**Title: Gut microbiota development during infancy: impact of introducing allergenic foods**

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**Abstract**

**Background:** The gut microbiota potentially plays an important role in the immunological education of the host during early infancy.

**Objective:** We sought to determine how the infant gut microbiota evolve during infancy,

particularly in relation to hygiene-related environmental factors, allergic and atopic disorders, and a randomized introduction of allergenic solids.

**Methods:** 1,303 exclusively breastfed infants were enrolled in a dietary randomized controlled trial (EAT study) from three months of age. In this nested longitudinal study, fecal samples were collected at baseline with additional sampling of selected cases and controls at six and twelve months to study the evolution of their gut microbiota, using *16S ribosomal RNA* gene targeted amplicon sequencing.

**Results:** In the 288 baseline samples from exclusively breastfed infant at three months of age, the gut microbiota were highly heterogenous, forming three distinct clusters: *Bifidobacterium*-rich, *Bacteroides*-rich and *Escherichia*/*Shigella*-rich. Mode of delivery was the major discriminating factor. Increased *Clostridium sensu stricto* relative abundances at three months of age were associated with presence of atopic dermatitis (AD) on examination at three and twelve months of age. From the selected cases and controls with longitudinal samples (n=70), transition to *Bacteroides*-rich communities and influx of adult-specific microbes were observed during the first year of life. The introduction of allergenic solids promoted a significant increase in Shannon diversity and increased abundances of specific microbes, such as genera belonging to *Prevotellaceae* and Proteobacteria (e.g. *Escherichia/Shigella*), as compared to infants recommended to exclusively breastfeed.

**Conclusions:** Specific gut microbiota of three-month-old breastfed infants were associated with cesarean birth, while increased *Clostridium sensu stricto* was associated with AD. The randomized introduction of allergenic solids from three months of age alongside breastfeeding was associated with differential dynamics of maturation of the gut microbial communities.

**Key messages:**

* Cesarean delivery is associated with reduced gut microbiota diversity at three months.
* The infant gut microbiomes mature towards *Bacteroides*-rich communities that begin to transition during the latter half of the first year of life.
* Early introduction of solids into babies’ diets accelerates microbiota diversity and promotes different trajectory of maturation.
* *Clostridium sensu strictu* abundance at three months was associated with AD and AD severity.

**Capsule summary:**

Among exclusively breastfed infants, the introduction of allergenic solids alongside breastfeeding was associated with a differential maturation of the gut microbiota.

**Keywords:** atopic dermatitis, bacteria, diet, environment, food, microbiome, colonization, tolerance

**Abbreviations:**

AD Atopic dermatitis

CI Confidence Interval

cOR Crude Odds Ratio

EAT Study Enquiring About Tolerance Study

OTU Operational Taxonomic Unit

rRNA Ribosomal ribonucleic acid

TEWL Transepidermal water loss

**Introduction**

Patterns of gut microbiota acquisition during infancy, especially during the first year, are highly dynamic and associated with the development of various conditions in later life, such as atopic diseases (1-4). Mechanisms for how early colonization of differential microbes influences health and diseases are under active research, and evidence has suggested that colonization of microbes is indispensable for immune maturation and organ development. Major events – mode of delivery, feeding patterns, and antibiotic usage – have been shown to affect the early microbiota acquisition (5-11). Although many studies have been conducted to explore the effects of environmental factors on gut microbiome development, most of them are retrospective and observational, therefore, challenged by mixed confounders.

Mode of delivery has been identified as the prominent factor during the newborn period, as it determines colonizing microorganisms that are shared from mother to newborn and may also influence gut microbiota characteristics (5-7, 12). It has been commonly reported that cesarean delivered infants harbour less Bacteroides and more hospital-associated pathobionts, such as *Enterococcus* spp. and *Klebsiella* spp. An infant’s diet may also influence its gut microbiota. Several reports have highlighted gut microbiota differences between breastfed and formula-fed infants (8, 13, 14). Two small uncontrolled studies have described an increase in infants’ gut microbiota complexity when dietary solids are introduced (15, 16), and a non-randomized study of 98 maternal-infant dyads observed that cessation of breastfeeding was associated with maturation into an adult-like microbiota (8).

Here, we investigated a nested cohort of infants undergoing randomized introduction of allergenic solids as part of a randomized controlled trial to prevent food allergy. This study provided an opportunity to evaluate the effects of early introduction of solid foods and other factors including mode of delivery and environmental exposures on the shifts and maturation of the gut microbiota and the risk of developing atopic disorders (17, 18).

