Infant Gut Microbiota Development Is Driven by Transition to Family Foods Independent of Maternal Obesity

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ABSTRACT The first years of life are paramount in establishing our endogenous gut microbiota, which is strongly affected by diet and has repeatedly been linked with obesity. However, very few studies have addressed the influence of maternal obesity on infant gut microbiota, which may occur either through vertically transmitted microbes or through the dietary habits of the family. Additionally, very little is known about the effect of diet during the complementary feeding period, which is potentially important for gut microbiota development. Here, the gut microbiotas of two different cohorts of infants, born either of a random sample of healthy mothers (n = 114), or of obese mothers (n = 113), were profiled by 16S rRNA amplicon sequencing. Gut microbiota data were compared to breastfeeding patterns and detailed individual dietary recordings to assess effects of the complementary diet. We found that maternal obesity did not influence microbial diversity or specific taxon abundances during the complementary feeding period. Across cohorts, breastfeeding duration and composition of the complementary diet were found to be the major determinants of gut microbiota development. In both cohorts, gut microbial composition and alpha diversity were thus strongly affected by introduction of family foods with high protein and fiber contents. Specifically, intake of meats, cheeses, and Danish rye bread, rich in protein and fiber, were associated with increased alpha diversity. Our results reveal that the transition from early infant feeding to family foods is a major determinant for gut microbiota development.

IMPORTANCE The potential influence of maternal obesity on infant gut microbiota may occur either through vertically transmitted microbes or through the dietary habits of the family. Recent studies have suggested that the heritability of obesity may partly be caused by the transmission of “obesogenic” gut microbes. However, the findings presented here suggest that maternal obesity per se does not affect the overall composition of the gut microbiota and its development after introduction of complementary foods. Rather, progression in complementary feeding is found to be the major determinant for gut microbiota establishment. Expanding our understanding of the influence of complementary diet on the development and establishment of the gut microbiota will provide us with the knowledge to tailor a beneficial progression of our intestinal microbial community.

KEYWORDS: 16S rRNA sequencing, breastfeeding, complementary diet, family foods, infant gut microbiota, maternal obesity

Despite the temporal resilience and stability of the gut microbiota, long-term diet (1) and major diet shifts (2) are known to affect the human gut microbiota. Infancy and early childhood constitute a period in life in which the microbiota is characterized by relatively low stability and high responsiveness toward influencing factors. During this period, dietary factors have major implications for the establishment of the gut microbiota.
microbiota (3, 4). While many previous studies have focused on early infant diet (5, 6), particularly breastfeeding and formula feeding, only a few have addressed the effects of the complementary (solid-food) diet of infants in the period after 6 months of age (7). As the gut microbial population is not fully established until the age of 3 to 5 years (8, 9), it is important to understand how it is influenced by the transition from early infant feeding to family foods during the complementary feeding period, which is defined by the WHO as the period from 6 until 18 to 24 months of age (10). The established adult gut microbiota has been linked with a range of metabolic, autoimmune, and allergic diseases (9). Specifically, the intestinal microbiome has repeatedly been linked to obesity in animal models (11–15) as well as in human studies (16, 17). By transplantation of fecal microbial communities from human twins discordant for obesity into germfree mice, it has been shown that a greater increase in body mass and adiposity occurs in mice transplanted with obese donor microbiota than with the corresponding microbiota from the lean donor twin, suggesting the importance of gut microbes over human genetics in the etiology of obesity (18). Indeed, children of obese parents have a higher risk of developing obesity, and this is not explained solely by human genetic predisposition. Since gut microbes can be transferred from mother to infant during birth (19), an obesity-associated microbiota may be transferred from an obese pregnant woman to her offspring. It is well documented that obese mothers on average breastfeed for a shorter time than normal-weight mothers (20) and that breastfeeding has a protective effect on obesity in offspring (21). Further, parental obesity is associated with lower socio-economic status and specific dietary patterns (22) that may affect the type of complementary diet introduced to the infant and thereby the development of the gut microbiota (9), as well as contribute to future obesity risk (23). Indeed, diet-microbiota interactions have been shown to be key players in the development of obesity (18). Therefore, we compared the gut microbiota profiles of two different cohorts of Danish infants at the ages of 9 and 18 months, designated SKOT I (24) and SKOT II (25), respectively (SKOT is a Danish abbreviation for dietary habits and well-being of young children). SKOT I includes infants from a random sample of mothers (mean body mass index [BMI], 22.9 kg/m²), and SKOT II includes infants of obese mothers (mean BMI, 35.1 kg/m²). To elucidate the impact of (i) maternal obesity and (ii) dietary factors on infant gut microbiota development, associations between specific features of the gut microbiota and dietary factors were investigated with a focus on breastfeeding and complementary diet composition.

