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A Scoping Review of Gut Microbiome and *Bifidobacterium* Research in Zimbabwe: Implications for Future Studies

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Background: Gut microbiota play a key role in host health, with certain *Bifidobacterium* strains critical for immune development. The healthy gut of breastfed infants is dominated by these pioneer microbes, especially the strains that feed on human milk oligosaccharides.

Objective: This is a scoping review of gut microbiome research from Zimbabwe. It focuses on distribution and dynamic changes of bifidobacteria, and milk components that promote growth of microbes in infants, together with the distribution of associated gut microbes in adults.

Design: Online databases were searched for publications from 2000 to 2023.

Results and Analysis: Fourteen publications on microbiota of infants and adults were included in this scoping review. Most were cross-sectional, while three were clinical trials/cohort protocols. Publications focused on pediatrics (78.5%), pregnant women (14.3%), and men (7.2%). Zimbabwe has a high burden of HIV; hence 35.7% of study populations were delineated by HIV status. The laboratory methods used included shotgun metagenomics (62%) or 16S rRNA gene amplicon sequencing. Almost 85% of the studies focused on total microbiome profiles and rarely reported the distribution of different *Bifidobacterium* species and variants. None of the papers studied human breast milk composition. There were reports of reduced abundance of beneficial genera in pregnant women, children, and adolescents living with HIV. Additionally, gut microbiota was reported to be poorly predictive of child growth and vaccine response, though this was not conclusive.

Conclusion: There are few studies that characterize the gut microbiome by Zimbabwe-based researchers. However, studies on strain level diversity of *Bifidobacterium* and other key microbes, and their role in health during and beyond infancy, lag behind in Zimbabwe and other low- and middle-income countries. Such cohorts are needed to inform future mechanistic studies and downstream translational work such as next-generation probiotics and prebiotics.

Keywords: *Bifidobacterium*, gut microbiome, human milk oligosaccharides, LMIC, prebiotics, probiotics, Zimbabwe

Introduction

The Infant Gut Microbiome

Untangling what constitutes a healthy gut microbiome is timely, particularly in low- and middle-income (LMIC) settings where problems of infant nutrition, chronic diarrhea, and failure to thrive are high priority.^{1,2} Studying the infant gut microbiome is also crucial as many of the events capable of shaping microbial communities (eg birth mode, antibiotic use, and feeding type) take place during this phase of life.^{2,3} The healthy infant gut microbiota is known to be dominated by the genus *Bifidobacterium* with certain species particularly dominant during lactation: *B. breve*, *B. bifidum*, *B. longum* subsp. *longum*, and *B. longum* subsp. *infantis*.^{2,4} Numerous studies have indicated that certain species and strains have immune-modulatory roles, eg *B. breve* (also a common breast milk-associated species). This species has been reported to

have anti-allergic properties and is generally absent from allergic infants.^{5,6} Additionally, various prospective studies performed in cohorts of allergic infants and formula-fed infants found less bifidobacteria, and this may link to negative health outcomes. Interestingly, certain adult-type *Bifidobacterium* species, mainly *B. adolescentis*, *B. catenulatum*, and *B. pseudocatenulatum*, appear to be more prevalent in allergic and formula-fed infants, but this requires further inquiry.^{2,6,7} Within LMICs, previous work has indicated that malnutrition in children causes depletion of bifidobacteria, with enhanced colonization of potential pathogens such as *Streptococcus* spp., *Fusobacterium mortiferum*, and *Escherichia coli*, which is associated with diarrhea and malabsorption of essential nutrients.⁸ In adults, presence of *Bifidobacterium* is also associated with “health”, and particular strains have potential roles in the reduction of serum cholesterol, alleviation of lactose intolerance, and treatment of inflammatory bowel diseases, acute diarrhea, colorectal cancer, and other intestinal infections.^{7,9}

Sources of Beneficial Bacteria or “Probiotics”

Probiotics have been isolated from breast milk and fecal samples of infants, eg *Bifidobacterium*, *Lactobacillus*, and the yeast *Saccharomyces boulardii*.¹⁰ Other good sources of beneficial *Lactobacillus* are dairy and dairy-related products, including fermented milk and kefir grains. Non-dairy fermented substrates, for instance Nigerian fermented foods, brines of naturally fermented Aloreña green table olives, meat, and fruits, are also sources of potential probiotic strains, eg *Lactobacillus*, *Staphylococcus carnosus*, or *Weissella*.¹⁰ Indeed, *Lactobacillus buchneri* P2, isolated from pickled juice, has probiotic properties, such as cholesterol reduction, and acid tolerance.^{9,10} Animals also harbor beneficial bacteria, and several species, eg pigs, rats, and poultry, insects, eg honeybees, and fish are potential probiotic strain sources.¹⁰

The Potential Roles of Probiotics and Prebiotics

The major difference between vaginally delivered and exclusively breastfed infants, and those who are not, is their access to natural prebiotics (eg human milk oligosaccharides) and probiotics (eg maternal and breast milk microbes).^{9,11} The human milk oligosaccharides (HMOs) found in breast milk are not digested by the infant, but rather they favor the growth of certain *Bifidobacterium* species and strains, eg *B. longum* subsp. *infantis* and *B. bifidum*.^{12–14} HMOs have other important roles, eg acting as antivirals and antimicrobials, modulating the host immune response, improving gut barrier function, preventing pathogen attachment to mucosal surfaces, developing the immune system, modulating intestinal cell responses, and lowering the abundance of *Enterococcus* spp. and *Escherichia coli*.^{1,8,13}