**Methods**

*EAT cohort*

In the EAT study, 1303 exclusively breastfed, healthy infants from England and Wales, aged between 12 and 17 weeks, were enrolled into a randomized controlled trial to examine whether the regular consumption of allergenic solids reduced the prevalence of food allergy when compared to continued exclusive breastfeeding up to six months of age (ISRCTN: 14254740) (17). Participants were randomly assigned to either the regular consumption of six allergenic foods (boiled hen’s egg, peanut, cow’s milk (yoghurt), wheat, white fish and sesame) beginning at three months of age twice weekly alongside continued breastfeeding (early introduction group) or exclusive breastfeeding until six months (standard introduction group). Beyond six months, food consumption was at parental discretion. Food specific IgE to each of the six foods was measured at enrollment, one and three years of age in both groups using ImmunoCap (Phadia) assays (cut off ≥0.35kU/L to determine food sensitization). The primary outcome of the original EAT Study was the prevalence of challenge-proven food allergy to one or more of the six study foods between one and three years of age. Parents completed online questionnaires eliciting exposure data, such as mode of delivery, sibship size, pet ownership, and antibiotic usage.

All infants were examined for atopic dermatitis (AD) at their enrolment visit at three months of age and twelve months, using the UK diagnostic criteria–based photographic protocol of the International Study of Asthma and Allergies in Childhood (ISAAC) Phase Two (19). AD severity was determined by the Scoring Atopic Dermatitis (SCORAD) index (20). Skin barrier function was assessed by measuring trans-epidermal water loss (TEWL) using the Biox Aquaflux AF200 (Biox, London, UK) closed condenser chamber device on unaffected skin of the left volar forearm. Venous blood samples were screened for the 6 commonest *FLG* mutations (TaqMan allelic discrimination assays for mutations R501X, 2282del4, R2447X, S3247X; ABI 7900 HT; Applied Biosystems, Foster City, Calif) or by sizing of fluorescent PCR products on an Applied Biosystems 3130 DNA sequencer (mutations 3673delC and 3702delG). These 6 mutations detect 99% of *FLG* mutation carriers in the UK population.

Ethical approval for the EAT Study was provided by the St Thomas’ Hospital Research Ethics Committee (REC Reference 08/H0802/93) and informed consent was obtained from the parents of all children enrolled in the study.

*Gut microbiota sample collection, DNA extraction and sequencing*

Beginning the 1st of October 2011, consecutively enrolling families (n=359) were asked to provide stool samples (see Fig E1). Stool samples were collected from 288 of enrolled participants with follow up samples requested at six and twelve months of age. Stool samples were retrieved from diapers and transferred to the King’s College London Molecular Microbiology Research Laboratory for storage at -80°C until DNA extraction. The cell lysis and DNA extraction protocol utilized both phenol-chloroform liquid-liquid phase separation and bead-beating, followed by -80°C storage (see Supplementary Appendix). To evaluate the assembly of gut microbial communities during infancy, a total of 428 samples were sequenced (of the total 288 subjects providing baseline samples, 218 participants provided baseline samples only, and a subset of 70 individuals provided samples from multiple time points of three, six, and twelve months). The subset of cases and controls (42 in the standard-introduction group and 28 in the early-introduction group) were selected based on intervention assignment and atopy (either challenge-proven food allergy or AD on examination) status. DNA extracts were amplified using primers targeting the hypervariable V4 (515F-806R) region of the *16S* ribosomal RNA gene and performed on the MiSeq instrument (21).

*Sequence analysis*

The open-source software mothur (22) and DADA2 (23) were used to process the raw sequencing data (see online supplement). Reads were aligned and classified down to the genus level with ribosomal database project naïve Bayesian classifier (24) or species level with Greengenes database v13.8 (25). Operational taxonomic units (OTUs) were defined at 97% similarity using the opticlust algorithm (26). We also undertook comparative analyses with 977 samples from the TwinsUK (adults from the same UK region) (27, 28), which were sequenced using the same primers (Accession No. ERP006339 and ERP006342; 9 samples were removed during preprocessing) and TEDDY (large, multi-centre infant cohort) (11) studies. TEDDY dataset samples were selected based on i) exclusive breastfeeding before solid food introduction and ii) longitudinal data (2 samples before 6 months of age) to permit comparable analyses.