RESULTS
Gut microbiota development during the complementary feeding period is independent of maternal obesity. To assess the impact of maternal obesity on gut microbiota establishment in offspring, we sequenced the V3 region of 16S rRNA genes from fecal samples of 227 individuals at both 9 and 18 months of age in the two SKOT cohorts. These cohorts are different with respect to maternal obesity and generally differ in terms of socio-economic status, C-section prevalence, and early infant feeding but differ only slightly with respect to infant body composition measures (Table 1). Between-sample diversity (beta diversity) of the gut microbiota in the two cohorts was investigated by principal-coordinate analysis (PCoA) of the Bray-Curtis dissimilarity indices and showed clustering according to age rather than cohort (Fig. 1A). Distances to the group centroid for each point, as an estimate of beta diversity, illustrated no differences between cohorts (Fig. 1A). However, greater beta diversity was observed at 9 months than at 18 months in both cohorts, in line with previous reports (26–28). Levels of within-sample diversity (alpha diversity), as estimated by the Shannon index, the number of observed genera, and Pielou’s evenness index of the communities, were not significantly different between the two cohorts at either 9 months or 18 months of age (Fig. 1B). However, there was a significant increase in these alpha diversity measures from 9 to 18 months in both cohorts. On a compositional level, the gut microbiotas across time and cohorts were dominated by four phyla, Firmicutes (64.2%), Actinobacteria (23.4%), Bacteroidetes (7.7%), and Proteobacteria (4.3%), while less than
0.5% belonged to other phyla or were unclassified. On average, 98.3% of the communities belonged to 24 bacterial families (Fig. 2). Despite large interindividual variation, average bacterial communities assessed at the phylum level as well as at the family level at 9 months and 18 months were highly similar between SKOT I and II (Fig. 2). Indeed, according to PCA of family-level composition, samples clustered according to age rather than cohort and showed the relative contributions of bacterial families to the variation in the data set (Fig. 3A). After correction for multiple testing, we found no significant differences between cohorts with respect to relative abundances of bacterial phyla, families, or genera at either 9 or 18 months, and no differences in the changes occurring from 9 to 18 months between the two cohorts were identified (Fig. 3B). In contrast, over time, Lachnospiraceae, Ruminococcaceae, Eubacteriaceae, Rikenellaceae,
and *Sutterellaceae* were significantly increased in both cohorts, and *Bifidobacteriaceae*, *Actinomycetaceae*, *Veillonellaceae*, *Enterobacteriaceae*, *Lactobacillaceae*, *Enterococcaceae*, *Clostridiales incertae sedis* XI, *Carnobacteriaceae*, and *Fusobacteriaceae* were significantly decreased in both cohorts (Fig. 3B; see Table S1 in the supplemental material). This is in agreement with a previous study of the SKOT I cohort using quantitative-PCR (qPCR)-based microbiota assessment (8) and with studies involving other cohorts (27–30). These results suggest that maternal obesity *per se* does not influence gut microbiota development during the complementary feeding period. The high gut microbiota similarity between the two cohorts, independently sampled during different time periods, allowed a high-powered characterization of infant gut microbiota development.
ota development and identification of the main factors explaining variation in gut microbiota.

Limited influence of C section, gestational age at birth, and prior use of antibiotics. The mode of delivery (31), gestational age at birth (within a normal range of full-term delivery) (32), and use of antibiotics (33) have all previously been shown to impact the infant gut microbiota. In the SKOT cohorts, neither microbial community compositions nor alpha diversity measures at 9 months were significantly different between individuals born by C section and those born vaginally (see Table S2 in the supplemental material). We did, however, note a decreased relative abundance of Bacteroidaceae ($P = 0.003$, false-discovery-rate-corrected $P$ values $[q] = 0.072$) in infants born by C section in SKOT II (Table S2), in line with results of previous studies (27, 31, 34). Gestational age at birth was not associated with gut microbiota composition or alpha diversity at 9 months (Table S3), and the use of antibiotics 2 weeks before the sample was taken (current antibiotic use was an exclusion criterion) could not explain the variation in gut microbial diversity at 9 or 18 months (Table S4). All infants in the present study were delivered at full term (range, 37 to 42 weeks), C-section prevalence was low in SKOT I (Table 1), and for only a few infants was the use of oral antibiotics during the 2 weeks prior to sampling registered (10 individuals in total). Further, the relative late sampling point (9 months of age) may explain discrepancies with prior studies.