Most HMOs contain a poly lactosamine or lacto-N-biose core and lactose at the reducing end. The core and lactose structures are often linked to fucose (70%) and sialic acid (30%).¹³ HMO metabolism is observed in various members of the *Bifidobacterium* genus, eg *B. longum* subsp. *infantis* and *B. bifidum*, which is linked to specific HMO degradation enzymatic gene clusters. Such HMO consumers contain all the glycosyl hydrolases required for catabolizing a whole spectrum of HMO linkages.^{13,14} However, some individual strains of *B. breve* display poor growth on HMOs but are capable of utilizing more simple structures such as lacto-N-tetrose, and some adult-type strains such as *Bifidobacterium adolescentis* are incapable of digesting HMOs.^{8,13,14} Alongside *Bifidobacterium*, HMO assimilation pathways have been described for a number of gut microbes belonging to other taxa, eg *Bacteroides* and *Akkermansia*.^{15,16}

Randomized controlled trials of prebiotic or probiotic interventions have reported significant weight gain in infants fed formula milk containing *Bifidobacterium animalis* subsp. *lactis*, and fructogalacto-oligosaccharides.¹⁷ *B. animalis* subsp. *lactis* is a common probiotic species believed to provide health benefits when consumed, and also improves bone density,¹⁷ while fructogalacto-oligosaccharides are prebiotics that have structural similarities to certain HMOs, that promote the growth of normally abundant, beneficial microbes in the human intestine.^{13,17} Hence, microbiome modulations using HMOs and some *Bifidobacterium* strains are potential nutritional and therapeutic targets.¹³

Some strains of *Bifidobacterium* are included as probiotics in functional dairy products, food supplements, and pharma products.¹⁸ This is because of work indicating they may promote health, and the fact that alteration in their number and composition is a frequent feature in intestinal diseases such as inflammatory bowel disease, colorectal cancer, or irritable bowel syndrome, as well as extra-intestinal pathologies, such as those affecting the liver or the respiratory tract (eg, allergy, bronchial asthma, and cystic fibrosis).¹⁹ However, the degree of scientific evidence regarding the effectiveness of probiotics is still insufficient, while evidence from African populations is still intermittent.

It has been hypothesized that perturbation of gut *Bifidobacterium* levels and diversity accounts for differences in immune development and vaccine response.^{2,20} Understanding the distribution and dynamics of *Bifidobacterium* species and the roles of different *Bifidobacterium* strains and HMOs is important for design of context-specific interventions, focusing on immune conditions. For instance, the introduction of *Bifidobacterium* in clinical studies showed that supplementation with breast milk-metabolizing *B. bifidum* was associated with long-term establishment and increased *Bifidobacterium* diversity in the preterm gut.²¹ In contrast, adding complementary foods to breastfeeding babies has been shown to radically alter the infant intestinal community, leading to a mature, adult-like microbiota.^{2,22} In light of the above evidence, this scoping review was carried out to systematically map the research performed in the growing area of gut microbiome in Zimbabwe, with focus on *Bifidobacterium* composition, as well as to identify any knowledge and methodological gaps.

Objectives

The following research question was formulated: What is known from published literature about gut *Bifidobacterium* and associated gut microbiota, from Zimbabwe? What is the evidence of the gut microbiota evolution in response to diet, infection, and environmental exposures, and what is its contribution to infant growth, susceptibility to viral infections, oral vaccine response, and adult health? What are the implications of a perturbed microbiome for both infant and adult health and disease, respectively? Are there any reports on the fortification of foods with specific *Bifidobacterium* strains, other potential probiotics, and human milk oligosaccharides?

Method

Protocol Development

Our protocol was drafted using the Preferred Reporting Items for Systematic Reviews and Meta-Analysis Extension for Scoping Reviews (PRISMA-ScR)²³ and reviewed by two members of the research team (see Figure 1). Ethics approval was not required for this scoping review.

Eligibility Criteria

To be included in this scoping review, papers needed to have been: published between 2000 and 2023, involve human participants from Zimbabwe, measure or incorporate one or two laboratory methods used in gut microbiome studies, and either be an original research or protocol (not review).

Information Sources

Searches were performed in databases and trials or study registries relevant to the research question, ie Science Direct, PubMed, and Google Scholar. The reference lists or bibliographies of included studies were searched, and for unpublished data or grey literature we contacted content experts and authorities in the field. Two co-authors then screened titles and abstracts. We made a final decision on the inclusion criteria for studies one month before completion of this scoping review.

Search Strategy

The following search terms were used: “gut microbiome research Zimbabwe”. Appropriate controlled vocabulary and free text terms were used to refine searches in the online databases.

Results and Analysis

Data Charting and Data Analysis

Data items relevant to the review questions were extracted. Data charting involved the charting of authors' or studies' key information, year of publication, country where the study was conducted, aims/purpose, population and sample size, methodology, study design, and findings that relate to the scoping review questions. Data analysis was undertaken via