*Statistical analysis*

The beta-diversity and linear discriminant statistical analyses were performed using R software, with libraries including vegan, ape, and bios2mds. Thetayc distance (theta) metric was used to measure distance between communities (beta diversity) (29), and principal coordinate analysis (PCoA) was performed based on theta distance, unless otherwise indicated. Significance of differences between groups was tested by various methods, including Wilcoxon rank-sum, Kruskal-wallis, and the analysis of molecular variance (AMOVA) (30). Covariates of community variation were evaluated by calculating correlation between indicated PCoA ordination and metadata or genus abundance (envfit function in vegan R package; 10,000 permutations with Bonferroni correction).

**Results**

Of enrolled families, 80.2% (288 of 359) contributed baseline samples, a subset of which (n=70) also contributed follow-up samples. Within the subset, cases were comprised of twelve participants with allergies to egg (one additionally allergic to peanut), one milk allergy, and one codfish allergy, and thirteen with AD (SCORAD over 15 at either three or twelve months).

*Exclusively breastfed infants’ gut microbial communities form 3 distinct clusters*

To understand the landscape of the gut microbiome during breastfeeding, we analyzed baseline (3-month) samples (n=288, and a mean age of 105 days and interquartile range 100-112 days). The most abundant genera established in the microbial communities at three months were *Bifidobacterium* (median abundance (MA): 38.75%), *Bacteroides* (MA: 5.43%) and multiple genera belonging to Firmicutes and Proteobacteria (Fig 1, A and B). The gut microbial communities of exclusively breastfed three-month-old infants was diverse, in examining the PCoA with theta distance (29) and Bray-Curtis distance based on genus-level classification (Fig 1, A and see Fig E2, A). k-means clustering (see Fig E2, A and B) revealed that the microbial communities of exclusively breastfed infants clustered into 3 groups, in which cluster with Bifidobacterium-rich cluster #1, Bacteroides-rich cluster #2 and cluster #3 predominated by *Escherichia*/*Shigella* and genera belonging to Firmicutes phylum (Fig 1, A and B). Although minor genera, such as *Streptococcus*, *Dorea* and *Rothia*, also contributed to the community variation, the three major genera were sufficient to reproduce clustering, underscoring their importance in community variation (Fig 1, C). Community diversity was lower in cluster #1 than in clusters #2 and #3 (alpha diversity; Shannon index, p<0.005; Wilcoxon-rank sum test) due to the predominance of *Bifidobacterium* (Fig 1, D). Of note, the age of participants did not have a significant effect on clustering (see Fig E2, C, p>0.05). PCoA and k-means clustering based on theta distance calculated from species-level (see Fig E2, D) and *de novo* OTU (97% similarity) abundance (see Fig E2, E) and on Bray-Curtis distance (see Fig E2, A) were highly comparable to the genus-based clustering (Fig 1, B).

*Infants’ gut microbial communities mature in first year*

Data from six- and twelve-month samples (n=70 samples per each time point; see methods) were superimposed onto the existing three-month PCoA coordinates (see Fig E3, A). Microbiota from six-month-old infants were similarly dispersed as three-month communities, whereas twelve-month samples were exclusively overlaid on cluster #2, indicating that gut microbial communities mature towards *Bacteroides*-rich communities at 1 year of age, in keeping with the previous literature (7, 10). Interpersonal similarity increased during aging (see Fig E3, B) due to the convergence to Bacteroides-rich communities. We then investigated whether the infant gut microbiomes matured towards an adult-like pattern. Since gut microbiomes can vary in adult populations from distinct geographic regions (31), we compared our dataset to a representative dataset of fecal microbiota from adults in a geographic region analogous to our infant cohort, revealing that infants begin to transition towards adult gut microbiomes during six to twelve months of age (see Fig E4). Collectively, these results demonstrated the extensive transition and maturation of gut microbiota during the first year of life.

*Early environmental exposures are associated with the infant gut microbiota*

Among the environmental exposures that were evaluated, mode of delivery was the most significant factor contributing to microbiome community variation (p<0.01) at 3 months of age (Fig 2, A), with underrepresentation of *Bacteroides* in individuals born by cesarean section (Fig 2, B and C), as shown in previous reports (5, 32, 33). Since the mode of delivery profoundly contributed to microbiome variation, PCoA and clustering were re-analyzed after stratification by delivery methods. Gut microbial communities of vaginally delivered infants also formed 3 clusters, with the same three key genera – *Bifidobacterium*, *Bacteriodes*, and *Escherichia/Shigella* – as the major contributing taxa (see Fig E5, A), indicating contribution of other exposures in defining microbiome communities. Therefore, pair-wise associations between major bacterial genera (mean abundance >1%) and environmental factors were investigated (see Fig E5, B). Results highlighted additional bacterial genera associated with delivery mode (e.g. Bifidobacterium, *Clostridium sensu stricto,* Enterobacteriaceae, and Lachnospiraceae were enriched in infants with cesarean-delivered infants) underscoring the importance of delivery mode in shaping infant gut microbiomes.