FIG 2 Composition of gut microbiota in SKOT I and SKOT II. Relative abundances of bacterial phyla (small panels) and families (large panels) in the SKOT I cohort ($n = 114$) at the ages of 9 months (A) and 18 months (B) and in the SKOT II cohort ($n = 113$) at the ages of 9 months (C) and 18 months (D). Boxes indicate 25th to 75th percentiles, with mean relative abundances marked as lines and whiskers indicating the range (minimum/maximum) multiplied by the interquartile range (25th to 75th percentiles) from the boxes. Bacterial families are ranked by average relative abundances at the age of 9 months. Detailed information can be found in Table S1 in the supplemental material.
Duration of exclusive breastfeeding, rather than age at introduction of complementary feeding, is reflected in late-infancy gut microbiota. Danish mothers are advised to exclusively breastfeed their infants until the age of approximately 6 months and to continue partial breastfeeding until the infant is about 1 year old. It is additionally recommended to introduce complementary foods (apart from infant formula) at about the age of 6 months but not before the age of 4 months (35). As we have previously reported (25), infants in the SKOT I cohort were both exclusively and partially breastfed significantly longer than infants in the SKOT II cohort. Additionally, age at the introduction of complementary foods was significantly lower in SKOT II than in SKOT I (Table 1). Despite the fact that no infants in either of the cohorts were exclusively breastfed beyond the age of 6 months, the recorded duration of exclusive breastfeeding was associated with the relative abundance of specific bacterial taxa at the age of 9 months. This was most pronounced in SKOT I, possibly due to the longer average duration of exclusive breastfeeding in this cohort (Fig. 4A and B). However, differences in the effects of duration of exclusive breastfeeding on microbiota between cohorts were modest and not large enough to evoke detectable significant differences between the two cohorts at the age of 9 months (Fig. 1 to 3). In both cohorts, the duration of exclusive breastfeeding was negatively correlated with Lachnospiraceae (e.g., the genera Dorea, Coprococcus, Blautia, Pseudobutyrivibrio, and Roseburia) and genera within Ruminococcaceae (e.g., Ruminococcus, Anaerotruncus, Oscillibacter, Clostridium IV, and Butyricicoccus), encompassing species known to utilize plant-derived complex carbohydrates and resistant starch introduced with solid foods (36). Also, Erysipelotrichaceae, Peptostreptococcaceae, and Eubacteriaceae were negatively affected...
by the duration of exclusive breastfeeding (Fig. 4A). Positive correlations with exclusive breastfeeding were observed in both cohorts for *Bifidobacteriaceae* (*Bifidobacterium*), which are known to utilize the lactose and human milk oligosaccharides found in breast milk (37), and *Veillonellaceae* (e.g., *Veillonella* and *Megasphaera*), known lactate utilizers (38, 39). In addition, *Pasteurellaceae* (*Haemophilus*) abundances were positively correlated with the duration of exclusive breastfeeding (Fig. 4A and B). Although not significant in both cohorts, lactic acid bacteria (*Lactobacillaceae*, *Enterococcaceae*, *Streptococcaceae*) and other bacteria known to be present in human milk, like *Prevotella* (40), and on breast tissue, like *Enterobacteriaceae* (*Escherichia* and *Klebsiella*) (41), were positively correlated with duration of exclusive breastfeeding (Fig. 4A and B). At the age of 9 months, 97 infants ($n_{SKOT\ I} = 59$, $n_{SKOT\ II} = 38$) were still partially breastfed. Additionally, the estimated average daily breast milk intake at the age of 9 months was strongly correlated with gut microbiota composition and confirmed the associations obtained for the duration of exclusive breastfeeding (see Table S5 in the supplemental material). Consistently with our previous report (8), the effects of breastfeeding on microbial composition were limited at 18 months (Table S6). Some infants are fed with infant formula as a replacement or a supplement to breastfeeding for a period prior to the introduction of complementary foods. However, age at the introduction of complementary foods (range, 3 to 6 months) did not correlate with abundances of specific bacterial families at 9 months (Table S7). Furthermore, alpha diversity measures at 9 months were negatively correlated with the duration of exclusive breastfeeding, whereas age of introduction to complementary feeding was generally not correlated with alpha diversity measures, although a weak negative association with observed

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**FIG 4** Duration of exclusive breastfeeding is reflected in late-infancy gut microbiota. Hierarchical clustering of Spearman’s rank correlations of duration of exclusive breastfeeding with gut microbial composition at 9 months of age at the family (A) and genus (B) levels in SKOT I and II. *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$. (C) Spearman’s rank correlations of alpha diversity measures (Shannon index, observed genera, and Pielou’s evenness index) to duration of exclusive breastfeeding (0 to 6 months) and age at introduction of complementary feeding (3 to 6 months) for compiled data from SKOT I and II. Boxes indicate 25th to 75th percentiles, with mean values marked as a line and whiskers indicating minimum and maximum values.
Complementary diet affects late-infancy gut microbiota composition. (A) Macronutrient intake in SKOT I and II is expressed as a mean percentage of total energy intake (E%) from carbohydrates, protein, and fat, whereas fiber intake is expressed in milligrams per kilojoule, with error bars indicating standard deviations. (B and C) PCA plot of consumption of the 23 food groups (grams per day per kilogram of body weight) for each individual in SKOT I and II (n = 217), resulting in principal component 1 (PC1), designated family food, and PC2, designated health-conscious food. (D) Heatmap illustrating hierarchical clustering of Spearman’s rank correlations between relative abundances of family-level gut microbes and macronutrient intake. *, P < 0.05; ***, P < 0.001; ***, P < 0.001. (E and F) Spearman’s rank correlations between family food and relative abundances of Bifidobacteriaceae and Lachnospiraceae. 

genera at 9 months was observed (Fig. 4C). These results suggest that breastfeeding duration, rather than the timing of introduction of complementary foods, is reflected in gut microbiota composition during late infancy, as recently proposed in a study of gut microbiome data from Swedish infants (27).

Composition of complementary diet during late infancy affects gut microbiota composition. A validated 7-day food registration was performed by the parents when the infants were 9 months old (42). On the macronutrient level, no significant differences in fat or carbohydrate intake were observed between the two cohorts; however, protein intake was significantly higher (P < 0.0001, Student’s t test) in SKOT II, while SKOT I infants had a significantly higher (P = 0.016, Student’s t test) fiber intake (Fig. 5A). To capture the complete picture of the complementary diet of the infants in both cohorts at 9 months of age, we previously (25) divided the complete dietary recordings into 23 food groups (defined in Table 2). By PCA of the compiled SKOT I and II subsets of data included in this study (n = 217), the previously defined (25) principal components named family foods (PC1) and health-conscious food (PC2) were generated (Fig. 5B). The family foods component describes the transition from early infant foods (with low loadings of breast milk, formula, and porridge) to foods introduced during late infancy (with high loadings of meat, milk, cheese, animal fat, and rye bread).

