C). We conclude that infants who are not colonized by bifidobacterial species during their first months of life have an immune system state at three months of life characterized by elevated markers of intestinal inflammation, activated immune cell populations and a perturbed immune cell network.

B. infantis EVC001 supplementation silenced intestinal inflammation in newborn children

In an attempt to improve immune system education and silence the chronic inflammatory responses seen in children devoid of infant-adapted bifidobacterial species, we evaluated fecal cytokines in exclusively breastfed, term infants, half of whom were fed *B. infantis* EVC001 daily from day 7 to day 28 postnatal and half that were not fed a probiotic. Baseline fecal cytokine levels were not significantly different between the two groups; however, infants fed *B. infantis* EVC001 had significantly lower levels of Th2-inducing cytokine, IL-4 and IL-13, proinflammatory cytokine, IL-17A, and regulatory and chemotactic cytokines, IL-21, IL-31, IL-33, and MIP3a by day 60 postnatal (*P*=0.022, 0.009, 0.046, 0.0002, 0.0038, 0.00001, and 0.0038, respectively; Figure 5B). Conversely, IFNβ concentrations were significantly increased in infants fed B. infantis EVC001 (P=0.017; Figure 5B). This suggested a modulatory effect of proinflammatory and Th2 responses by B. infantis EVC001 colonization, so we performed pairwise correlation tests for all 40 individual stool samples at day 60 of life comparing microbial composition and fecal cytokine concentrations (Spearman correlation with Benjamini-Hochberg false discovery rate correction a < 0.02), which identified three taxa, including Clostridiaceae, Enterobacteriaceae, and

Staphylococceae significantly correlated with proinflammatory cytokine production. Specifically, Clostridiaceae abundance correlated with increased IL-21, IL-33, and IL-4, Enterobacteriaceae correlated with increased IL-13, IL-17A, IL-21, IL-33, and IL-4 production, while Staphylococceae correlated with higher levels of IL-21 (Figure 5C). Additionally, we used a Random Forest analysis to identify taxa associated with increased IFNβ production and predictors with the 10 lowest mean depth values confirming that Bifidobacteriaceae, Streptococcaceae, Veillonellaceae, and Staphylococcaceae most often contributed to IFNβ concentration (Supplementary Figure 1A).

We next evaluated whether IFN β concentration at day 60 correlated with early colonization of *Bifidobacteriaceae* at day 21 postnatal. Fecal samples containing *Bifidobacterium* at day 21 had significantly increased levels of IFN β compared to samples that did not contain *Bifidobacterium* by day 60 postnatal (Supplementary Figure 1B), and at a the species level, *B.longum* abundance at day 21 correlated with IFN β levels at day 60 postnatal (Supplementary Figure 1C; P = 0.0028, rho = 0.48). Importantly, no other *Bifidobacterium* species correlated with increased IFN β levels indicating that *B. longum* abundance may drive increased IFN β production. These findings indicate that supplementation with a Bifidobacterial strain, adapted to expand in breastfed infants, may silence proinflammatory responses in the gut and potentially induce a healthier immune system priming.



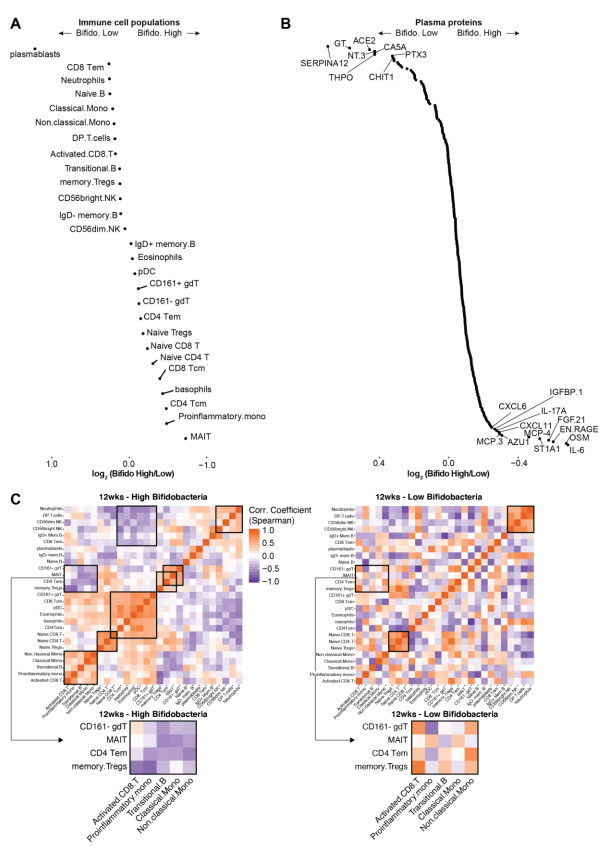


Figure 4. Immune dysregulation in infants with low v.s high bifidobacteria. (A) Fold-change immune cell frequencies at 3 months of life in infants with high v.s. low fecal bifidobacteria. **(B)** Fold-change plasma protein levels at 3 months of life in infants with high v.s. low fecal bifidobacteria. **(C)** Spearman correlation matricies of immune cell frequencies at 3 months of life in children with high v.s. low fecal bifidobacteria. Black boxes highlight modules of co-regulated immune cell populations.

