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Persistent effects of early infant diet and associated microbiota on the juvenile immune system

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Keywords: breast-milk, rhesus macaque, gut microbiota, T cell activation, T_H17 cells

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Early infant diet has significant impacts on the gut microbiota and developing immune system. We previously showed that breast-fed and formula-fed rhesus macaques develop significantly different gut microbial communities, which in turn are associated with different immune systems in infancy. Breast-fed animals manifested greater T cell activation and proliferation and harbored robust pools of T helper 17 (T_H17) cells. These differences were sustained throughout the first year of life. Here we examine groups of juvenile macaques (approximately 3 to 5 y old), which were breast-fed or formula-fed in infancy. We demonstrate that juveniles breast-fed in infancy maintain immunologic differences into the fifth year of life, principally in CD8⁺ memory T cell activation. Additionally, long-term correlation networks show that breast-fed animals maintain persistent relationships between immune subsets that are not seen in formula-fed animals. These findings demonstrate that infant feeding practices have continued influence on immunity for up to 3 to 5 y after birth and also reveal mechanisms for microbial modulation of the immune system.

Introduction

The nutrition an infant receives can have lifelong effects, which are partly mediated by dietary influence on the infant microbiota.^{1–7} Previous studies have shown that breast- vs formula- fed

human infants have significantly different gut microbiotas.^{8,9} A study in rhesus macaques demonstrated that differences in early infant diet altered the gut microbiota and cytokine production.¹⁰ We recently carried out the first detailed study of possible effects of the infant microbiota on a wide range of immune cell phenotypes and functions. We showed that breast-fed and formula-fed rhesus macaque infants have significant differences in the gut microbiota that persist through the first year of life.¹¹ For example, breast-fed infants developed more robust pools of T_H17 cells, as well as increased T cell activation and proliferation. These differences in immunologic development may partly explain inter-individual variation in immune responses to infection or vaccination, in infancy or later life.

As a follow-up to our previous study, we have now assessed for how long early infant diet may shape immune cell phenotypes. We investigated the immune systems of previously breast- or formula-fed juvenile rhesus macaques between 3–5 y of age and compared the data to those from our previous infant cohort. The immune systems of juvenile animals were affected by early dietary differences, although the differences between rearing groups were more subtle than was observed in infancy. Striking relationships between immune subsets, shaped by early diet, were documented in network diagrams and these important relationships were sustained into juvenile life. Our findings show that early infant diet has long-standing effects on the immune system,