FIG 5 Complementary diet affects late-infancy gut microbiota composition. (A) Macronutrient intake in SKOT I and II is expressed as a mean percentage of total energy intake (E%) from carbohydrates, protein, and fat, whereas fiber intake is expressed in milligrams per kilojoule, with error bars indicating standard deviations. (B and C) PCA plot of consumption of the 23 food groups (grams per day per kilogram of body weight) for each individual in SKOT I and II (n = 217), resulting in principal component 1 (PC1), designated family food, and PC2, designated health-conscious food. (D) Heatmap illustrating hierarchical clustering of Spearman’s rank correlations between relative abundances of family-level gut microbes and macronutrient intake. *, P < 0.05; ***, P < 0.001; ***, P < 0.001. (E and F) Spearman’s rank correlations between family food and relative abundances of Bifidobacteriaceae and Lachnospiraceae. 

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The health-conscious food component describes the amount of health-conscious food choices with low loadings of sweets/cake, sugary drinks, and fast food and high loadings of fruits, vegetables, fats (vegetable), potatoes, and fish (Fig. 5C). As previously shown (25), there was no difference between the two cohorts with respect to intake of family food \( (P = 0.481, \text{Mann-Whitney test}) \); however, SKOT I had significantly higher scores for health-conscious food \( (P < 0.0001, \text{Student’s t test with Welch’s correction}) \), corresponding to a higher intake of fruits, vegetables, and potatoes (Table 2). Clustering of Spearman’s rank correlations between macronutrient types and gut microbiota compositions revealed that bacterial groups associated with breast milk and early infant feeding, namely, \textit{Bifidobacteriaceae}, \textit{Enterococcaceae}, and \textit{Lactobacillaceae},

**TABLE 2** Food group definitions, average intake, and correlation to gut microbial alpha diversity at 9 months of age in SKOT I and SKOT II

<table>
<thead>
<tr>
<th>Food group</th>
<th>Definition</th>
<th>Avg intake (g/day/kg body wt)</th>
<th>Spearman correlation to Shannon diversity (SKOT I + SKOT II)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SKOT I ((n = 114))</td>
<td>SKOT II ((n = 103))</td>
<td>SKOT I + SKOT II ((n = 217))</td>
</tr>
<tr>
<td>Porridge</td>
<td>Cereal gruel and porridge (homemade or already prepared)</td>
<td>18.21</td>
<td>15.03</td>
</tr>
<tr>
<td>Breakfast cereals</td>
<td>Oatmeal, muesli, Cornflakes, sugar puffs, and sugary cereals</td>
<td>0.13</td>
<td>0.17</td>
</tr>
<tr>
<td>Wheat bread, whole grain</td>
<td>Grainty bread and crisp bread</td>
<td>1.05</td>
<td>1.73</td>
</tr>
<tr>
<td>Wheat bread, no whole grain</td>
<td>White bread and biscuits</td>
<td>0.57</td>
<td>0.89</td>
</tr>
<tr>
<td>Rye bread</td>
<td>Rye bread with and without seeds</td>
<td>1.28</td>
<td>1.40</td>
</tr>
<tr>
<td>Pasta/rice</td>
<td>Pasta and rice</td>
<td>0.90</td>
<td>0.97</td>
</tr>
<tr>
<td>Potato</td>
<td>Potatoes that were boiled, baked, mashed, or prepared in potato salad</td>
<td>4.48</td>
<td>1.51</td>
</tr>
<tr>
<td>Fruit</td>
<td>Fresh fruit, fresh berries, and fruit porridge, soup, or compote (homemade or already prepared)</td>
<td>12.47</td>
<td>8.23</td>
</tr>
<tr>
<td>Vegetable</td>
<td>All vegetables eaten raw, cooked, or mashed alone or in a dish</td>
<td>6.84</td>
<td>3.26</td>
</tr>
<tr>
<td>Fish</td>
<td>All fish and fish products eaten as a sandwich spread or in a dish</td>
<td>0.81</td>
<td>0.71</td>
</tr>
<tr>
<td>Meat</td>
<td>All meat and meat products eaten as a sandwich spread or in a dish, except poultry and fish</td>
<td>1.63</td>
<td>1.74</td>
</tr>
<tr>
<td>Poultry</td>
<td>All poultry and poultry products eaten as a sandwich spread or in a dish</td>
<td>0.43</td>
<td>0.35</td>
</tr>
<tr>
<td>Egg</td>
<td>All egg and egg products eaten as a sandwich spread or in a dish</td>
<td>0.16</td>
<td>0.14</td>
</tr>
<tr>
<td>Fats (animal)</td>
<td>Butter, spreadable butter, and sauce made from butter</td>
<td>0.98</td>
<td>1.01</td>
</tr>
<tr>
<td>Fats (vegetable)</td>
<td>Oil, margarine, mayonnaise, remoulade, ketchup, and low-fat sauce</td>
<td>0.56</td>
<td>0.38</td>
</tr>
<tr>
<td>Cheese</td>
<td>All cheese and cheese products eaten as a sandwich spread or in a dish</td>
<td>0.55</td>
<td>0.65</td>
</tr>
<tr>
<td>Milk</td>
<td>All milk and milk products eaten alone or in a dish, except human milk or infant formula</td>
<td>12.95</td>
<td>17.90</td>
</tr>
<tr>
<td>Formula</td>
<td>Infant formula and follow-up formula</td>
<td>32.71</td>
<td>27.77</td>
</tr>
<tr>
<td>Breast milk</td>
<td>Human milk from the mother</td>
<td>11.62</td>
<td>8.06</td>
</tr>
<tr>
<td>Fruit/nut/snack</td>
<td>Cereal bar, nuts, almonds, dried fruit, fruit spread, jam, honey, peanut butter, and seeds</td>
<td>0.31</td>
<td>0.16</td>
</tr>
<tr>
<td>Sweets/cake</td>
<td>Ice cream, chocolate, licorice, soufflé, croissant, Danish pastry, cookies, cream cake, pancake, and cream puff (mix of light/not light versions)</td>
<td>0.21</td>
<td>0.16</td>
</tr>
<tr>
<td>Sugary drink</td>
<td>Soda, juice, lemonade, chocolate milk, milk shake, and yogurt drink (mix of light/not light versions)</td>
<td>1.12</td>
<td>0.39</td>
</tr>
<tr>
<td>Fast food</td>
<td>Fried potatoes, French fries, hot dogs, pizza, burgers, spring rolls, and chips</td>
<td>0.59</td>
<td>0.50</td>
</tr>
</tbody>
</table>