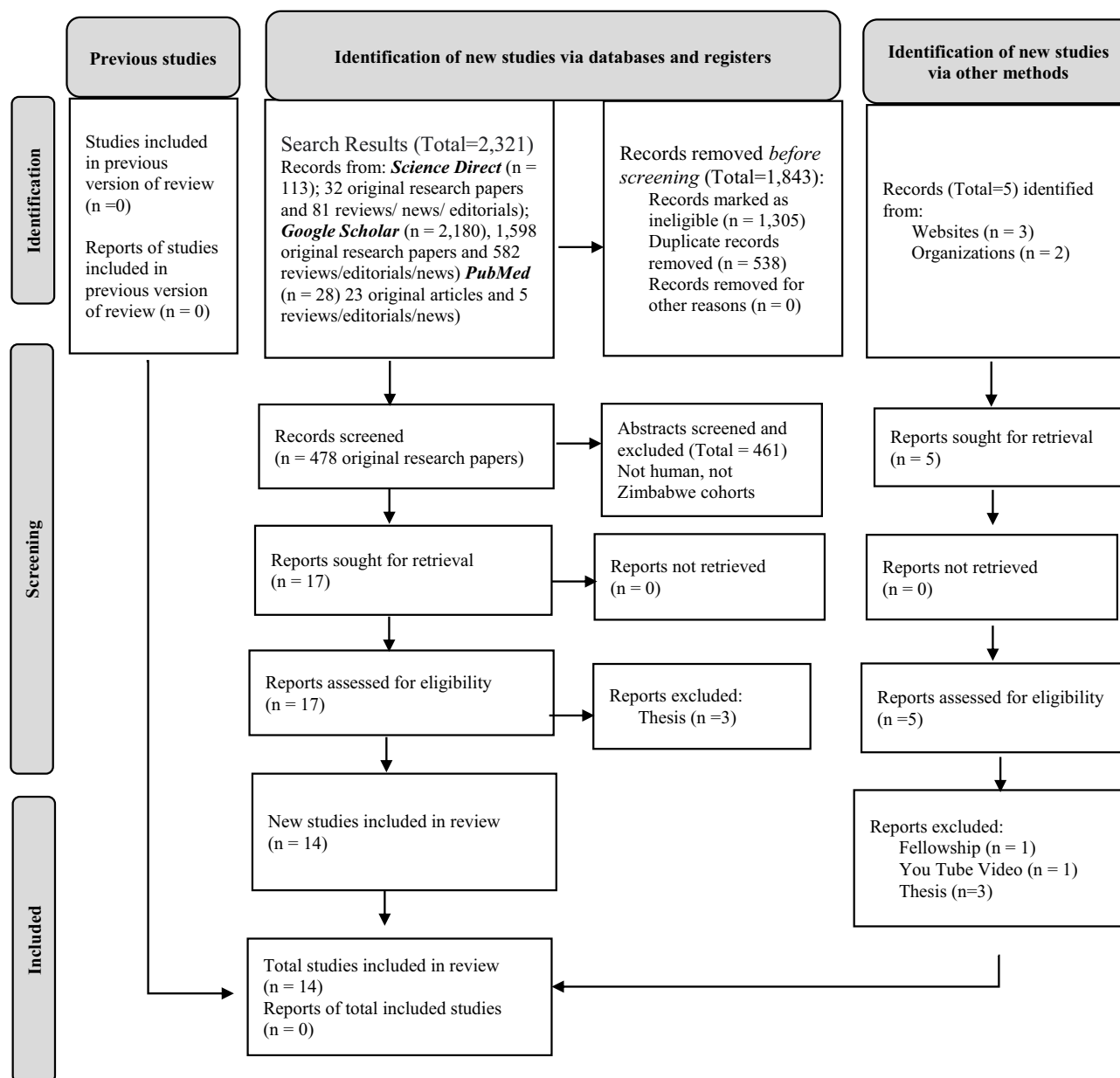


Figure 1 Prisma Flow Diagram for Scoping Reviews.

Notes: Adapted from Page MJ, McKenzie JE, Bossuyt PM, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. *BMJ*. 2021; 372:n71.²³

frequency counts and basic content analysis. The data were presented using tables, boxes, and narrative summaries to address the scoping review's objectives, and to answer the questions of the review.

Summary of Results

Our searches returned over 2320 manuscripts. We removed duplicates and ineligible studies. Two authors then screened 461 remaining abstracts for eligibility. Most of the screened abstracts were ineligible for being either from elsewhere in Africa, other than Zimbabwe, or being focused on wild animals. Out of 17 eligible publications, three were thesis reports and were excluded, therefore only 14 unique publications were retrieved. Of the 14 studies included, two are clinical trial protocols, one is a prospective cohort protocol, nine are cross-sectional study reports, and two are cohort study reports. A total of 1711 stool specimens were analyzed for gut microbiota composition, by various researchers, in collaboration with experts from South Africa, Switzerland, UK, and USA. None of the microbiota assays were carried out locally,

which is a reflection of both skills and technological gaps, in the Zimbabwe setting. As a result of a dearth of research laboratories the total number of relevant papers was low by international standards. Studies on prebiotics (human milk oligosaccharide) and food sources of probiotics are not yet apparent. The few publications that reported on *Bifidobacterium* suggested reduced abundance and no association of these beneficial genera with child growth and vaccine response, in Zimbabwean populations.^{24–26} Each of the 14 publications is summarized in Table 1 and in the narrative summaries below.

The 2015 publication by Gough et al presented the protocol for the Sanitation Hygiene Infant Nutrition Efficacy (SHINE) Trial in Zimbabwe, and how it was designed to provide a platform for examining linear growth, anemia, and the evolution of the gut microbiota in response to diet, infection, and environmental exposures.²⁴ The protocol outlines the hypotheses guiding the project and the analytic approaches to be used, such as the longitudinal design, randomized interventions, and advanced DNA sequencing technologies including shotgun DNA (metagenomic) and RNA (meta-transcriptomic) sequencing. The follow-up paper by Humphrey et al described the protocol in greater detail and shared

Table 1 Showing Results of the Systematic Literature Search; Fourteen Original Research Papers Were Retrieved.^{24–37}