B. infantis EVC001 metabolites influence T-cell polarization

Classically, it is suggested that newborn infants have an immune system biased towards Th2 responses rather than Th1 proinflammatory responses, yet this is mostly based on data from cord blood cells collected prior to any postnatal environmental exposures, or mouse model systems (Brodin, 2020). Our evaluation of enteric cytokines showed significantly decreased Th2 cytokines, IL-4 and IL-13, Th17 cytokine IL-17A, as well as increased IFNβ concentrations in infants colonized with *B. infantis* EVC001 compared to those that were not; therefore, we investigated the potential for microbial metabolites and enteric cytokines from two infant populations that were exclusively fed human milk but were discordant for *B. infantis* colonization to skew T-cell polarization in vitro. We flow-sorted naïve CD4⁺ T-cells from a healthy adult donor and polarized these cells using standard cytokine combinations (Cano-Gamez et al., 2020) in combination with fecal waters (1:100 dilution) collected from newborns fed B. infantis EVC001 or control children lacking B. infantis (Figure 6A). To evaluate the polarization and cellular states of the T-cells in the cultures we applied a targeted multiomics approach (Rhapsody, BD Biosciences) in which we combined the analysis of 259 mRNA by targeted sc-mRNA-sequencing and 10 proteins using oligo-coupled antibodies (BD Abseq[™], BD Biosciences)(Mair et al., 2020). In this way we were able to assess transcription factor and cytokine gene expression often more difficult to detect in untargeted sc-mRNA-sequencing applications, and although the overall UMAP distribution of cells is similar across conditions, subtle differences in cell states were observed (Figure 6B). Using a graph abstraction and clustering method, PAGA (Wolf et al., 2019), we found that T-cell states differ by polarizing condition as expected, but we also observed differences between cells cultured in the presence of B. infantis EVC001 fecal water compared to fecal water lacking *B. infantis* (Figure 6C). Specifically, Th1, Th2 and iTregs are largely comparable between B. infantis EVC001 and no B. infantis fecal water (Figure 6C). Conversely, Th0 cells cultured without additional polarizing cytokines and no B. infantis fecal water induced a cell state similar to that of Th2polarized cells, while the *B. infantis* EVC001 induced a Th0 cell state that more closely reflect Th1-induced conditions (Figure 6C). Furthermore, we assessed the differentially regulated genes among cultured Th0 cells exposed to fecal waters from the two

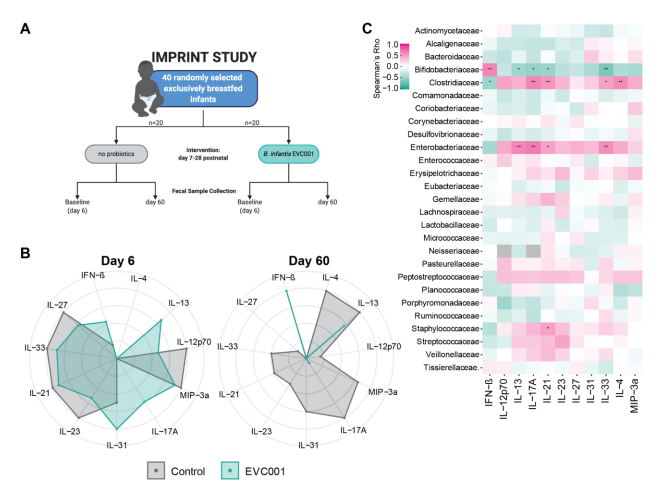


Figure 5. B. infantis EVC001 supplementation is associated with reduced intestinal inflammation. (A) IMPRINT study design and randomization, **(B)** Fecal cytokines at baseline (Day 6) and post-treatment with B.infantis EVC001, or no supplementation. **(C)** Spearman correlation coefficients between fecal cytokine levels and bacterial abundance. *P=0.05. **p=0.01.

different groups and found upregulation of GZMA, GZMB, TNF and STAT1 in the cells skewed towards a Th1-phenotype after exposure to *B. infantis* EVC001 fecal water while IL23R expression in the cells exposed to no *B. infantis* fecal water, similar to the Th2-like cells (Figure 6D). These findings suggest *B. infantis* EVC001 specific metabolites and/or enteric cytokines induced in the intestine by the presence of *B. infantis* EVC001 polarizes naïve CD4⁺ T-cells towards the Th1 lineage, and are of particular interest given the association between Th2-type cells and the development of immune-mediated diseases such as asthma and allergies in young children in which a polarizing effect away from Th2 towards Th1 might prove beneficial in the future.

Apart from the skewing of Th0 cells by fecal water, we also saw a slight difference in the cell states among Th17-polarized cells when comparing *B. infantis*

EVC001 and no *B. infantis* fecal water effects (Figure 6C). Specifically, naïve T-cells polarized towards Th17 in the presence of no *B. infantis* fecal water expressed elevated markers of activation and proliferation (Ki67) compared to naïve T-cells polarized towards Th17 in the presence of *B. infantis* EVC001 fecal water (Figure 6E). Collectively, these results imply that during the first weeks of life there are transient immune activation events, centered on the mucosal surfaces, primarily in the gut. The colonization and formation of the gut microbiome plays an integral role in this process and is likely, itself influenced by this process. Specific bacteria, such as *B. infantis* EVC001, influence this process by dampening intestinal inflammation, shaping immune cell-cell regulatory relationships, and polarizing T-cells away from Th2 to Th1-type cells, with potential long-term consequences for immune imprinting and the risk of developing immune-mediated diseases.