- The food group definitions are from reference 25. Gut microbial alpha diversity is the Shannon index. Significant \( P \) and q values are highlighted in bold. Data from 10 individuals from SKOT II were missing.
formed a cluster characterized by negative associations with fiber as well as protein intake (Fig. 5D). The families *Erysipelotrichaceae*, *Peptostreptococcaceae*, *Lachnospiraceae*, *Clostridiaceae*, *Sutterellaceae*, and *Ruminococcaceae* formed a cluster positively associated with protein intake, while *Eubacteriaceae*, *Pasteurellaceae*, *Prevotellaceae*, *Veillonellaceae*, and *Fusobacteriaceae* were all positively associated with fiber intake (Fig. 5D). In both cohorts, fiber intake was significantly positively correlated with *Pasteurellaceae* abundance. Additionally, compiling of cohorts revealed that *Pasteurellaceae* correlated positively (*q*/*H* 0.012) with health-conscious foods (Table S5). In both of the independent cohorts, protein intake was significantly positively correlated with *Lachnospiraceae* but significantly negatively correlated with *Bifidobacteriaceae*, probably reflecting the amount of complementary food in the infant’s diet (Fig. 5D). Indeed, the family food dietary pattern, reflecting progression of an infant’s diet toward family foods, was negatively associated with *Bifidobacteriaceae* abundance (Fig. 5E) but positively associated with *Lachnospiraceae* abundance (Fig. 5F). Further, family food correlated negatively (*q* = 0.019) with *Enterococcaceae* abundance and positively correlated (*q* = 0.009) with *Sutterellaceae* abundance (Table S5). These results suggest that the progression from early infant food to family foods with higher protein and fiber contents is the major driver of gut microbial changes during late infancy.

**Progression toward family foods increases the alpha diversity of the infant gut microbiota.** Next, we investigated how the complementary diet affected the gut microbial alpha diversity. At the age of 9 months, protein and fiber intake was significantly positively associated with the Shannon index, while fat intake was negatively correlated with the Shannon index. No significant association was observed between carbohydrate intake and the Shannon index (Fig. 6A to D). Both the dietary pattern family food and to a lesser extent the dietary pattern health-conscious food (E and F) correlated negatively (*r* = 0.322, *p* < 0.0001) with *Family food* and positively correlated (*r* = 0.142, *p* = 0.040) with *Health Conscious food* (PC1) (9.5%, PC2) (13.2%) with macronutrient intake (A to D) and the dietary patterns family food and health-conscious food (E and F) at 9 months of age (SKOT I, green; SKOT II, purple).
These correlations reflect that foods with high loadings in the family food dietary pattern (Fig. 5C), including cheese, meat, and rye bread, were positively associated with alpha diversity, while breast milk consumption at the age of 9 months was negatively correlated with alpha diversity (Table 2). After correction for multiple testing, the remaining 19 food groups did not correlate significantly with alpha diversity (Table 2), indicating that the progression of an infant’s diet toward family foods, characterized by the transition from breastfeeding to more nutritionally diverse family foods rich in fiber and proteins, is the main driver of gut microbial alpha diversity development.