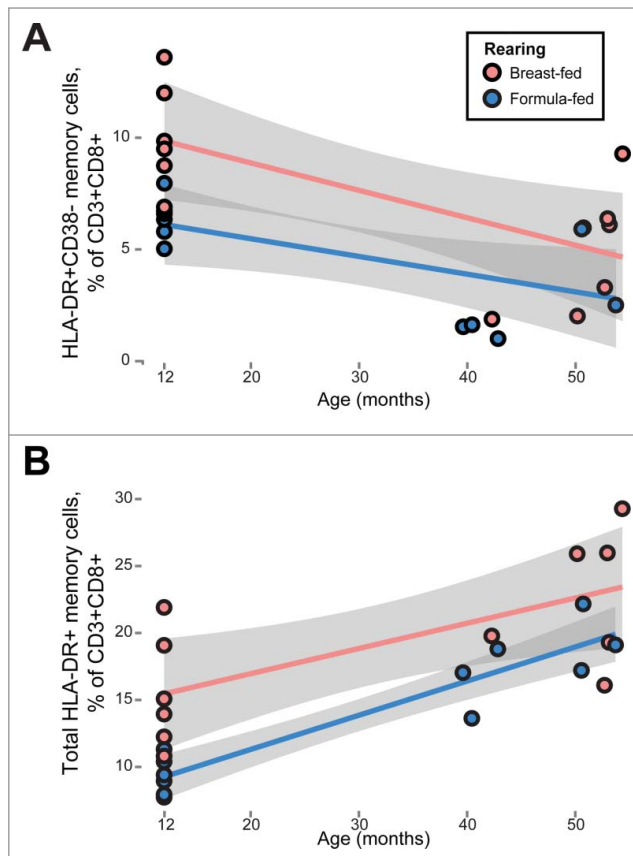


Figure 1. Persistent differences in T cell activation between previously breast-fed and formula-fed rhesus macaque juveniles. Multivariate mixed-effects regression revealed differences in T cell activation between breast- and formula-fed animals (red and blue markers, respectively), while controlling for age and sex. Breast fed animals manifested higher fractions of (A) CD8⁺HLA-DR⁺CD38⁻ memory T cells ($p = 0.004$) and (B) the total HLA-DR⁺ fraction among CD8⁺ memory T cells ($p = 0.0008$). Gray bands represent 95% confidence intervals.

which could result in different responses to immunologic challenges in later life.

Results

Differences in immune subsets shaped by early diet are maintained into juvenile life

To investigate the long-term effects of early infant diet on immune cell subsets, we isolated peripheral blood mononuclear cells (PBMCs) from 12 juvenile rhesus macaques, which were 3–5 y in age (6 breast-fed, 6 formula-fed in infancy). We performed extensive immunophenotyping via flow cytometry, including antibody panels defining more than 115 phenotypes of interest, including antigen-presenting cell and T cell surface, activation, and homing markers. We compared

the data obtained to those from 12-month-old infant macaques (6 breast-fed and 6 formula-fed) in our previous study.

We used multivariate regression to determine immune subsets that were significantly associated with rearing status and/or age, recognizing age as an important covariate.¹¹ Some differences in immune subsets found in infancy were maintained in this older cohort. For example, activated T cell subsets were significantly associated with both rearing status and age, indicating that T cell activation remained higher in breast-fed older animals (e.g. CD8⁺HLA-DR⁺CD38⁻ memory T cells, % of CD8⁺; $p = 0.004$; Fig. 1A). These and other activated subsets had also been seen to be higher in 12-month-old breast-fed macaques, as compared to formula-fed animals. Total CD8⁺HLA-DR⁺ memory T cells

followed a similar pattern (% of CD8⁺; $p = 0.0008$, Fig. 1B), except that age was positively associated with the frequency of these cells.

Other subsets found to be different between rearing groups in our 12-month-old infants were no longer different in the juvenile cohort. These populations include proliferative subsets (Ki-67⁺) and memory and naïve cells within the CD4⁺ T cell compartment (Fig. 2). In both CD4⁺ and CD8⁺ T cells and in the corresponding effector-memory T cells, 12-month-old breast-fed infants manifest higher Ki-67 expression than formula-fed infants. In our juvenile cohort these differences disappear, with the frequency of these cells in previously breast-fed juveniles apparently declining to match that in their previously formula-fed counterparts (Figs. 2A and 2B). In the cases of CD4⁺ memory and naïve T cells (Fig. 2C), similarly, there are significant differences in these subsets at 12 months: the ratio of memory to naïve cells is much higher in breast-fed infants. In previously formula-fed juveniles, this ratio seems to “catch up” with that seen in breast-fed peers, indicating development of CD4⁺ memory T cells to an extent eventually matching that in breast-fed animals (Fig. 2C).

Similar “catch up” seemed to occur with T_H17 cells, in that no significant differences in T_H17 cell pools remained between 3- to 5-year-old macaques that were previously breast- or formula-fed ($p = 0.53$).

Relationships between immune subsets shaped by early diet are sustained into juvenile life

We then assessed the relationships between immune subsets in early and later life. We calculated Spearman correlations between immune variables separately in breast- and formula-fed infant and juvenile animals, that is, in 4 groups separately. We then used Cytoscape (“Merge Networks” tool) to identify correlations that were persistent over time within each rearing cohort. Finally, we looked at the differences between these 2 merged networks (breast-fed vs formula-fed; Fig. 3).