Discussion

It is increasingly clear that early-life gut dysbiosis influences newborn children's risk of developing autoimmune and allergic diseases later in life; however, the exact mechanisms driving perturbations in the developing immune system have not been fully elucidated. Results here confirm previous findings that highlight a stereotypic postnatal adaptive immune response to environmental exposures, and extend to show an immunological sequence of events, triggered by microbial colonization, resulting in altered patterns of intestinal inflammation. In particular a lack of bifidobacterial early in life is associated with elevated markers of intestinal inflammation and a perturbed immune cell regulatory network.

The transient immune cell expansion events described here are reminiscent of the transient reactions to gut microbes in mice at the time of weaning, termed "the weaning reaction" (Nabhani et al., 2019). It has been postulated that such timed reactivity to the colonizing microbes in mice is regulated by Goblet cell-associated passages (GAPs) and serve important functions in establishing a healthy immune-microbe interface in the gut and the induction of tolerance (Knoop et al., 2017). There are several notable differences between these transient events reported in mice and the

process uncovered in human newborns in our current report. First, the timing of transient immune cell reactions is completely different and uncoupled from weaning since the breastfeeding regimen differs between children and most breastfed children in our cohort continue to nurse beyond the three-month window during which the dramatic immune cell changes occur. Secondly, the type of reactions seen in human newborns seem qualitatively distinct and involve different cell populations as compared to the changes seen in mice. This is not only due to the difference between blood-focused analyses in human newborns and intestinal analyses performed in mouse model organisms (Nabhani and Eberl, 2017, 2020; Nabhani et al., 2019), because when analyzing blood immune cell changes in mice, the events remain distinct from human immune cell developmental changes (Brodin lab, manuscript in preparation). The comparison between human newborns and young mice colonized with a more natural gut microbiome will be informative and likely more similar in this regard as shown for other immune system processes (Rosshart et al., 2017).

Furthermore, we report that feeding *B. infantis* EVC001 modulated of immune dysregulation by silencing intestinal Th2 and Th17 immune responses, while inducing IFNβ production, and in vitro its metabolites skewed T-cell polarization *in vitro*, from Th2, towards Th1 phenotype. Thus, given that the gut microbiome seems to play a major role in early immune system development, feeding *B. infantis* to breastfed infants may improve immune imprinting during the first critical months of life. No molecular mechanisms have been identified that directly link microbiome composition or function with the key steps of immune development in humans; however, infants that lack *Bifidobacterium* are more likely to develop autoimmune and asthmatic diseases (Arrieta et al., 2015, 2017; Vatanen et al., 2016). One proposed candidate mechanism involves the over-production of Th17-inducing cytokines leading to increased numbers of Th17 cells as one mechanism of elevated disease risk (Guo et al., 2008; Harrington et al., 2005).

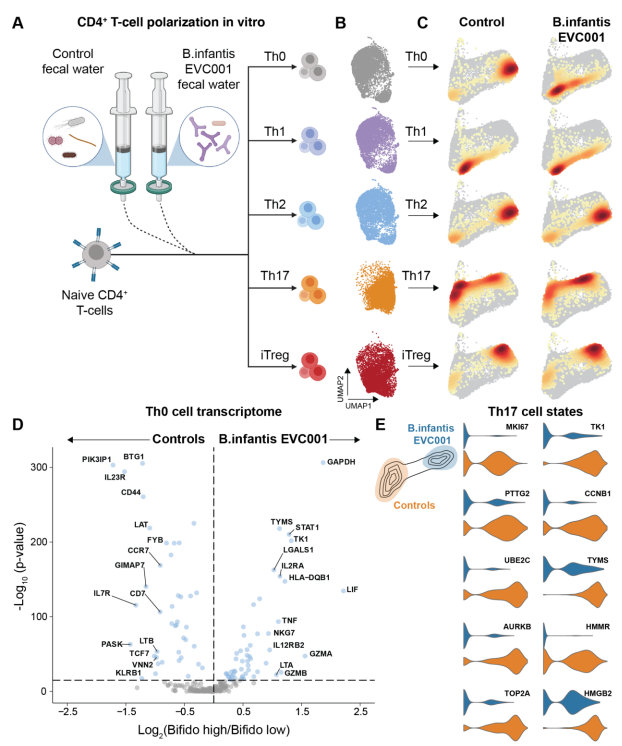


Figure 6. CD4* T-cell polarization under the influence of microbial metabolites. (A) CD4* T-cell polarization in vitro in the presence of fecal water from infants supplemented with B.infantis EVC001 or control (No supplement). **(B)** UMAP plots of polarized T-cells analyzed by targeted sc-mRNA-seq. **(C)** PAGA plots of T-cells polarized in the presence of fecal water from fecal water from infants given B.infantis EVC001 supplementation or control. Coloring by cell density from grey (low) to red (high). **(D)** Volcano plot showing differentially expressed mRNA in Th0 cells cultured with fecal water from infants given B.infantis EVC001 supplementation or control. **(E)** Top 10 differentially expressed mRNA in Th17 cell states induced by fecal water from infants given B.infantis EVC001 supplementation or control.