**DISCUSSION**

Elucidating the influence of dietary factors on the development of the gut microbiota during early life is necessary in order to understand potential links between childhood diet and microbiota-associated disease risks later in life. Indeed, the early life period is considered a window of opportunity for microbial manipulation, due to the relatively unstable configuration of the incompletely developed gut microbiota (9). This study reveals that maternal obesity per se does not affect the development of the gut microbiota during the complementary feeding period. Rather, the gradual process leading to the establishment of a complex microbiota, which occurs during infancy and early childhood, is connected primarily to the transition from breastfeeding to family-like foods rich in fiber and protein and apparently independent of other potential effects, such as lifestyle and genetic disposition related to maternal obesity. Until now, only a few studies have investigated potential links between maternal obesity and offspring microbiota (43, 44). Collado et al showed that levels of *Staphylococcus* and *Bacteroides* were significantly higher in infants (aged 1 and 6 months) of overweight mothers (BMI \(> 25\)) than in infants of normal-weight mothers, whereas levels of *Bifidobacterium* spp. were lower (43). Galley et al. found higher alpha diversity and lower beta diversity in children (aged 18 to 27 months) of obese mothers than in children of nonobese mothers but primarily in a subgroup of individuals of higher socio-economic status. These findings were not confirmed by the present study. Previously reported evidence has revealed differences in gut microorganisms between lean and obese individuals (16, 45); however, the infants/children in the present study were not obese at the sampling points. Nonetheless, it is plausible that the relatively late sampling point (9 months) used in this study explains the lack of observed differences in the infants, since potential initial differences in maternally transferred gut microbes may be overruled by the effect of breastfeeding and complementary diet by that age. Regardless of this, we find it remarkable that the communities developed so similarly in the two different cohorts (discordant for maternal obesity), which were independently sampled and originated from infants of different socio-economic backgrounds (Table 1). Despite differences between cohorts in breastfeeding duration and complementary feeding practices (Table 1), these differences did not cause significant differences in gut microorganisms between cohorts within the given sensitivity and taxonomic resolution of our method. Importantly though, the high similarity between cohorts enabled us to cross-validate associations. We observed that *Lachnospiraceae*, *Ruminococcaceae*, *Eubacteriaceae*, *Rikenellaceae*, and *Sutterellaceae* constitute the major bacterial families that increase in abundance from infancy to the toddler age (early childhood) and vice versa for *Bifidobacteriaceae*, *Actinomycetaceae*, *Veillonellaceae*, *Enterobacteriaceae*, *Lactobacillaceae*, *Enterococccaceae*, *Clostridiales incertae sedis XI*, *Carnobacteriaceae*, and *Fusobacteriaceae*, which is largely in agreement with previous studies (27, 29, 30). In contrast to what was anticipated, we found limited effect of C section and gestational age on gut microbiota, but this is probably explained by the fact that our first point of sampling was not until the age of 9 months, at which time such effects are likely to have been diluted by other influencing factors, including breastfeeding and complementary feeding. The relative abundance of gut bacterial taxa as well as alpha diversity measures at the age of 9 months was explained by breastfeeding duration rather than by time of introduction to complementary (solid) foods, in line with what was recently suggested by Bäckhed et al. (27). Specifically, we
found *Bifidobacterium*, *Veillonella*, *Megasphaera*, *Haemophilus*, and members of lactic acid bacteria and the *Enterobacteriaceae* to be positively affected by breastfeeding duration, while the reverse was true for members of *Lachnospiraceae* and *Ruminococcaceae*, known to utilize complex carbohydrates (36). Further, at the age of 9 months, the level of progression toward family foods, reflected in a higher consumption of meat, cheese, and rye bread, rich in fiber and protein, was clearly associated with gut microbiota composition and alpha diversity measures. Particularly, *Lachnospiraceae* abundance increased with the amount of family foods in the diet, while the opposite was found for *Bifidobacteriaceae*. This probably reflects the shift from breastfeeding associated with *Bifidobacteriaceae* to late-infancy foods, resembling the food of the family with higher fiber and protein content, which selects for species within the *Lachnospiraceae*. The fact that Shannon diversity, richness, and evenness correlate with the transition to family foods indicates both that the number of different microbes increases and that their mutual distribution even with the progression toward family foods. This might be interpreted as a sign of increased gut microbial stability. In adults, whey protein and meat were recently found to be positively correlated with gut microbial alpha diversity (46), and a high fiber content (e.g., arabinoxylans) of rye bread has been shown to increase butyrate concentration in feces (47) and plasma (48) and may also contribute to an increased alpha diversity of the microbiota by overall increasing the diverse group of bacterial butyrate producers (49). Consumption of both protein and fiber increases during the complementary feeding period and might therefore represent new energy sources for gut microbes at an infant age of 9 months. This may result in selective advantages for specific microbes to establish in the gut, which will increase alpha diversity.

Although we also investigated associations between anthropometrical data and gut microbiota features, we found no significant correlations after correction for multiple testing. This might be explained by the relative healthy conditions of infants included in the SKOT cohorts. However, the body of recent evidence for a relationship between early gut microbial diversity and metabolic, autoimmune, and allergic diseases (9) emphasizes that our findings are relevant for elucidation of the relationship between complementary diet, gut microbiota establishment, and disease risk. In summary, we conclude that maternal obesity per se does not impact gut microbial changes during late infancy and early childhood substantially. Rather, variation in late-infancy gut microbiota is affected by breastfeeding, complementary diet, and the transition toward family foods with high protein and fiber contents, such as meat, cheese, and rye bread. An increased understanding of the influence of the complementary diet on the development and establishment of the infant gut microbiota provides us with tools to tailor a beneficial progression of our intestinal microbial consortium.

**MATERIALS AND METHODS**

**SKOT cohorts.** The present study used samples from the SKOT cohorts, in which 311 (SKOT I) and 184 (SKOT II) Danish children were followed for the first 3 years after birth, with the overall aim of investigating relationships between early diet, growth development, and later disease risks, especially obesity and metabolic syndrome. The study protocols were approved by the Committees on Biomedical Research Ethics for the Capital Region of Denmark (H-KF-2007-0003 and H-3-2010-122). In SKOT I, infants from a random sample of mothers were recruited, whereas in SKOT II, only infants of obese mothers (BMI > 30 kg/m²) were recruited (25). Inclusion criteria in both cohorts were single birth and full-term delivery, an absence of chronic illness, and an age of 9 months ± 2 weeks at the first visit. Recruitment of participants for SKOT I was done by postal invitations sent to 2,211 randomly selected mothers of infants based on the National Civil Registration System (24). A positive response was obtained from 330 (15%), but 19 dropped out before the first examination; thus, 311 individuals were included in the cohort. With exclusions of individuals with fecal samples taken at only one of the two time points (9 or 18 months of age), individuals using antibiotics, or individuals for which we had inappropriately stored fecal samples (e.g., samples stored at room temperature, for which the storage time was >24 h, or that defrosted before delivery to the university), a randomly selected subset of 114 samples was used in the present study. In SKOT II, 184 infants of the invited 208 obese pregnant women participating in the TOP (treatment of obese pregnant women) intervention study were recruited (50). Using the same exclusion criteria for fecal samples that were used in SKOT I, a subset of 113 infants was included in this study. Collection of data was carried out from 2007 to 2010 and from 2011 to 2014 for SKOT I and SKOT II, respectively. Official Danish guidelines for infant feeding did not change during this period. Participants...
in both cohorts were examined at 9 months (±2 weeks), 18 months (±4 weeks), and 36 months (±12 weeks) of age, and fecal samples and information on body composition, food questionnaires, and household interviews were collected at these examinations during the study. Several studies, not including microbiota assessments, have been published previously on these cohorts (24, 25). For SKOT I, one paper reporting on the microbiota assessed by a qPCR-based approach (8) and one study reporting on the relation between microbiota data from SKOT I and nondietary factors affecting gut microbiota and prevalence of atopic symptoms (51) are available.