Author, Year, Study Name, if Applicable	Participants N, Median Age (IQR) or Mean (SD) or Range	Study Design, % Female, and Participants and Site Description, Gene Profiling Method(s) and Other Lab Tests	Reported	General Findings	Specific Findings and Shortcomings
Gough et al, 2015, The Sanitation Hygiene Infant Nutrition Efficacy (SHINE) Trial Protocol ²⁴	N = 5280 women recruited, and 3791 live-born children were assessed (Mother–baby pairs).	Trial in rural area without specified sex distribution for infants, participants randomized to improved water, sanitation, and hygiene (WASH), and improved infant and young child feeding (IYCF), Shotgun metagenomics sequencing.	SHINE trial tested the independent and combined effects of improving infant diet and household WASH on length-for-age Z-score and hemoglobin (primary outcomes) at 18 months of age.	Secondarily tested the gut microbiome composition of mothers and babies in various publications on gut microbiome. ^{24,25,28}	N/A
Gough et al, 2021, The SHINE Trial Sub-study ²⁵	N=207, Mothers Mean ages: 31.2 (5.9) for those living with HIV, 27.2 years (6.7) for those without HIV.	Cross-sectional sub-study in a trial, 100% of them female. Participants being rural-based mothers and HIV-exposed and unexposed infants, Shotgun metagenomics sequencing.	Relationship between mothers' microbiome and infants' gestational age, birth weight, or neonatal growth.	Maternal fecal microbiome enriched for metabolizers of resistant-starch, including <i>Ruminococcus bromii</i> and <i>Faecalibacterium prausnitzii</i> , plus <i>Prevotella copri</i> abundant in populations feeding on high fibre diets.	Few <i>Bacteroides</i> indicating meat deficiency, resistant-starch degraders, were an important contributor to birth weight and neonatal growth, and to a lesser extent gestational age. A large proportion of DNA was unknown or unidentifiable.
Robertson et al, 2021, The SHINE trial Sub-study ²⁶	N=158, infants Median age = 43 days (35.0–68.0).	Cross-sectional sub-study of infants, from a rural trial, 43.6% female, within 30 days of an oral rotavirus vaccine (RVV), Shotgun metagenomics sequencing and anti-rotavirus IgA.	Stool metagenomes and their association with RVV immunogenicity.	RVV was poorly immunogenic, no association between gut microbiome and RVV seroconversion. However, maternal recall for some metadata led to low-quality data.	<i>Bifidobacterium longum</i> most abundant species, followed by <i>Escherichia coli</i> , <i>B. bifidum</i> , and <i>B. pseudocatenulatum</i> among others. Cross-sectional study design has inherent flaw of not distinguishing momentary changes from enduring ones in the gut microbiota.

(Continued)

Table I (Continued).

Author, Year, Study Name, if Applicable	Participants N, Median Age (IQR) or Mean (SD) or Range	Study Design, % Female, and Participants and Site Description, Gene Profiling Method(s) and Other Lab Tests	Reported	General Findings	Specific Findings and Shortcomings
Robertson et al, 2023, The SHINE Trial Sub-study ²⁷	N=335 infants, Age range = 1–18 months.	Cohort sub-study in a trial, infants, 43.9% female, from rural area, 1/3 of mothers are living with HIV, Shotgun metagenomics sequencing and nutritional status.	Analyzed gut microbiome of infants in a trial of improved WASH and IYCF.	<i>Bifidobacteria</i> , <i>E. coli</i> , and <i>Bacteroides fragilis</i> species most abundant <12 months of age.	<i>Bifidobacterium longum</i> predominant species up to 12 months of age. Maternal HIV status associated with reduced abundance of <i>Bifidobacterium</i> , and over-maturity, and over-diversification of the early-life gut microbiome in the uninfected children.
Smith et al, 2021, The CHAIN Trial Protocol ²⁸	N=192, Infants aged between 5 and 6 months.	Trial with infants, sex unspecified, from Shurugwi district, in rural Zimbabwe, Unspecified gut microbiome methods.	Infant and young child feeding (IYCF) and gut microbial composition.	Not yet published.	N/A
Flygel et al, 2020, The BREATHE Trial Sub-study ²⁹	N=177 children living with HIV and 103 HIV-uninfected controls, aged 6–16 years.	Cross-sectional sub-study in a trial of children, 47% females, from urban Harare, 16S rRNA (V4 hypervariable region) gene sequencing.	HIV, ART, and gut microbial composition.	Children living with HIV (CLWH) had significantly lower alpha-diversity and higher beta-diversity. Prolonged ART-treatment (≥ 10 years) associated with a richer gut microbiota by alpha diversity.	Enriched levels of <i>Corynebacterium</i> , <i>Finexgoldia</i> , and <i>Anaerococcus</i> in CLWH and enrichment of <i>Enterobacteriaceae</i> in participants with low CD4+ counts.
Osakunor et al, 2020 ³⁰	N=113 preschoolers, 1–5 years, 3.7 \pm 1.1 years.	Cross-sectional study of 49.6% females, made up of children from a rural pre-school, Shotgun metagenomics sequencing.	Structure and diversity of the gut microbiome (bacteria, fungi) and resistome.	Infection with <i>S. haematobium</i> associated with alterations in the gut microbial but not with AMR gene abundance and diversity.	The most abundant bacteria phyla in decreasing order were <i>Bacteroidota</i> (previously <i>Bacteroidetes</i>), <i>Faecalibacterium</i> , <i>Clostridium</i> , and <i>Proteobacteria</i>
Kay et al, 2015 ³¹	N=139 children aged six months to 13 years old.	Cohort study, of children (50.9% female), from rural area with a high burden of <i>S. haematobium</i> (prevalence=27% in study participants), 16S rRNA (V3-V4 hypervariable region) gene sequencing.	Diversity and abundance of different gut bacteria by age, sex, infection with helminths parasites causing schistosomiasis and exposure to anti-helminthic praziquantel.	Some OTUs including <i>Bacteroides</i> and <i>Helicobacter</i> were more abundant in children ≤ 1 year old compared to older children.	Pre- and post-treatment the most abundant phyla were <i>Bacteroidetes</i> , <i>Firmicutes</i> , and <i>Proteobacteria</i> . Gut microbiome differed in schistosome-infected vs uninfected children.
Pfavayi et al, 2021 ³²	N=116 preschoolers (≤ 5 years old).	Cross-sectional study of rural-based children, 50.9% females, Shotgun metagenomics sequencing.	Gut mycobiome and related it to fungal sensitization and seroreactivity.	Mycobiome was <1% of the gut microbiome. <i>Protomyces</i> was most abundant.	<i>Schistosoma haematobium</i> had an effect on fungal genera. All produced Abs to the fungi.