*Clinical outcomes are linked with specific infant gut microbiota*

We next examined potential links between clinical outcomes and the 3-month gut microbiome clusters; however, no statistically significant associations were identified. Therefore, associations between major bacterial genera (mean abundance >1%) and clinical manifestations were also investigated (Fig, E6, A). Interestingly, SCORAD at 12 months of age was significantly associated with the abundance of *Clostridium sensu stricto* and *Haemophilus* at enrolment. Equally, raised TEWL at 3 months showed a positive association with *Haemophilus*. Raised TEWL at 12 months was also linked to an increased abundance in *Veillonella* (Fig E6, A). Linear discriminant analysis confirmed higher relative abundances of *Clostridium sensu stricto* at three months of age were observed in infants who had AD at three months and twelve months of age (see Fig E6, B and C). Family history of allergic disease and food allergy/sensitization did not significantly differ according to the microbiome variation or clustering.

*The introduction of allergenic foods alters infants’ gut microbiota*

The EAT Study randomized exclusively breastfed infants to dietary introduction of six allergenic foods (‘early introduction’, n=28) versus continued exclusive breastfeeding (‘standard introduction’, n=42) until 6 months of age (17). Therefore, the maturation in the gut microbiota was investigated relative to the timing of introduction of allergenic solids. Participants in the intervention group had a significantly increased Shannon microbial diversity index at six months as compared to their enrolment samples (Fig 3, A, and Fig E7, A). PCoA revealed the gut microbial communities of the early and standard introduction groups were largely indistinguishable (Fig 3, B, P non-significant; AMOVA) at three and twelve months. In contrast, these microbial communities significantly differed at six months of age (Fig 3, B, middle panel. p<0.05; AMOVA), demonstrating the differential maturation trajectories of early versus standard introduction groups at six months of age. Interestingly, early intervention group infants predominantly transitioned along the second principal coordinate (y-axis) from three to six months of age (Fig 3, C). Abundances of *Prevotella* and *Escherichia/Shigella* were positively correlated with the second principal coordinate (p<0.05, and r2=0.52 and 0.44, respectively). Similarly, linear discriminant analysis revealed *Paraprevotella* and various genera belonging to Proteobacteria and Firmicutes phyla were enriched in the early intervention group (see Fig E7, B and D) at six months, but not evident at 12 months of age (see Fig E7, C and E). Infants in both groups moved along with the first principal coordinate (x-axis) and converged towards *Bacteroides-*rich communities during six to twelve months (Fig 3, D). To validate that the introduction of solids altered the gut microbiota in exclusively breastfed infants, we analyzed gut microbiome changes in exclusively breastfed infants after similar dietary changes within the TEDDY cohort (see Methods for detailed criteria) (11) and superimposed their data onto the same PCoA dimensions of (see E8, A). TEDDY study infants with early introduction of solids (n=48, no formula and started solid foods before 150 days of life) also transitioned along with the second principal coordinate from 3 months to 6 months, whereas TEDDY study infants mimicking the standard introduction group (n=45, no formula and started solid foods after 150 days of life) did not follow the same gut microbiome transition (Fig 3, E).

**Discussion**

There remains deep interest in understanding what contribution external factors may contribute to the development of atopic and allergic disorders. This cohort provided a unique opportunity to examine the relationship between the gut microbiome and many factors including dietary intervention, clinical outcomes, and hygiene-related factors. Gut microbiota are widely heterogenous amongst exclusively breastfed three-month-old infants and have lesser abundance of *Bacteroides* species if born by cesarean section. After three months of age, the gut microbiota underwent major transformation in diversity, complexity, and predominance of ‘adult-like’ bacteria. Babies’ gut microbiota diversified when allergenic foods were introduced and matured towards *Bacteroides*-rich communities at twelve months of age, with significant changes observed at a younger age in infants with earlier introduction of allergenic solids beginning at three months of age. While there were significant microbiota-related associations with AD and AD severity at three and 12 months, none of the observed gut microbiota changes were associated with the development of food allergy or sensitization.