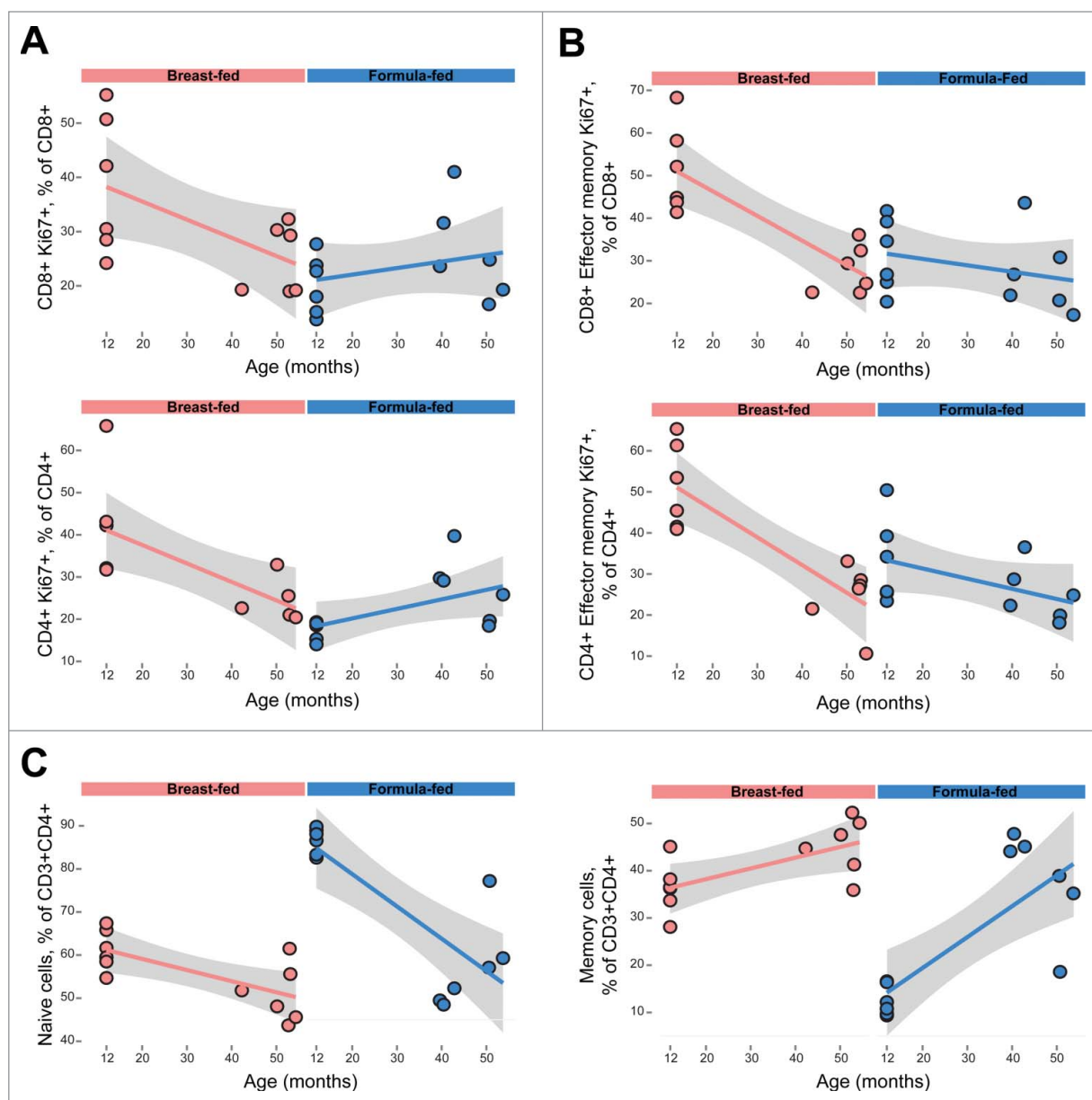


Figure 2. Relative equalization of some immune parameters between rearing groups in juveniles, as compared to infant animals. While infant animals show significant differences between breast and formula fed animals, previously breast- and formula-fed juvenile rhesus macaques show little difference in (A) T cell proliferation ($CD4^+Ki67^+$ and $CD8^+Ki67^+$) (B) Effector memory T cell proliferation ($CD8^+$ and $CD4^+$ shown above and below, respectively) or (C) $CD4^+$ naive and memory T cells. Gray bands represent 95% confidence intervals.