Here we report additional possible mechanisms of immune system imprinting, possibly influencing the risk of developing immune mediated diseases later in life. Our data indicated low abundance of Bifidobacterium colonization correlated with increased enteric inflammation both in fecal water and in circulation and is associated with a perturbed cell-cell regulatory network. In particular, memory Tregs frequency was inversely correlated with proinflammatory monocytes and activated T-cell populations in children with abundant *Bifidobacterium*, but this regulatory relationship is lost in children devoid of such beneficial microbes. Indeed, Bifidobacteria are efficient producers of SCFA (Frese et al., 2017), which are in turn important inducers of regulatory T-cells in the gut (Smith et al., 2013). Additionally, we uncovered another possible mechanism of beneficial imprinting by beneficial Bifidobacteria. Breastfed infants supplemented with B. infantis EVC001 upregulated intestinal IFNB, which is known to and promote the development of regulatory T cells (Kotredes et al., 2017). Moreover, IFNβ inhibits Th17 cell differentiation and consequently lowers IL-17 production (Durelli et al., 2009; Ramgolam et al., 2009) and children supplemented with B. infantis EVC001 had significantly lower levels of fecal IL-17A as compared to control children. Exogenous IFNβ is given to patients with Multiple Sclerosis (MS) and is believed to improve Tregmediated immunological tolerance (González-Navajas et al., 2012), inhibit IL-23dependent Th17 cell expansion (Harrington et al., 2005) and TGB-β/IL-6-mediated Th17 development (Guo et al., 2008). Additionally, there is preclinical evidence of Treg cell mediated protection in a rodent colitis models whereby both intestinal inflammation and cytokine secretion is decreased by exogenous IFNβ (González-Navajas et al., 2012). Collectively these data indicate a previously unappreciated role for IFNB in early life immune system imprinting by beneficial microbes such as B. infantis.

Microbially-derived metabolites are also likely to be critical in immune system development (Lavelle and Sokol, 2020). Indeed, perturbations to the gut microbiome early in life due to antibiotic use lead to changes in the production of bacterial metabolites such as short-chain fatty acids (SCFAs) and other microbial-derived metabolites involved in immune maturation and homeostasis, as well as host energy metabolism, and intestinal integrity (Lavelle and Sokol, 2020). For example, 12-13

DIHOME, a microbial-derived metabolite has recently been shown to correlate with lower risk of asthma development in susceptible infant populations (Levan et al., 2019). Moreover, indole-3-lactic acid (ILA), which is produced in high concentration in breastfed infants colonized with B. infantis has been shown to decrease enteric inflammation through activation of Aryl hydrocarbon receptor and NF erythroid 2-related factor (Nrf2) (Ehrlich et al., 2018; Laursen et al., 2020; Meng et al., 2020). Importantly, differences in *B. infantis* HMO utilization loci have been identified (Albert et al., 2019) and strains missing specific genes involved in HMO utilization are unlikely to confer the same benefits to infants as we observed here, as the conversion of indigestible HMOs to infant accessible short chain fatty acids and indole-3-lactic acid are key functions of a healthy gut microbiome (Duar et al., 2020). Similarly, acetate and propionate have been implicated in the reduction of lung inflammation in animal models (Trompette et al., 2014) and development of food allergy in human children (Sandin et al., 2009). Notably, a previous intervention study showed significantly increased SCFAs and lactate production in infants fed B. infantis (Frese et al., 2017); due to its unique evolution to consume human milk oligosaccharides (HMOs)(Sela et al., 2008) and a recent study of preterm infants supplemented with Bifidobacterium and Lactobacillus showed increased concentrations of acetate and lactate as compared to controls not given any microbial supplementation (Alcon-Giner et al., 2020).

Finally, the finding that fecal water from children supplemented with *B. infantis* EVC001 influence T-cell polarization *in vitro* towards Th1 cell states, while fecal water from control children induce Th2-like phenotypes is intriguing and suggest that specific metabolites exert a skewing effect on naïve CD4 $^+$ T-cells. This effect is probably not mediated by IFN β alone since cultures supplemented with IFN β did not show this same pattern (Supplementary Figure 2). Instead, we hypothesize that IFN β may provide a tolerizing effect in already polarized T-cells. Future metabolomic profiling of the fecal water and functional testing during T-cell polarization, including analyses of epigenetic changes during polarization will be valuable to mechanistically understand this imprinting effect further.

Collectively, the data provided here extends our current understanding of immune system imprinting by beneficial microbes such as *B. infantis* EVC001 and proposes possible explanations for the known differences in risk of developing immune mediated diseases associated with such early life imprinting effects. Randomized controlled trials targeting these additional mediators and mechanisms proposed will be required to establish a way forward towards preventive interventions to lower the burden of immune mediated diseases such as asthma, allergies, and autoimmunity in the future.

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STAR Methods

Born-Immune newborn cohort study

The study was performed in accordance with the declaration of Helsinki and the study protocol was approved by the regional ethical board in Stockholm, Sweden (DNR: 2009/2052 – 31/3 & 2014/921-32). After obtaining informed consent from parents, blood samples from newborns and parents were collected at the Karolinska University Hospital. We also collected fecal samples from infants, either at the time of clinical visits and frozen directly at -80 C or collected at home frozen at -20 C and brought to the clinic by parents. Clinical metadata such as mode of delivery, nutrition, growth and medications were gathered in a clinical database.