*Study strengths and limitations*

Strengths of the EAT study cohort were strictly controlled eligibility of exclusively breastfed infants, population-representative sampling from the general population across England and Wales, and a randomized intervention with allergenic solid introduction between three- and six-months age, rather than an observational study design with no control of food intake (17). The nested observational study within the EAT cohort allowed comparisons of the effects of the dietary intervention on the gut microbiota over the first year of life. We validated these findings in exclusively breastfed infants with comparable introductions of solid foods in the TEDDY cohort, recapitulating these observations in other populations with westernized diets. Detailed phenotyping of this cohort enabled assessment of alterations in infant gut microbiota in association with exposures, such as mode of delivery, antibiotic prescribing, communal child care, sibship size, rural vs urban residence, and pet ownership. Validated scores were used to examine participants for AD, and diagnosis of food allergy was challenge-based, rather than relying on parental report. A limitation of our study is that we were unable to evaluate the gut microbiota before three months of life, which would have allowed an even earlier characterization of microbiota development.

*Neonatal environmental exposures may influence gut microbiota at three months of age*

Delivery by cesarean had the most profound effects on the shaping of the gut microbiota during early infancy with reduction in *Bacteroides,* similar to other cohorts (5, 11, 32, 33). Colonization with *Bacteroides* in the neonatal period may have considerable health benefits, especially immune maturation (34). *Bacteroides fragilis* has Polysaccharide A molecular moieties which stimulate the development of T-regulatory cells in the lamina propria (35) and samples from human fecal microbiota have found that at least 16 Bacteroides taxa contribute to this phenomenon *in vivo* (36). Therefore, the abundance of *Bacteroides* spp. may relate to the nature of early life immunological priming and warrants further investigation. In addition, if these mechanistic studies are fruitful, intervention trials may consider targeting infants born by cesarean to try and alter *Bacteroides* relative abundances and/or *Bacteroides* diversity (7).

Although we did not find other exposures contributing to the heterogeneity of microbiome at three months of infancy, there may be mixed effects of multiple factors and/or hidden factors which were not included in our metadata; therefore, an optimized model for discovering associated factors might be necessary. With the current dataset, we trained a random forest classifier with metadata to predict 3 clusters at three months. While the classifier did not perform well (out-of-box error rate > 40%, data not shown), it calculated maternal age at birth as a second important factor for classifying clusters, suggesting the importance of maternal/household factors in microbiome development at infancy. Similarly, studies have reported the long-term colonization of vertically transmitted microbes (12).

*Diet contributes to evolution of the infant gut microbiota*

Infants’ diet also had significant effects on the shaping of the gut microbiota during early infancy. Several reports have demonstrated differential microbiome colonization in breastfed versus formula-fed infants, with a higher prevalence of *Bifidobacterium* spp. (*B. longum* and *B. breve*) and *Lactobacillus* in breastfed infants (8, 9, 11, 14). In addition, the introduction of solid foods has been proposed as a key contributor to the maturation of the gut microbiota. For example, a Scandinavian observational study concluded that the gut microbiota matured after the cessation of breastfeeding (8).

Since studies randomizing dietary interventions in infants are very unusual, this study offered a unique opportunity to study the effects of introducing allergenic solids on the gut microbiome. In the EAT study, early peanut and egg introduction, if consumed in sufficient quantity, was shown to protect against the development of peanut and egg allergies between one and three years of age (17, 18). We have demonstrated that the early introduction of allergenic foods alongside ongoing breastfeeding between three and six months of age led to an increase in overall gut microbiota Shannon diversity, in particular promoting an influx of various microbes including *Prevotellaceae* and *Escherichia/Shigella*. Interestingly, presence of *Prevotella* has been shown to be associated with high-fiber diet including remote villages with less frequent chronic inflammatory disorders (37-40). Additionally, preceding studies have noted that the cessation of breastfeeding may promote further diversification of infants’ gut microbiota(8), whereas our results showed that the introduction of a variety of allergenic foods in the setting of exclusive breastfeeding increased diversity of the gut microbiota. Of note, six-month gut microbiota Shannon diversities were not statistically significantly different between the early and standard introduction groups, indicating that the increased microbial community diversity after early introduction of allergenic foods was comparable to exclusively breast-fed infants (Fig E7, A). However, the combined findings from this study demonstrate that early introduction of solid foods elicits considerable and consistent changes in gut microbial composition and diversity from three to six months of infancy. Therefore, controlled dietary interventions in infancy may promote selective colonization of desired microbes and potentially exerts health and developmental benefit to the host. However, we found no evidence that the observed *changes* in the gut microbiota driven by the early introduction of allergenic foods impact on the development of AD.