Animals that were previously breast-fed sustained long-term correlation networks not seen in formula-fed animals (Fig. 3). Activated $CD8^+$ T cells ($CD38^+HLA-DR^+$; purple nodes in Fig. 3A) were positively correlated with IL-17-producing $CD4^+$ T cells (T_H17 cells) and memory T_H17 cells producing interferon gamma ($IFN\gamma$; light pink nodes; Fig. 3A). Frequency of these T_H17 cells, in turn was correlated with that of IL-17-producing $CD8^+$ T cells

(T_C17 cells, orange nodes). In addition, in this same breast-fed group, we identified sustained correlations between proliferative T cells ($Ki67^+$, green nodes) and extracellular markers of gut-homing T_H17 cells ($CCR4^+$ and $CCR6^+$ cells in Figure 3B, darker pink nodes). None of these networks were found in the previously formula-fed group, indicating that the relationships involved were not as dominant or consistent after formula feeding.

Discussion

In this follow-up to our early infant diet cohort study, we found that some immunologic differences seen in infancy are sustained into juvenile life. Markers of T cell activation remained associated with the rearing status and age of an animal, especially those marking activation of $CD8^+$ memory T cells. Other immunologic features became more uniform between the infant and juvenile periods,

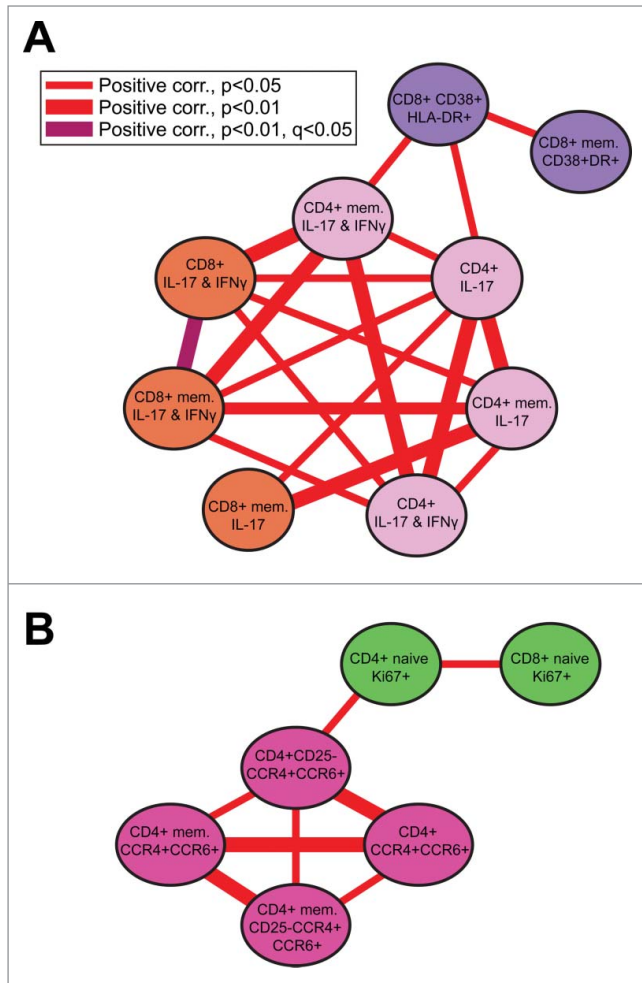


Figure 3. Breast-fed animals maintain correlation networks that formula-fed animals do not. Correlations between immune subsets seen in our breast-fed cohorts (infant and juvenile) were not observed in formula-fed animals. **(A)** Activated T cells (purple) correlate positively with T_H17 cells and memory T_H17 cells producing $IFN-\gamma$ (light pink), as well as with T_C17 cells (orange). **(B)** Proliferating T cells (green) correlated positively with extracellular markers of gut-homing T_H17 cells (dark pink). Formula-fed animals do not maintain either network.

suggesting that the effects of other exposures came to dominate those that were earlier attributable to infant feeding practices. Markers of T cell proliferation and relative proportions of memory and naïve T cells had equalized between rearing groups by the juvenile stage. The exact timing of and impetus for these changes are unknown but will be revealed by careful longitudinal studies.