Blood immune cell profiling by Mass cytometry

Blood samples drawn from newborns and parents were mixed with a stabilizer (Brodin et al., 2019) (one of the components of Whole blood processing kit; Cytodelics AB, Sweden) either immediately or within 1-3 hours post blood draw and cryopreserved as per the manufacturer's recommendations. Samples were thawed, and cells were fixed/RBCs lysed using WASH # 1 and WASH # 2 buffers (Whole blood processing kit; Cytodelics AB, Sweden) as per the manufacturer's recommendations. This was performed a few days prior to barcoding and staining of cells. Post fix/lysis of cells, ~1-2x10⁶ cells/sample were plated onto a 96 well round bottom plate using standard cryoprotective solution (10% DMSO and 90% FBS) and cryopreserved at -80°C. At the time of experimentation, cells were thawed at 37°C using RPMI medium supplemented with 10% fetal bovine serum (FBS), 1% penicillin-streptomycin and benzonase (Sigma-Aldrich, Sweden). Briefly, cells were barcoded using automated liquid handling robotic system (Agilent technologies)(Mikes et al., 2019) using the Cell-ID 20-plex Barcoding kit (Fluidigm Inc.) as per the manufacturer's recommendations. Samples were pooled batch wise by keeping together the longitudinal samples from each new born baby or parent in the same batch. Cells were then washed, FcR blocked using blocking buffer (in-house developed recipe) for 12 min at room temperature, following which cells were

incubated for another 30 min at 4°C after addition of a cocktail of metal conjugated antibodies targeting the surface antigens. Cells were washed twice with CyFACS buffer (PBS with 0.1% BSA, 0.05% sodium azide and 2mM EDTA) and fixed overnight using 2% formaldehyde made in PBS (VWR, Sweden). The broad extended panel of antibodies used are listed in Supplementary Table 1. For acquisition by CyTOF, cells were stained with DNA intercalator (0.125 μM Iridium-191/193 or MaxPar® Intercalator-Ir, Fluidigm) in 2% formaldehyde made in PBS for 20 min at room temperature. Cells were washed twice with CyFACS buffer, once with PBS and twice with milliQ water. Cells were mixed with 0.1X Norm Beads (EQTM Four Element Calibration Beads, Fluidigm) filtered through a 35μm nylon mesh and diluted to 1000,000 cells/ml. Samples were acquired using super samplers connected to our CyTOF2 mass cytometers (Fluidigm Inc.) using CyTOF software version 6.0.626 with noise reduction, a lower convolution threshold of 200, event length limits of 10-150 pushes, a sigma value of 3, and flow rate of 0.045 ml/min.

Antibodies and reagents for mass cytometry

The panel of monoclonal antibodies used for this study are indicated in the Key Resources Table. Monoclonal antibodies were either purchased pre-conjugated from Fluidigm or obtained in carrier/protein-free buffer as purified antibodies that were then coupled to lanthanide metals using the MaxPar X8 polymer conjugation kit (Fluidigm Inc.) as per the manufacturer's recommendations. Following the protein concentration determination by measurement of absorbance at 280nm on a nanodrop, the metallabeled antibodies were diluted in Candor PBS Antibody Stabilization solution (Candor Bioscience, Germany) for long-term storage at 4°C.

Born-immune Plasma protein profiling

Plasma protein data was generated using Olink assays, a proximity extension assay (Olink AB, Uppsala)(Lundberg et al., 2011). For analysis, 20µL of plasma from each sample was thawed and sent for analysis, either at the plasma protein profiling platform, Science for Life Laboratory, Stockholm or Olink AB in Uppsala. In these assays, plasma proteins are dually recognized by pairs of antibodies coupled to a cDNA-strand that ligates when brought into proximity by its target, extended by a polymerase and

detected using a Biomark HD 96.96 dynamic PCR array (Fluidigm Inc.). Three Olink panels have been used as indicated in Key Resources Table, capturing a total of 267 unique proteins in each plasma sample.

Born-immune fecal metagenomics

DNA was extracted as in IHMS DNA extraction protocol #8 (Costea et al., 2017). According to protocol recommendations 0,2g of faeces were used. Briefly, samples were treated with lysozyme solution and subjected to bead beating using zirconium beads. After centrifugation, DNA is extracted from supernatants with QIAamp Fast DNA Stool Mini Kit (Qiagen, Cat No. 51604). After DNA extraction, collected DNA was quantified with Qubit (ThermoFisher, Cat No. Q32851) and 10ng were subjected to mechanical fragmentation with the Covaris Focused-ultrasonicator to ensure that fragment sizes were compatible with Illumina sequencing (~300bp average). Sequencing adapters and sample barcodes were incorporated to the DNA fragments using ThruPLEX DNA-seg kit (Rubicon Genomics, Cat No. R400406). ThruPLEX DNAseg products were purified and size selected by AMPure beads (Beckman Coulter, Cat No. B23318), and DNA concentration and size distribution were inspected with the Qubit dsDNA HS Assay Kit (ThermoFisher, Cat No. Q32851) and the Agilent 2100 Bioanalyzer High Sensitivity kit (Agilent Technologies, Cat No. 5067-4626), respectively. Purified ThruPLEX DNA-seq products were then equimolarly pooled in 4 lanes and subjected to NovaSeg 6000 S4 Illumina Sequencing at the Science for Life Laboratory, Stockholm, Sweden.