**Anthropometry and body composition.** Birth weight and length measurements were taken by midwives and obtained from health records. Weight, length, waist circumference, and subscapularis skinfold thickness measurements at 9 and 18 months were taken at the Department of Nutrition, Exercise and Sports, University of Copenhagen, by trained research staff. Using a digital scale (at 9 months, Sartorius® 65 [Sartorius AG, Göttingen, Germany]; at 18 months, Lindeltronic 8000 [Samhall Lavi AB, Kristianstad, Sweden]), weight was measured, without clothes, to the nearest 0.1 kg. Recumbent length was a mean of three measurements carried out with a digital measuring board (Force Technology, Brøndby, Denmark), which made readings to the nearest 0.01 cm. Skinfold thickness was measured to the nearest 0.1 mm by a Harpenden skinfold caliper (Chasmos Ltd., London, United Kingdom), and we used the mean of the three measurements. Recumbent waist circumference was measured to the nearest millimeter at the level of the umbilicus with a nonstretchable tape measure (Lasso; Child Growth Foundation, London, United Kingdom). Weight, length, subscapularis skinfold thickness, and BMI were converted to 2 scores, with the World Health Organization growth standards used as a reference and with the software program World Health Organization Anthro (52).

**Food questionnaire.** As described previously (25), the infant diet was recorded by parents at the age of 9 months using validated 7-day food records (42). Portion sizes were estimated with household measures and food photograph series and noted in a precoded food diary. All intakes of energy, nutrients, and food items recorded in the precoded food record were calculated for each individual using the software system GIES (version 1.000d; National Food Institute, Søborg, Denmark), a system developed at the National Food Institute, Technical University of Denmark, and the Danish Food Composition Databank (version 7; National Food Institute [http://www.Foodcomp.dk]). Quality control was carried out by trained research staff before data were entered in the database. Possible over- and underreporters were identified on the basis of the estimated daily energy requirement of 338 kJ/kg for both genders, an average between the 6- and 12-month estimates and cutoff values of ±46% (53). The food groups (Table 2) were selected on the basis of nutritional knowledge in an attempt to cover most aspects of the official recommendations, nutrition evidence, and typical infant diet in Denmark. Food groups were named with a short, compressed description, such as “RyeBread” and “SugaryDrink.” Intake of breast milk was calculated as the number of breastfeedings per day, using a rough estimate of 99 g per feeding (54). The intake (grams/day) of all food groups was divided by total body weight (in kilograms) for each participant.

**Information extracted from parental background interviews.** Information on sex, socioeconomic status, education level, and household income, prevalence of C section, gestational age at birth, prior use of antibiotics and other medication, durations of exclusive and total breastfeeding, and age at introduction to complementary feeding (Table 1 and see Tables S2 to S4, S6, and S7 in the supplemental material) were collected from background interviews with parents at the 9- and 18-month visits. Use of antibiotics was recorded 14 days prior to the 9- and 18-month visits. Exclusive breastfeeding was defined as receiving only breast milk, water, and vitamins.