(Continued)

Table I (Continued).

Author, Year, Study Name, if Applicable	Participants N, Median Age (IQR) or Mean (SD) or Range	Study Design, % Female, and Participants and Site Description, Gene Profiling Method(s) and Other Lab Tests	Reported	General Findings	Specific Findings and Shortcomings
Katsidzira et al, 2019 ³³	N=20 adults, >50 years of age.	Cross-sectional study, 50% females, equal numbers of urban and rural adult participants, 16S rRNA whole gene sequencing.	Microbiota of urban and rural Zimbabweans, with focus on colorectal cancer.	There were differences in the relative abundance of a few genera.	No difference in the abundance of <i>Fusobacteria</i> , which has been associated with tumor formation in the colon. Some differences in <i>Streptococcus bovis</i> and <i>Oscillospira guilliermondii</i> between urban and rural participants.
Bourke et al, 2019, The ARROW Trial Sub-study ³⁴	N=140 stool samples from children, age unspecified, from Zimbabwe.	Cross-sectional sub-study in a trial, Children randomized to continue versus stop cotrimoxazole, Shotgun metagenomics sequencing for 20 VGS-positive samples.	Association of cotrimoxazole and the gut microbiome and biomarkers of intestinal inflammation.	Global microbiome composition was no different, but viridans group <i>Streptococci</i> (VGS) were lower among children continuing cotrimoxazole.	20 samples were positive for at least one of the VGS. Six samples were from children continuing and 14 were from children stopping cotrimoxazole. Individual VGS were present less often from children continuing cotrimoxazole.
Gough et al (2020), The ARROW Trial Sub-study ³⁵	N=20 children, age unspecified, from Zimbabwe.	Cross-sectional sub-study in a trial. Stool from six children who continued and fourteen from children who stopped cotrimoxazole and were positive for VGS.		<i>S. salivarius</i> strains in children living with HIV who are on ART who continued cotrimoxazole use are no different from those who stopped.	<i>S. salivarius</i> strains are different from those of children from high-income countries.
Duri et al ³⁶	N=1200 pregnant women (50% living with HIV) and their babies	Protocol describing an observational cohort. Maternal urine, stool, plasma, cord blood, amniotic fluid, placenta and milk plus infant plasma, dried blood spot, and stool collected at enrolment and follow-up visits.	An assessment of the maternal and infant gut microbiome and other factors of HIV-exposed, uninfected infants' adverse outcomes.	Some results are described in paper.	
Chandiwana et al, 2023, UZ-CHS Birth Cohort Sub-study ³⁷	N=94 Mothers, Median age = 28 years (22.3–32.0)	Cross-sectional sub-study in a cohort, Participants are 100% female, ie pregnant women, from urban area, 16S rRNA gene sequencing (segments spanning V5 and V6 regions).	Association of microbiota taxa with HIV infection and disease progression	HIV associated with lower α - and β -diversity.	HIV associated with lower abundance of <i>Bacteroides</i> and <i>Bifidobacterium</i> .

some initial findings.³⁸ Water, sanitation, and hygiene (WASH) interventions included reduced soil ingestion, improved sanitation, water treatment, handwashing, and hygienic preparation of food.^{24,38,39} The infant and young child feeding (IYCF) interventions included 20 g of a small-quantity lipid-based nutrient supplement per day from 6 to 18 months in

addition to counseling that was targeted on key barriers to optimal infant feeding.²⁴ The IYCF interventions increased the mean length-for-age Z-score and the mean hemoglobin concentration by 2.0 g/L (1.3–2.8), reduced stunting and anemia, and improved ponderal growth compared with the non-IYCF interventions. The WASH intervention had no effect on these outcomes. Neither intervention reduced child diarrhea or mortality.²⁴ Subsequent research papers on the infant and maternal microbiota from this cohort are reported below.^{25–27}

In 2021, Gough et al discuss evidence for a persistent public health challenge of preterm birth and low-birth-weight infants in LMICs.²⁵ To identify more effective targets for intervention, the researchers performed shotgun metagenomics sequencing of maternal fecal specimens collected during pregnancy and at 1 month post-partum, from the SHINE Trial. The authors reported that pregnancy gut microbiome taxa accurately predicted birth weight and weight-for-age Z-scores at 1 month. The authors reported remarkably few *Bacteroidetes* species and that this may reflect dietary or functional microbiome differences. The authors did not discuss specimen storage conditions as a possible factor, but it could be theorized that *Bacteroidetes* is highly labile and susceptible to long-term storage.³⁹