IMPRINT study design and cytokine quantification

A total of 80 fecal samples (ClinicalTrials.gov: NCT02457338) from 40 subjects were analyzed at three time points—day 6 (Baseline), day 40, and day 60 postnatally. Individual subjects were chosen at random and made up a subset of the original study participants. All aspects of the study were approved by the University of California Davis Institutional Review Board (IRB Number: ID 631099) and all participants provided written informed consent. Details of the study design and procedures used to collect these samples has been reported (Frese et al., 2017). Briefly, exclusively breastfed

term infants were randomly selected to receive 1.8 × 10¹⁰ colony-forming units (CFU) B. infantis EVC001 daily for 21 days (EVC001) starting at day 7 postnatal or to receive breast milk alone (control) and followed up to postnatal day 60 (Frese et al., 2017). All mothers received lactation support throughout the study. The demographic information (e.g., age, sex, and gestational age) was collected from each participant. Here stool samples from randomly selected infants who were fed EVC001 (n = 20) and control infants (n = 20) on days 6 (Baseline) and 60 postnatal were collected and quantified from 80 mg of frozen stool diluted 1:10 in Meso Scale Discovery (MSD; Rockville, MD) diluent using the U-PLEX Inflammation Panel 1 (human) Kit. Samples were measured in duplicate and blank values were subtracted from all readings according to the manufacturer's instructions. All detectable biomarker values were included as continuous data in the analyses; however, values below level of detection (< 30% of all cytokines measurements) were generated below the level of quantification to justify parametric statistics. Fecal cytokine concentrations were determined using calibrations curves to which electrochemiluminescence signals were backfitted. Final concentrations were calculated using the Sector Imager 2400 MSD Discovery Workbench analysis software.

IMPRINT study – fecal microbiome analyses

DNA was extracted from 296 stool swab samples stored in DNA/RNA shield lysis tubes (Zymo Research, Irvine CA) using the ZymoBIOMICS 96 MagBead DNA kit (Zymo Research). Extracted DNA was quantified using QuantIT dsDNA Assay kit, high sensitivity (ThermoFisher Scientific, Waltham, MA) according to the manufacturer's protocol. 3 samples were omitted from downstream analysis due to failure to meet input requirements for library preparation. Libraries were prepared for each sample using the Illumina Nextera DNA Flex library kit (Illumina, San Diego, CA) with unique dual indexes according to manufacturer guidelines. Libraries were pooled and submitted to UC Davis DNA Technologies core for sequencing on the Illumina NovaSeq S4 flow cell. (Illumina, San Diego, CA). Each lane of the S4 flow cell contained 96 libraries.

IMPRINT-Absolute quantification of *B. infantis* by Quantitative Real-Time PCR

Quantification of the total *B. infantis* was performed by quantitative real-time PCR using Blon_2348 sialidasetgene primers Inf2348F (5'- ATA CAG CAG AAC CTT GGC CT -3'), Inf2348_R (5'- GCG ATC ACA TGG ACG AGA AC -3'), and Inf2348_P (5'- /56-FAM/TTT CAC GGA /ZEN/TCA CCG GAC CAT ACG /3IABkFQ/-3')(Lawley et al., 2017) Each reaction contained 10µL of 2× TaqMan Universal Master Mix II with UNG master mix (Applied Biosystems), 0.9 µm of each primer, 0.25 µM probe and 5 µL of template DNA. Thermal cycling was performed on a QuantStudio 3 Real-Time PCR System and consisted of an initial UNG activation step step of 2 minute at 50 followed by a 10 minute denaturation at 95°Csucceeded by 40 cycles of 15 s at 95 °C and 1 min at 60°C. Standard curves for absolute quantification were generated using genomic DNA extracted from a pure culture of *B. infantis* EVC001.

IMPRINT-Quality filtering and removal of human sequences

Demultiplexed fastq sequences were quality filtered, including adaptor trimming using Trimmomatic v0.36 (Czajkowski et al., 2018) with default parameters. Quality-filtered sequences were screened to remove human sequences using GenCoF v1.0 (Czajkowski et al., 2018) against a non-redundant version of the Genome Reference Consortium Human Build 38, patch release 7 (GRCh38_p7; www.ncbi.nlm.nih.gov). Human sequence-filtered raw reads were deposited in the Sequence Read Archive (SRA; https://www.ncbi.nlm.nih.gov/sra) under the reference number, PRJNA390646.

IMPRINT-Fecal water preparation

Historical fecal samples from term infants either colonized with *B. infantis* EVC001 or not were collected and stored in -80C until processing. Pooled fecal samples (minimum of 3) from infants colonized with *B. infantis* EVC001 or not were weighed and diluted 25% w/v in sterile PBS and vortexed for 1 min allowing stool to thaw and form a homogenous slurry. Fecal slurries were then centrifuged for 30min at 4,000 RPM at 4C. The supernatant was collected and spun again for 3 hours at 12,000 RPM at 4C. The supernatant was further collected and then serially filtered (40um cell strainer, 1um, .45um, and .22um). Filtered fecal waters were stored in -80C until use.

IMPRINT-fecal cytokine measurements

Interleukin (IL)-1 β , IL-2, IL-4 IL-5, IL-8, IL-10, IL-13, IL-17A, IFN γ , and TNF α were quantified from 80 mg of stool diluted 1:10 in Meso Scale Discovery (MSD; Rockville, MD) diluent using the U-PLEX Inflammation Panel 1 (human) Kit according to the manufacturer's instructions as previously published (ref). Standards and samples were measured in duplicate and blank values were subtracted from all readings. The concentration of fecal calprotectin and MPO were quantified using MSD R-PLEX Human Calprotectin Antibody Set. The samples were plated in duplicate and the assay was performed twice. The plates were then read on a Sector Imager 2400 MSD Discovery Workbench analysis software.