**Fecal samples, DNA extraction, and PCR amplification of the V3 region of the 16S rRNA gene.** Fecal samples obtained at 9 and 18 months of age were freshly delivered on the morning of visitation or had been stored in the participant’s home, in provided freezer containers, either in the freezer (−18°C) or in the refrigerator (4°C) for maximally 24 h before delivery to the University of Copenhagen’s Department of Nutrition, Exercise and Sports, where they were stored at −80°C until DNA extraction. Samples were randomized across cohorts (nSKOT 1 = 10, nSKOT 2 = 10) for each DNA extraction round (nplate = 20). DNA was extracted (12855-100 PowerLyzer PowerSoil DNA isolation kit; Mo Bio) from 250 mg feces according to the protocol provided by the manufacturer with minor modifications: bead beating was performed at 30 cycles/s for 10 min (Retsch MM 300 mixer mill), and the initial centrifugation steps were performed at 10,000 × g for 3 min, as recommended for clay matter. DNA quantity and quality were measured by the Qubit double-stranded-DNA (dsDNA) BR assay (Invitrogen; Q32850) and with a NanoDrop 1000 (Thermo Scientific), respectively, yielding on average 32.7 ± 21.4 ng/μl DNA with an A260/A280 equal to 1.81 ± 0.12 and an A260/A926 equal to 1.60 ± 0.39. The PCR amplification of the V3 region of the 16S rRNA gene was performed with 5 ng community DNA as the template, using 0.2 μl Phusion high-fidelity (HF) DNA polymerase (Fisher Scientific; F-553L), 4 μl HF buffer, 0.4 μl dNTPs, and 0.4 μl deoxyribonucleoside triphosphate (dNTP) (10 mM of each base), 1 μM forward primer (PBU [primer bacterial universal] 5′-A-adapter-TACAG-barcode-CCTACCGGAGGCGACAGC-3′) and 1 μM reverse primer (PRB [primer bacterial reverse] 5′-trp1-adapter-ATTACCGCGGCTGCTGG-3′) in a 20-μl total reaction volume. Both primers include sequencing adaptors, and the forward primer additionally includes a unique 10- to 12-bp barcode (IonXpress barcode adapters). The PCR program included 30 s at 98°C, 24 cycles of 15 s at 98°C and 30 s at 72°C, and then 5 min at 72°C. The PCR product was purified by use of HighPrep PCR magnetic beads (MagBio; AC-60005) with the 96-well magnet stand (MagBio; MyMag 96), according to the prescribed procedure. DNA quantity was measured using the Qubit dsDNA HS assay (Invitrogen; Q32851), and samples were pooled to obtain equimolar libraries containing up to 90 samples (randomized across cohorts and age) in each library.
DNA sequencing and data handling. Sequencing of the 16S rRNA gene libraries was performed using the Ion OneTouch and Ion personal genome machine (PGM) systems with an Ion 318 chip kit, generating 5 to 7 million reads per chip with a median length of 180 bp. Sequencing data were imported into CLC Genomic Workbench (version 7.0.3, CLC bio; Qiagen, Aarhus, Denmark), reads were quality controlled, demultiplexed, and trimmed to remove low-quality sequences ($P_{base calling error} = 0.05$), ambiguous nucleotides (maximum of 2 allowed), primers, and barcodes and to discard reads below 110 bp and above 180 bp. The sorted and trimmed FASTA files were run through the Ribosomal Database Project classifier (55), with a bootstrap cutoff of 50% as recommended for sequences shorter than 250 bp (56). Chimera removal was not performed, since short amplicon length and a low number of PCR cycles reduces chimera prevalence (57). The total number of reads for each sample was on average 47,544 ± 18,656 (range, 12,749 to 121,070) and was used to calculate the relative abundances of bacterial taxa at the phylum (99.8% classified), family (98.3% classified), and genus (75.2% classified) levels. In the further analysis, a cutoff of 0.01% in mean relative abundance at either 9 or 18 months was set. Based on the detection limit (1 read), a threshold was set to 0.001% (~0.48 reads) and samples with zero reads for a given bacterial taxon were assigned this value. Based on reads assigned to the genus level (average, 36,012 zero reads for a given bacterial taxon were assigned this value. Based on the detection limit (1 read), a threshold was set to 0.001% (~0.48 reads) and samples with zero reads for a given bacterial taxon were assigned this value. Based on reads assigned to the genus level (average, 36,012 ± 16,221; range, 4,384 to 92,980), 8 samples were excluded due to low depth (<10,000 reads) for calculation of alpha and beta diversity measures. In the remaining 446 samples, sequences were rarefied (average of 100 subsamplings) to 10,000 reads/sample. Binary Bray-Curtis dissimilarity and alpha diversity measures (Shannon index, number of observed genera, and Pielou’s evenness index) were calculated for each individual in each cohort at the ages of 9 and 18 months using the R package vegan. Based on principal-coordinate analysis (PCoA) of Bray-Curtis dissimilarities, the distances to group centroids were used as measures of beta diversity, using the function betadisper within vegan.

Statistical tests and correlations. Correlation analyses and statistical tests were done with the GraphPad Prism software (version 5.0.3; GraphPad Software Inc., La Jolla, CA) and R (version 3.1.0, R Core Team 2014 RA language and environment for statistical computing; R Foundation for Statistical Computing, Vienna, Austria). Principal-component/coordinate analyses (prcomp/betadisper) and heatmaps (heatmap.2) were performed in R using the packages ggbiplot, gplot, and vegan. Normal distribution of data was evaluated by the Shapiro-Wilk normality test and visual inspection of histograms. Cohort characteristics were compared by Fisher’s exact test/chi-square test for categorical data and an unpaired t test or the Mann-Whitney test for continuous data. A paired Wilcoxon signed-rank test or paired t test was used to examine the changes in alpha diversity measures and bacterial composition across time, whereas the Mann-Whitney test or unpaired t test was used to compare measures of alpha diversity and bacterial composition between cohorts at the ages of 9 and 18 months. Turkey’s honestly significant differences test was used to evaluate differences in beta diversity across time and cohorts. Spearman’s rank test/Pearson’s correlations and the unpaired t test/Mann-Whitney test were used to investigate associations between gut bacterial composition and alpha diversity measures with birth mode, gestational age at birth, prior use of antibiotics, duration of exclusive/total breastfeeding, age at introduction to complementary feeding, complementary diet at 9 months, and anthropometrics at 9 and 18 months. When indicated, false discovery rate-corrected (58) $P$ values ($q$ values) were applied to correct for multiple testing, with a threshold of 0.05.

Accession numbers. Sequencing data are deposited in NCBI’s Sequence Read Archive with the accession number SRP052851 under BioProject number PRJNA273694.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at http://dx.doi.org/10.1128/mSphere.00069-15.

- Figure S1, EPS file, 2.6 MB.
- Figure S2, EPS file, 2.6 MB.
- Table S1, DOCX file, 0.03 MB.
- Table S2, DOCX file, 0.02 MB.
- Table S3, DOCX file, 0.02 MB.
- Table S4, DOCX file, 0.01 MB.
- Table S5, DOCX file, 0.02 MB.
- Table S6, DOCX file, 0.02 MB.
- Table S7, DOCX file, 0.02 MB.

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