In conclusion, we confirmed that cesarean delivery is associated with reduced gut microbiota diversity at three months. The early introduction of solids into infants’ diets accelerates maturation of microbiota diversity as well as increases the relative abundances of *Prevotella* and *Escherichia/Shigella*. This manipulation of the gut microbiota in early life towards diversity and maturity, in particular in those delivered by cesarean section or at risk of allergy and atopic diseases, should be examined in future studies to examine potential wider health benefits.

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***Data Access***

All sequencing data were deposited and are available at the NCBI Sequence Read Archive (SRA) under BioProject number PRJNA597342.

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**List of figure captions**

**FIG 1. Gut microbiota characteristics at three months of age**

A, Principal coordinate analysis (PCoA) of the gut microbiome at 3 months. Pair-wise distances (theta distance) among all samples were calculated and two major axes (PC1 and PC2) from the multi-dimensional distance space were calculated and depicted on a scatter plot. Colors indicate different clusters, according to k-means clustering. Arrows and letters indicate specific genera significantly correlated with PCoA ordination (p<0.05, lengths of arrows are proportion to R2, calculated from EnvFit function from R). Corresponding genera for letters are showing on the right panel of B.

B, Averaged stacked bar chart of infants’ microbiome within each cluster. Genera that are significantly correlated with PCoA ordination are depicted as colored bar (legend on right) and others are merged as ‘Other bacteria’.

C, Triangle plot showing relative abundances of the three key genera (*Bifidobacterium*, *Bacteroides*, and *Escherichia*/*Shigella*) in the gut microbiota of baseline samples. The colors indicate clusters (same as Fig 1A).

D, Boxplot of bacterial community diversity (Shannon index) differences according to microbiome community clusters. Shannon diversity of Cluster #1 community is significantly lower than Cluster 2 and 3. (\*\*p<0.01, \*\*\*p<0.001; Wilcoxon rank-sum test, after Kruskal-wallis test)

**FIG 2. Evaluation of associations between gut microbiota characteristics and cesarean delivery**

A, Association analysis of clinical, hygiene and life-style factors (‘covariates’) with microbiome variations. Among total of 48 covariates tested, 11 covariates with r2>0.01 are plotted here. Each bar indicates the amount of variance explained by each covariate, calculated by EnvFit function from R. (\*\*p<0.01; Bonferroni corrected) After correction for multiple comparison, mode of delivery remained as a significant factor that associated to microbiome variation.

B, PCoA of microbiomes at 3 months with delivery mode delineated. Color indicates cluster (same as Fig 1, A) and the shape indicates delivery mode. Arrows demonstrate direction of each covariate in the ordination space. Infants born by cesarean section tend to cluster on the left portion of scatter plot.

C, Boxplot showing the difference of relative abundances of *Bacteroides* by mode of delivery. *Bacteriodes* are largely missing in infants born by cesarean section. (\*\*\*p<0.001; Wilcoxon rank-sum test)

**FIG 3. Impact of allergenic solid introduction on infants’ gut microbiota**

A, Boxplot comparing Shannon diversity changes amongst participants’ longitudinal samples according to randomized allocation to continued exclusive breastfeeding (standard introduction group) or the introduction of allergenic solids (early introduction group) (\*p<0.05; Wilcoxon rank-sum test).

B, PCoA scatter plot demonstrating longitudinal transition from 3 to 6 month of age. Grey arrows connect samples from the same individuals. Yellow and purple arrows on the sides indicate the average shift of the microbiota in each PCoA axis.

C, PCoA scatter plot demonstrating longitudinal transition from 6 to 12 month of age. Grey arrows connect samples from the same individual. Black arrows on the sides indicate average shift of the microbiota in each PCoA axis.

D, PCoA plot showing longitudinal transition of the gut microbiome from 3 to 12 months of age in the different dietary intervention groups. At 6 months, the microbiota of early introduction is significantly different from standard introduction (p<0.05 by AMOVA).

E, Boxplot showing changes of microbiome from 3 to 6 months, in different dietary intervention groups and cohorts (EAT and TEDDY). Early introduction in both studies led to an increase along the second principal coordinate axis (\*p<0.05 and \*\*p<0.01; paired Wilcoxon rank-sum test).