IMPRINT - Statistical Analysis

All statistical analyses were performed in R v3.6.2. A Kruskal-Wallis one-way analysis of variance coupled with an FDR or Bonferroni correction was used for statistical comparisons between individual genes, cytokines and taxa amongst groups. Statistical analysis to assess total resistome or enterotype composition by group was performed using a Mann-Whitney or Holm-adjusted Dunn's test. Rarefaction curves were computed to estimate the diversity of the identified ARGs across samples. A nonparametric two-sample t-test was used to compare rarefaction curves using Monte Carlo permutations (n=999). Enterotype analysis was performed as previously described (Arumugam et al., 2011). Cytokines and *B. infantis*-specific qPCR were correlated with the Spearman method with FDR correction. The P-values throughout the manuscript are represented by asterisks (*, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.001). A random forest model was created to identify taxa most often used in prediction of IFN concentration. Family-level taxa relative abundance at days 40 and 60 and age (day of life) were used as predictors. The depth at which each taxon appeared within a tree was collected and averaged across the ensemble of 500 trees.

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AUTHOR CONTRIBUTIONS

BMH and PB conceived the study. BMH, JTS, MAU, JBG, SAF, and PB designed the study. RDM, SC, JP, and HKB generated data presented in the manuscript. BMH, JTS, JBG, SAF, and PB analyzed the data. BMH, AME, and PB wrote the manuscript. All authors approved the final manuscript for publication.

DECLARATION OF INTERESTS

PB, JM and TL are founders and shareholders of Cytodelics AB (Stockholm, Sweden). PB is an advisor to Scailyte AG (Zurich, Switzerland). RDM, SC, JP, HKB, SAF, and BMH are employees of Evolve BioSystems, a company focused on restoring the infant microbiome. JTS received funding to conduct the IMPRINT trial and AME received funding to assist in writing the manuscript. JBG is a co-founder of Evolve BioSystems. SAF and BMH serve as Adjunct Assistant Professors in Food Science & Technology Department, University of Nebraska Lincoln.



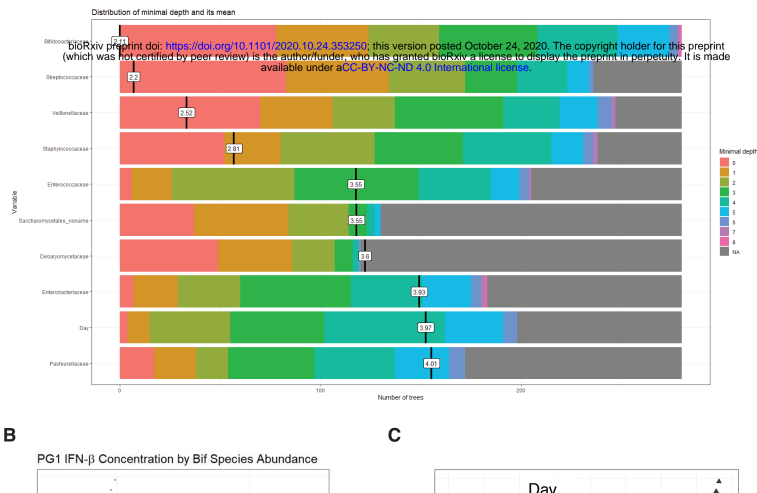
KEY RESOURCES TABLE

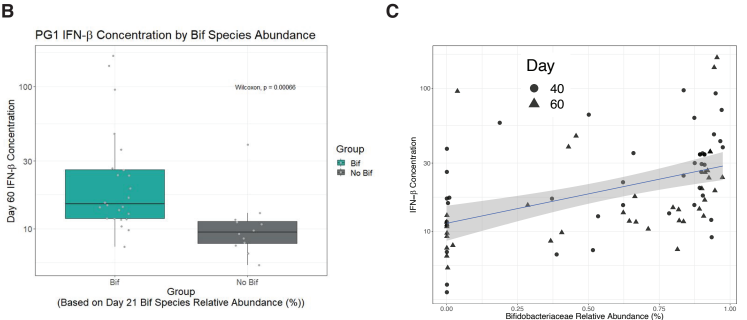
REAGENT or RESOURCE	SOURCE		IDENTIFIER			
Software and Algorithms						
Seven Bridges Platform	_		https://ww	ww.sevenbridges.com/platform/		
R 4.0.2	R C	R Core Team, 2020 https://www.r-project.org/				
Python 3.7		, , , , , , , , , , , , , , , , , , ,	https://www.python.org/			
Seurat			https://cra project.or html	an.r- g/web/packages/Seurat/index.		
DESeq2	Love	ove et al., 2014 https://bioconductor.org/packages/relse/bioc/html/DESeq2.html				
tmod 0.46.2				an.r-project.org/package=tmod		
kallisto	Bray	y et al., 2016	https://github.com/pachterlab/kallisto			
Scanpy	Wol	f et al., 2018	https://github.com/theislab/Scanpy			
PAGA	Wol	f et al., 2019	https://github.com/theislab/paga			
BD Rhapsody antibodies and reagents						
BD AbSeq Hu CCR7(CD197)	BD		Cat # 940394			
BD AbSeq Hu CD137	BD		Cat # 940055			
BD AbSeq Hu CD28	BD		Cat # 940017			
BD AbSeg Hu CD11a	BD		Cat # 940077			
BD AbSeq Hu TCR galma/delta	BD		Cat # 940057			
BD AbSeq Hu CD103	BD		Cat # 940067			
BD AbSeq Hu CD27	BD		Cat # 940018			
BD AbSeq Hu CD39	BD		Cat # 940073			
BD AbSeq Hu CD62L	BD		Cat # 940041			
BD AbSeq Hu CD4	BD	Cat # 940001		0001		
BD AbSeq Hu CD8	BD		Cat # 940003			
BD AbSeq Hu CD3	BD		Cat # 940000			
BD AbSeq Hu CD16	BD	Cat # 940006		0006		
BD AbSeq Hu CD56	BD		Cat # 940007			
BD AbSeq Hu CD161	BD		Cat # 940070			
BD AbSeq Hu CD45RA	BD		Cat # 940011			
BD AbSeq Hu CD38	BD		Cat # 940013			
BD AbSeq Hu HLA-DR	BD		Cat # 940010			
BD single cell multiplexing kit	BD		Cat # 633781			
BD Rhapsody cartridge reagent kit	BD		Cat # 633731			
BD Rhapsody cDNA kit	BD		Cat # 633773			
BD Rhapsody targeted mRNA and AbSeq amplification kit	BD		Cat # 633774			
BD Pharmingen Stain Buffer (FBS)	BD		Cat # 554656			
DRAQ7	BD C			Cat # 564904		
BD Rhapsody Human T-Cell Expression Panel	BD C		Cat # 633	Cat # 633751		
Critical Commercial Assays						
irconium beads 0,1 mm Qiag		Qiagen		13118-400		
Lysozyme	ThermoFis			89833		
Proteinase K		Qiagen		19133		
ThruPLEX® DNA-seq 96D	96D Takara			R400676		