When we investigated the relationships between immune subsets over time, we found that breast-fed animals maintained correlation networks that formula-fed animals did not. Breast-fed animals maintained positive correlations between T cell

activation and proliferation and T_H17 cells. These sustained relationships are likely driven by the influence of breast-feeding in infancy. Possibly, for example, T_H17 cell development is driven in breast-fed animals by characteristics of the early infant microbiota, via effects on T cell activation, while in formula-fed animals later influences such as viral infection are more important. We did not analyze the gut microbiota for the juvenile cohort. Studies in humans have shown that the gut microbiota differences imposed by different early diets dissipate over time.^{12–15} Possibly, homogenization of the gut microbiota between rearing groups (all

animals in this study received identical diets) could explain homogenization of many immune cell subsets.

Possible consequences of early infant feeding for infection and inflammation in later life

These new data suggest that effects of early infant diet may be different at various subsequent life stages.

Infancy

The largest immunologic effects of early infant diet were observed at 9–12 months of age, a time period that in humans is considered to be part of infancy. Our studies demonstrate a link between breast feeding, the associated breast feeding-specific microbiota, and T cell activation and proliferation, as well as increases in T_H17 cell pools. T_H17 cells are important modulators of intestinal immunity whose development is induced by gut commensal bacteria.¹⁶ Larger pre-existing T_H17 cell pools have been associated with lower viral loads when rhesus macaques are infected with SIV.¹⁷ The increased T cell activation and proliferation set points may be caused by stimulation of the T cell compartment by a more diverse gut microbiota or may be associated with specific bacterial colonizers or their metabolites.¹¹

Juvenile to early adulthood

By 3–5 y of age, previously breast- vs. formula-fed macaques demonstrated no detectable differences in T cell proliferation, or in levels of memory and naïve T cells. However, differences in T cell activation were sustained, suggesting that infant feeding could be determinative of the quality of certain immune responses even after a long period of time. Hosts with increases in these cells may manifest more rapid responses against infection.^{18,19} Alternatively or in addition, the altered T cell activation may be reflective of changed antigen presenting cell function, perhaps due to durable changes in the microbiota.²⁰

Later adult life

We have not yet assessed immunologic differences in adult macaques that may be ascribed to infant feeding. Given the data

reported here on juvenile animals, which are suggestive of waning effects of infant feeding and possibly greater effects of more proximal exposures, a large study may be required to understand lifelong effects of infant diet. However, the long-term connections between immune subsets, reflected in correlation networks, may indicate that early diet has a long-term impact that is sustained by long-term effects on the microbiota. Different microbiotas and associated drivers of adaptive immune cell subsets would suggest the likelihood of different responses to interventions targeting the immune system. Among previously breast-fed animals, but not formula-fed animals, T cell activation (HLA-DR⁺) and proliferation (Ki-67⁺) were positively associated with T_H17 cells. Thus, future therapies that might induce T_H17 cells through T cell activation and proliferation may be more effective in breast-fed individuals.²¹

Breastfeeding provides important protection against infectious diseases, which account for over two-thirds of the 12 million annual deaths in children younger than 5 y in less developed countries.²²⁻²⁷ Nonetheless, we know little about the specific immunophenotypic changes established by breast-feeding that confer such advantages, or the mechanisms by which those differences shape an infant's risk of disease. We can infer specific risks or protective effects where mechanisms are well understood.²⁸ For example, animals with increased T cell activation (as a result of previous breast-feeding) may have more rapid responses to infection or quantitatively greater responses to vaccination.^{18,19} Such T cell activation, however, might be detrimental in the setting of chronic HIV infection.²⁹ Setting aside such specific cases, we still know relatively little about how natural immune variation in healthy individuals contributes to disease risk overall. To understand the impact of such inter-individual variability, more longitudinal and interventional experiments must be performed.