Taxa that are associated with resistant starch degradation, specifically members of the *Ruminococcaceae*, *Lachnospiraceae*, and *Eubacteriaceae* families, were important predictors of birth weight and length-for-age Z-scores.²⁵ The authors conclude that pregnancy gut microbiome in rural Zimbabwe is characterized by resistant starch degraders, and that these may be an important metabolic target to improve birth weight and neonatal growth (Box 1). There was no mention of *Bifidobacterium* in the gut microbiome of the studied women,²⁵ which is unusual as *B. adolescentis* is one of the major starch degraders.⁴⁰ According to the authors, the prediction accuracy for all outcomes was improved when functional enzyme relative abundances were used, demonstrating the potential value of whole-metagenome shotgun sequencing for investigating the human microbiome and health.²⁵

In the same year (2021), Robertson et al published a manuscript from observations also made in the SHINE Trial.²⁶ The authors first discussed the rotavirus as the leading cause of diarrheal morbidity and mortality in children and that oral rotavirus vaccine (RVV) immunogenicity is considerably lower in low- versus high-income populations. They hypothesized that gut microbiota may have a role because of its established contribution to the maturation of early-life immune function. To verify this, the researchers performed shotgun metagenomics sequencing on stool samples and measured antirotavirus immunoglobulin A in corresponding plasma samples, on a subset of infants. The authors reported presence of diverse *Bifidobacterium*, dominated by *B. longum*, followed by *B. bifidum* and *B. pseudocatenulatum*. Among the 158 infants with stool samples and anti-rotavirus IgA titers, the authors reported that only 34 (22%) were RVV seroconverters. The authors concluded that further research is warranted to examine the mechanistic role of the gut microbiome in poor oral RVV efficacy in LMICs. Besides the “to-be-expected” presence of *B. pseudocatenulatum* in the infants, they reported a potential association of *Bacteroides thetaiotaomicron* with vaccine responsiveness, as it was the only species associated with anti-rotavirus IgA titer.²⁶ The authors debated their findings which failed to find a clear infant microbiome signature that distinguished RVV seroconverters from non-seroconverters, despite using gold-standard metagenome sequencing. This suggests a need for further studies using a larger cohort and long-read metagenomics methods such as MinIon[®] nanopore sequencing (Box 2).

In 2023, the SHINE Trial Team (first author Robertson RC) published a manuscript, which started by describing traditional determinants of stunting and child undernutrition, then proposed some yet unestablished determinants.²⁷ The authors then described their study of early-life fecal microbiome composition and function, in which stool samples were collected longitudinally from infants. After shotgun metagenomics sequencing, the authors found that *B. longum* was the

Box 1 Question About Potential Benefit of Starch Degrading Microbiome

Could resistant starch degraders be an important metabolic target to improve birth weight and neonatal growth?

Box 2 Long-Range Sequencing Methods

Could long-read sequencing help to distinguish infant microbiome signatures of RVV seroconverters and non-seroconverters?

predominant species at all time-points up to 12 months of age. Four other *Bifidobacterium* species (*B. breve*, *B. bifidum*, *B. pseudocatenulatum*, and *B. kashiwanohense*) were consistently among the most abundant species at the earlier time points before the children attained 12 and 18 months of age, before being outnumbered by different gut bacteria. The study also showed that children who were HIV-exposed but uninfected (HEU) exhibited reduced abundance of *Bifidobacterium* species compared to the HIV-unexposed. The authors proposed that this may partially explain deficits in growth in the HEU childhood and interventions may complement efforts to combat child undernutrition.²⁷ Of interest is the presence of *B. pseudocatenulatum* in the infants, associated with bottle (formula)-fed infants, which needs exploring (Box 3). Formula feeding is rare in Zimbabwe due to Ministry of Health and Child Care campaigns and the Zimbabwe Public Health (Breast-milk Substitutes and Infant Nutrition) Regulations, 1998, restricting the advertising and sales promotion of breast milk substitutes to the public.

According to a 2022 Child Health, Agriculture and Integrated Nutrition (CHAIN) protocol, the authors redesigned infant and young child feeding (IYCF) interventions using locally available foods to improve intake, uptake, and utilization of nutrients.²⁸ The objectives of the trial are to improve infant growth, and interrogate underlying pathogenic pathways during the complementary feeding period with the aim of reducing global burden of stunting. Results of this study are not yet published.

A 2020 paper by Flygel et al²⁹ compared the composition of gut microbiota of children and adolescents living with perinatal transmitted HIV, taking antiretroviral therapy, in the Bronchopulmonary function in REsponse to Azithromycin Treatment for chronic lung disease in HIV-infected children (BREATHE) Study, with that of children and adolescents without HIV. Rectal swabs were collected from 177 participants living with HIV and 103 controls living without HIV. Gut microbial composition was explored using 16S rRNA gene (V4 hypervariable region) amplicon sequencing. The authors reported enriched levels of *Corynebacterium*, *Fingoldia*, and *Anaerococcus* in participants living with HIV, and enrichment of *Enterobacteriaceae* in participants with low CD4⁺ counts (<400 cells/mm³). Prolonged ART-treatment (≥10 years) was significantly associated with a richer gut microbiota by alpha diversity.²⁹ They concluded that children living with HIV have altered gut microbiota and that prolonged ART may restore the richness of the microbiota closer to that of children and adolescents without HIV. This result may be crucial for high HIV-burdened settings like Zimbabwe, though more information on reasons for differences and presence of potential probiotic microbes and potential pathogens in the birth canal, breast milk, gut, and skin of mothers of HIV-exposed infants needs to be explored (Box 4).