Cell²res

Agencourt AMPure XP	Beckmann Coulter	<u>A63881</u>
Qubit™ Assay Tubes,	ThermoFisher	Q32856
Qubit™ dsDNA HS Assay Kit	ThermoFisher	Q32854
QIAamp Fast DNA Stool Mini Kit	Qiagen	51604
Agilent High Sensitivity DNA Kit	Agilent	5067-4626
96 microTUBE Plate	Covaris	520078

Metal tag	Marker	Clone	Vendor
89Y	CD45	HI30	Fluidigm
113ln	CD57	HCD57	BioLegend
115ln	HLA-A,B,C	W6/32	BioLegend
141Pr	CD49d	9F10	Fluidigm
142Nd	CD19	HIB19	Fluidigm
143Nd	CD5	UCHT2	BioLegend
144Nd	CD16	3G8	BioLegend
145Nd	CD4	RPA-T4	BioLegend
146Nd	CD8a	SK1	BioLegend
147Sm	CD11c	Bu15	Fluidigm
148Nd	CD31	WM59	BioLegend
149Sm	CD25	2A3	Fluidigm
150Nd	CD64	10.1	BioLegend
151Eu	CD123	6H6	BioLegend
152Sm	TCRgd	5A6.E9	Fischer Scientific
153Eu	Siglec-8	837535	R&D Systems
154Sm	CD3e	UCHT1	BioLegend
155Gd	CD33	WM53	BioLegend
156Gd	CD26	BA5b	BioLegend
157Gd	CD9	SN4 C3-3A2	eBiosciences
158Gd	CD34	581	BioLegend
159Tb	CD22	HIB22	BioLegend
160Gd	CD14	M5E2	BioLegend
161Dy	CD161	HP-3G10	BioLegend
162Dy	CD29	TS2/16	BioLegend
163Dy	HLA-DR	L243	BioLegend
164Dy	CD44	BJ18	BioLegend
165Ho	CD127	A019D5	Fluidigm
166Er	CD24	ML5	BioLegend
167Er	CD27	L128	Fluidigm
168Er	CD38	HIT2	BioLegend
169Tm	CD45RA	HI100	Fluidigm
170Er	CD20	2H7	BioLegend
171Yb	CD7	CD7-6B7	BioLegend
172Yb	IgD	IA6-2	BioLegend
173Yb	CD56	NCAM16.2	BD Pharmingen
174Yb	CD99	HCD99	BioLegend
175Lu	CD15	W6D3	BioLegend
176Yb	CD39	A1	BioLegend
191Ir	DNA Ir	Cell-ID DNA Intercalator	Fluidigm
193Ir	DNA Ir	Cell-ID DNA Intercalator	Fluidigm
209Bi	CD11b	Mac-1	Fluidigm

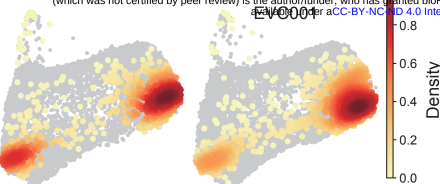




Supplementary Figure 1. Related to Figure 5. Gut microbes associated with fecal cytokine levels. (**A**) Random Forest Model output. (**B**) Box plot of day 60 fecal IFNb abundances. (**C**) Correlation (Pearson) between day 60 relative abundance of Bifidobacteria and fecal IFNb, r=0.55, p=8.9 x 10-8

Th0 cultured with IFN $\!\beta$

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Supplementary Figure 2. Related to Figure 6. CD4 $^+$ T-cell polarization under the influence of microbial metabolites and IFN β . CD4 $^+$ T-cell polarization in vitro in the presence of fecal water from children given B.infantis EVC001 or control and additional IFN β . T-cells at the end of the culture shown as 2D PAGA plots and separated by culture condition.