Our data show that early infant diet has a profound and long-lasting effect on the immune system. The initial divergence is powered by differing gut microbiotas

developing in breast- and formula- fed infants. While the effects do wane over time, various immune subsets seem to equalize at different rates. Factors other than the infant microbiota, such as cytomegalovirus infection, also contribute important variability to the immune system.³⁰ Thus, although work remains to be done in order to clarify the interactions between early and later exposures, large differences attributable to infant feeding doubtless explain some of the immune variation seen between individuals, particularly between infants.³¹

Materials and Methods

Ethics statement

This study was performed under strict compliance with the NIH Guide for the Care and Use of Laboratory Animals. Established policies of the Institutional Animal Care and Use Committee (IACUC) of the University of California, Davis were followed for sample collections, housing, and medical care. The study was performed at the CNPRC, which is one of 8 centers supported by the National Institutes of Health, Office of the Director (NIH/OD), and is accredited by the Association for the Assessment and Accreditation of Laboratory Animal Care International (AAALAC). The CNPRC houses more than 5,000 non-human primates, most of which are rhesus macaques (*Macaca mulatta*), in outdoor corrals or indoor cages.

Study design

Twelve infant and 12 juvenile rhesus macaques were used for study of immune cell phenotypes. The infants were the same used and described in ref. 11. Juvenile animals were housed in similar outdoor environments. Three of the previously formula-fed animals were specific-pathogen free (SPF) and were therefore seronegative for Herpes B virus, simian immunodeficiency virus (SIV), Type D retrovirus and simian T-lymphotropic virus.³² These 3 were not significantly different from non-SPF formula-fed juveniles in immune variables, as tested by Wilcoxon rank sum test. Our low sample size of previously formula-fed

juveniles limits that result, and may warrant further study. Other than SPF status, juvenile animals had no environmental differences between rearing groups besides their early feeding practices.

Sample collection

Blood samples were taken from each rhesus macaque infant 12 months after birth as well as juvenile macaques aged 3–5 years, as cited in ref. 11. Peripheral blood mononuclear cells (PBMCs) were isolated by gradient density purification using Lymphocyte Separation Medium (LSM; MP Biomedicals, LLC, Solon, OH), then washed in medium containing FBS (fetal bovine serum) and 10% DMSO, then cryopreserved in liquid nitrogen.

Immune cell phenotyping by flow cytometry

Frozen rhesus macaque PBMCs were thawed and stained for flow cytometry (including intracellular staining) as previously described in ref. 11.

Statistical analysis

Immunophenotyping data from 12-month-old rhesus macaques (obtained in our previous study) and from our juvenile cohort were used for analysis. All analysis was performed in R, and all plots were created using *ggplot2*. We used multivariate mixed-effects regression (using *lmer* in the *lme4* package) to assess variables significantly associated with age and rearing status, comparing to a null model testing age only. Significance of random effects was assessed based on the likelihood ratio-test between nested models. Differences between rearing groups at individual time points were calculated using Wilcoxon rank sum test.

To assess long-term correlations between immune subsets, we first separated data into 4 groups based on age cohort (infant vs juvenile) and rearing status. We calculated Spearman correlations between variables. We then used Cytoscape to merge networks based on rearing status and display the resulting networks using the “Merge Networks” tool.³³ Final networks were drawn in Illustrator based on Cytoscape results.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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Author Contributions

AA and DHO designed the study. AA, NN, and DHO performed the statistical analysis. NN, AA, GML, and DHO wrote the manuscript. GML performed the flow cytometry assay and analysis. DL and KKAVER collected the samples.

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