In 2020, Osakunor et al focused on preschool-aged infants living in a rural community endemic with bilharzia.³⁰ Bilharzia, also known as schistosomiasis, is caused by helminth parasitic worms and has been shown to have systemic effects within the host. The authors first discussed established factors associated with the composition of the gut microbiome, then deliberated about some yet-to-be-determined environmental factors such as protozoa and helminth parasites. In their study, the researchers tested the hypothesis that infection with the human helminth parasite, *Schistosoma haematobium*, is associated with changes in gut microbial and antimicrobial resistance gene abundance or diversity. They used shotgun metagenomics sequencing to characterize the gut microbiome and the resistome of Zimbabwean preschool infants aged 1 to 5 years, in which the *S. haematobium* infection prevalence was 15.9%. The most abundant bacteria phyla in decreasing order were Bacteroidota (previously Bacteroidetes), Bacillota (previously Firmicutes), and Pseudomonadota (previously Proteobacteria).³⁰ This study was one of the first to describe

Box 3 *Bifidobacterium pseudocatenulatum* and Formula Milk in Infants from Zimbabwe

What could be the explanation for the presence of *Bifidobacterium pseudocatenulatum* in the exclusively breastfed infants, which is usually associated with bottle (formula)-fed infants?

Box 4 Microbiota Differences in HIV-Exposed Children

Are microbiota differences observed in HIV-exposed children due to altered maternal microbiota, transformed human milk oligosaccharides from the mothers, prophylactic antiretroviral drugs OR prophylactic antibiotics, which may influence the seeding and succession of the infants' gut microbiome?

a metagenomics dataset of Zimbabwean preschool-aged infants and indicates an association between urogenital schistosome infection and abundance of Pseudomonadota in the gut microbiome. They concluded that the gut microbiome but not the resistome is associated with urogenital schistosomiasis in preschool-aged children. The study also provided insight into microbial composition and function of preschool infants, though more information about bacterial strain level diversity is required.

Kay et al carried out a study to characterize the gut microbiome of children also exposed to the helminth *S. haematobium*, then pre- and post-treatment with the drug praziquantel.³¹ Stool DNA was extracted at baseline and 12 weeks following anti-helminthic treatment with praziquantel (PZQ), as appropriate. The 16S rRNA gene amplicon (V3-V4 hypervariable region) sequencing included baseline and post-treatment samples, with profiles clustered into operational taxonomic units (OTUs). Pre-treatment, the most abundant phyla were Bacteroidota, followed by Bacillota and Pseudomonadota, respectively. The relative abundance of taxa among bacterial classes and communities showed limited variation by age group or sex. Some OTUs including *Veillonella*, *Streptococcus*, *Bacteroides*, and *Helicobacter* were more abundant in children ≤ 1 year old compared to older children. Furthermore, the gut microbiome differed in schistosome-infected vs uninfected children, with 27 OTUs occurring in infected but not uninfected children. PZQ treatment did not appear to alter microbiome structure in infected or uninfected children from that observed at baseline.³¹ The data suggest changes in the gut microbiota with age, which agrees with reports from other settings.

Pfavayi et al studied the association between gut microbiome and fungal allergic sensitization in young rural Zimbabwean children.³² They carried out the research to interrogate increased prevalence of allergic diseases over the last few decades. The gut microbiome of stool samples was characterized using shotgun metagenomics sequencing, and sensitization to common fungi was assayed using skin prick tests. The mycobiome formed $<1\%$ of the sequenced gut microbiome, and 228 fungal genera were identified. According to the authors, the most abundant fungal genera detected were *Protomyces*, *Taphrina*, and *Aspergillus*, and individuals were frequently sensitized to *Saccharomyces cerevisiae*. The paper concluded that further studies with larger populations are required to understand the role of the microbiome in allergic diseases.³²

Katsidzira et al explored gut microbiome signatures associated with colorectal cancer (CRC) in 10 male and 10 female adults from rural and urban Zimbabwe.³³ In their hypothesis, they propose that the rise in CRC incidence in Zimbabwe is related to shifts in the colonic microbiota and metabolome. Therefore, the researchers conducted an exploratory study comparing diet and fecal markers associated with CRC risk in the 20 apparently healthy adults, all aged >50 years. The full-length 16S rRNA gene was sequenced. The researchers reported *minor* differences in fecal microbiota composition between urban and rural participants. In particular, there were significant differences in the relative abundances of a few genera, ie *Blautia obeum*, *Streptococcus bovis*, and *Subdoligranulum* were more abundant among urban residents. *Oscillospira guilliermondii*, *Sporobacter termitidis*, and *Clostridia* were more abundant among rural participants. The study detected no difference in the abundance of *Fusobacteria* in feces, which has been associated with tumor formation in the colon. In addition, there was no difference in the abundance of *Bilophila wadsworthia*, a bacterium associated with experimental colitis. The authors concluded that the higher abundance of *Subdoligranulum* in the urban participants highlights the potentially beneficial impact of retained fibre intake in the urban study population.³³

In a 2020 publication, Gough et al described results of strain-level analysis of gut-resident pro-inflammatory viridans group Streptococci (VGS) for infants in the Anti-Retrovirus Research for Watoto (ARROW) Trial.³⁴ Six stool samples from children who continued and fourteen from children who stopped cotrimoxazole were positive for VGS, in an earlier study. The authors further explored strain-level patterns within the VGS identified as being suppressed by long-term cotrimoxazole prophylaxis, *Streptococcus salivarius* being the most prevalent. They used a strain-level metagenomic profiling tool – The Pangenome-based Phylogenomic Analysis (PanPhlAn) – to further characterize *S. salivarius* sub-species variants. They found that *S. salivarius* strains in HIV-positive children on ART who continued cotrimoxazole use are no different from those of children who stopped, but that the *S. salivarius* strains are different from those of children from high-income countries.³⁴

