Association of Secretor Status and Recent Norovirus Infection With Gut Microbiome Diversity Metrics in a Veterans Affairs Population

Jordan A Johnson, Emory University
Timothy Read, Emory University
Robert A, III Petit, Emory University
Vincent Marconi, Emory University
Kathryn L Meagley, Emory University
Maria C Rodriguez-Barradas, Michael E DeBakey VA Med Ctr
David O Beenhouwer, Veterans Affairs Greater Los Angeles Health System
Sheldon T Brown, James J. Peters Veterans Affairs Medical Center
Mark Holodniy, Veterans Affairs Palo Alto Health Care System
Cynthia A Lucero-Obusan, Veterans Affairs Palo Alto Health Care System

Only first 10 authors above; see publication for full author list.

Journal Title: OPEN FORUM INFECTIOUS DISEASES
Volume: Volume 9, Number 5
Publisher: OXFORD UNIV PRESS INC | 2022-05-01, Pages ofac125-ofac125
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1093/ofid/ofac125
Permanent URL: https://pid.emory.edu/ark:/25593/vw89q

Final published version: http://dx.doi.org/10.1093/ofid/ofac125

Copyright information:

© The Author(s) 2022. Published by Oxford University Press on behalf of Infectious Diseases Society of America.

This is an Open Access work distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License (https://creativecommons.org/licenses/by-nc-nd/4.0/rdf).
Association of Secretor Status and Recent Norovirus Infection With Gut Microbiome Diversity Metrics in a Veterans Affairs Population


1Department of Epidemiology, Emory University Rollins School of Public Health, Division of Infectious Diseases, Department of Medicine, Emory University School of Medicine, and Atlanta Veterans Affairs Medical Center, Atlanta, Georgia, USA; 2Infectious Diseases Section, Michael E. DeBakey Veterans Affairs Medical Center, Houston, Texas, USA; 3Department of Medicine, Baylor College of Medicine, Houston, Texas, USA; 4Veterans Affairs Greater Los Angeles Health System, Los Angeles, California, USA; 5James J. Peters Veterans Affairs Medical Center, Bronx, New York, USA; 6Veterans Affairs Palo Alto Health Care System, Palo Alto, California, USA; 7Division of Infection Control and Public Health, Public Health Surveillance and Research, Veterans Health Administration, Washington, DC, USA; 8Division of Infectious Diseases & Geographic Medicine, Stanford University, Stanford, California, USA; 9Department of Pathology and Laboratory Medicine, Emory University School of Medicine, Atlanta, Georgia, USA; 10Department of Molecular Virology and Microbiology, Baylor College of Medicine, Houston, Texas, USA; and 11Division of Viral Diseases, National Center for Immunization and Respiratory Diseases, Centers for Disease Control and Prevention, Atlanta, Georgia, USA

Norovirus infection causing acute gastroenteritis could lead to adverse effects on the gut microbiome. We assessed the association of microbiome diversity with norovirus infection and secretor status in patients from Veterans Affairs medical centers. Alpha diversity metrics were lower among patients with acute gastroenteritis but were similar for other comparisons.

Keywords. norovirus; microbiome; secretor status; acute gastroenteritis.

The human digestive tract is populated by a diverse array of bacterial species, though these are highly variable, differing with diet, geography, and genetics [1–4]. Dysbiosis in the gut microbiome have been associated with a wide range of chronic and infectious diseases [3, 5, 6].

Norovirus causes 19–21 million cases of acute gastroenteritis (AGE) in the United States annually [7], with persons >65 years of age at greatest risk for norovirus-associated deaths [8]. Norovirus infection can cause diarrhea and vomiting, disrupting the intestinal environment and reducing gut microbiome diversity [9, 10]. Additionally, the microbiome may offer protection against symptoms of norovirus infection [11]. Further examination of the association between norovirus and the microbiome could explain differences between symptomatic and asymptomatically infected individuals.

Host genetics may also affect microbiome diversity [4]. Individuals with a functional fucosyltransferase-2 (FUT2) gene have histo-blood group antigens (HBGAs) present in the mucosa of the gut and are known as “secretors” [12]. Gut microbiome diversity and composition may differ between secretors and nonsecretors [13–15] or may not [16, 17]. Susceptibility to norovirus infection is strongly associated with secretor status, with nonsecretors being resistant to some norovirus genotypes [12, 18, 19]. Noroviruses use HBGAs as attachment factors for cellular entry, though the exact mechanism is unknown [20, 21].