According to Bourke et al, in a publication from 2019, stunting in Zimbabwean infants is characterized by low-grade chronic inflammation.³⁵ The study therefore focused on cotrimoxazole, an antibiotic that reduces systemic inflammation in children living with HIV. The research used shotgun metagenomics sequencing of total fecal DNA from children

Box 5 Identity of Potential Microbes

As maternal HIV is associated with a lower abundance of *Bacteroides* and *Bifidobacterium*, are these two viable nutritional probiotics, in pregnancy?

randomized to continue versus stop cotrimoxazole, in the ARROW Trial. Randomized groups did not differ in species-level diversity. Bacterial community composition was also similar between groups. However, 7 bacterial species (*Alistipes onderdonkii*, *Eggerthella lenta*, *Clostridium bartlettii*, *Haemophilus parainfluenzae*, *Streptococcus mutans*, *Streptococcus parasanguinis*, and *Streptococcus vestibularis*) and 11 protein families mapping to *Streptococcus parasanguinis*, *Streptococcus salivarius*, and *Haemophilus parainfluenzae* were consistently less abundant in the continue versus stop group. The relative abundance of Enterobacteriaceae, which includes gastrointestinal pathogens (eg *Salmonella*, *E. coli*, and *Shigella*) that are frequently resistant to cotrimoxazole, was not affected by cotrimoxazole and was increased in those continuing versus stopping cotrimoxazole at week 96.³⁵

Duri et al published the protocol of The University of Zimbabwe College of Health Sciences (UZ-CHS) Birth Cohort study in 2020.³⁶ This is a prospective observational cohort study comparing infants born to 1200 mothers living with and without HIV, attending primary health care clinics in Harare's high-density suburbs. Half of the recruited pregnant women were living with HIV. This is a prospective observational cohort study comparing infants born to women living with HIV and those without. The pregnant women were recruited at ≥ 20 weeks of gestation between February 2016 and June 2019. Participants are being followed up as mother–infant pairs at birth, within 10 days of birth and at 6, 10, 14, 24, 48, 72, and 96 weeks post-partum. Questionnaires are administered and bio-samples for laboratory tests are acquired at each visit. Biomaterials are for specific tests including gut microbiome profiling, as well as tests for current infections, clinical biochemistry, full blood counts, plus immune status. One paper has been published on maternal microbiome profiles.³⁷

The paper from the sub-study of mothers from the University of Zimbabwe (UZ) Birth Cohort,³⁶ mentioned above, was published by Chandiwana et al in 2023.³⁷ It assessed the composition of the intestinal microbiota in pregnant women residing in Zimbabwe.³⁷ Pregnant women living with HIV (PWLWH) had a lower species diversity (α -diversity) than those without. The differences in microbiome composition between samples (ie β -diversity) also differed between them. High abundance of Spirochaetaceae, Veillonellaceae, and *Treponema* in the third trimester of pregnancy was associated with low birth weight in the infants. This study focused on beneficial bacteria in the pregnant women and reported that living with HIV was associated with a lower abundance of *Bacteroides* and *Bifidobacterium* (Box 5), and a higher abundance of *Actinomyces* and *Succinivibrio*.³⁷

Results and Analysis

We searched the literature for publications about the gut microbiota and *Bifidobacterium*. We found very few publications on this and expanded our search to include papers describing components of a healthy or an unhealthy gut microbiome of neonates, infants, and adults from Zimbabwe. Some of the published papers explored evolution of the gut microbiota in response to diet, infection (eg bilharzia and HIV), and environmental exposures (such as antibiotics and antiretroviral drugs). Some explored the contribution of the gut microbiota and attempted to unravel the implications of a perturbed microbiome to mostly infant health and disease. There were no reports on the fortification of foods with specific *Bifidobacterium* strains and human milk oligosaccharides in the Zimbabwe context.

Most publications leveraged randomized clinical trials, eg the SHINE Trial and ARROW Trial, and international collaborations. The microbiome studies were carried out in collaboration with researchers from Europe (UK and Switzerland), South Africa, and USA. The laboratory methods applied to the microbiome work included Illumina shotgun metagenome sequencing and 16S rRNA gene sequencing (mainly sequencing of the V3 hypervariable region).

Discussions and Conclusions

There are commendable efforts to characterize the gut microbiome by individual researchers and groups, leveraging on-going mother–infant cohorts in Zimbabwe. Most of the assays were carried out in laboratories outside of Zimbabwe, mainly due to technological and skills gaps. Efforts to include local research institutions in microbiome research are likely to receive local and global support because issues of diversity, equity, and inclusion have risen to the top of the list of critical concerns in

science and medicine. All of the published microbiome studies focused on short-read sequencing. There may be useful insights to be gained from long-read sequencing, eg by using Nanopore sequencing technologies. In the Zimbabwe context MinIon[®] Nanopore sequencers would be most attractive and relevant due to their low cost and small size.^{41,42}

Based on the few numbers of studies and participants, there are many important microbiome-focused questions that still need to be answered. For example, studies of strain level diversity of *Bifidobacterium* communities and their role in immune development and vaccine response ought to be carried out, in Zimbabwe and other LMIC. In order to benefit and leverage advances made in other settings, Zimbabwean researchers must carry out local cohort studies that compare findings to other LMIC and high-income country groups. Such cohorts are valuable when designing mechanistic studies and downstream translational work such as next-generation probiotics, and prebiotics. The dynamics of *Bifidobacterium* during early life, and across the lifespan, needs to be explored in experimental studies and clinical trials. Culturing of microbes and sequencing of isolates are important for identifying local strains for therapy development. This is relevant because strategies to counteract malnutrition, enhance vaccine response, and improve health are of significant public health interest.^{1,8,36,43} Other pertinent questions that remain unanswered are placed in boxes throughout this scoping review.

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