To date, few studies have examined the complex relationships between secretor status, norovirus infection, and the microbiome. Due to the increased risk of norovirus-associated morbidity and mortality in the older adult population and adverse effects of potential microbiome disruptions, further exploration of these relationships in this population is needed.

The ongoing SURveillance Platform for Enteric and Respiratory Infectious Organisms in the VA (SUPERNOVA) is a collaboration between Emory University, the Centers for Disease Control and Prevention, and Department of Veterans Affairs Medical Centers (VAMCs) in Atlanta, Georgia, Houston, Texas, Bronx, New York, Los Angeles, California, and Palo Alto, California. We analyzed the microbiomes of 100 SUPERNOVA patients to assess differences in microbiome diversity by recent norovirus infection and secretor status. Additional analyses explored diversity by recent AGE symptoms and microbiome composition between norovirus and secretor status groups.

METHODS

Recruitment into the SUPERNOVA study has been described elsewhere [22]. In brief, individuals were enrolled as cases in the SUPERNOVA study if they met the definition for AGE (Supplementary Methods) when presenting as outpatients or when hospitalized at participating VAMCs (Atlanta, Houston, Bronx, and Los Angeles). Controls without AGE presented to the same VAMC with an admission or outpatient visit date within a week of a matched case.

Patients provided stool and saliva samples within 10 days of symptom onset (cases) or enrollment (controls). All stool specimens were tested using the BioFire FilmArray Gastrointestinal...
Supplementary Table 4. However, norovirus-positive patients had a smaller age range (48/51; 94%), and 12 (24%) had other AGE pathogens detected (Supplementary Table 1). The most common capsid genotype among norovirus-positive patients was GII.4 (20/51; 39%), though not all norovirus-positive samples were genotyped.

Among norovirus-negative patients with AGE, 16/23 (70%) did not have pathogens detected on BioFire. Other detected pathogens included C. difficile (2 patients), Rotavirus, Shigella, multiple pathogens (2 patients), and 1 inconclusive result. Three norovirus-negative patients without AGE had pathogens detected, including C. difficile, enteropathogenic E. coli, and multiple pathogens (1 patient each).

Recent AGE Illness Associated With Reduced Microbiome Diversity
Across all samples, secretors had a lower mean richness compared with nonsecretors, with a mean number of detected ASVs of 224.6 among secretors and 247.7 among nonsecretors (Figure 1; Supplementary Table 2). Mean richness was similar between norovirus-positive secretors and norovirus-positive nonsecretors, with 228.1 and 223.3 ASVs, respectively. Shannon indices were similar in all comparisons (Supplementary Table 3).

Overall, richness and Shannon indices were similar between patients with and without norovirus (Figure 1; Supplementary Tables 2 and 3). However, norovirus-positive patients had a lower richness when compared with norovirus-negative patients without AGE (227.5 and 268.5, respectively), but higher when compared with norovirus-negative patients with non-norovirus AGE (227.5 and 200.3). Shannon diversity indices had a similar pattern.

Patients with AGE had substantially lower mean richness (216.2) and a lower Shannon index (3.773) than patients without AGE (270.2 and 4.096, respectively) (Figure 1; Supplementary Tables 2 and 3). This held true when restricting to norovirus-negative individuals.

No Differences in Microbiome Composition Seen by Any Variables of Interest
There were no differences found in microbiome composition between secretors and nonsecretors via ordination using the most abundant genera, including when restricting to individuals without AGE (Figure 2A and B). Points representing secretors were slightly more graphically dispersed than nonsecretors, but there was substantial overlap in the 2 groups. Norovirus-positive patients had similar microbiome compositions to norovirus-negative patients on ordination plots (Figure 2C). Further, overall compositions were similar when comparing norovirus AGE with non-norovirus AGE (Figure 2D). Patients with AGE appeared similarly in microbiome composition to patients without AGE, including when restricting to only norovirus-negative patients (Figure 2E and F).

Individual LDMs for secretor status, norovirus infection, and recent AGE did not show significant associations with microbiome composition (Supplementary Table 4). In the
full model with all variables and controlling for age and prescription drug use, no significant associations were detected (Supplementary Table 4).

**DISCUSSION**

Secretor status and norovirus infection were not independently associated with differences in either microbiome richness or Shannon diversity index. Additionally, no differences by secretor status or norovirus infection in microbiome composition were seen on NMDS ordination or LDM regression. No differences were detected despite a larger sample size compared with some similar studies [13–15, 17].

Due to the disruption of the gut environment during AGE [32], we expected to see lowered microbiome diversity for patients with AGE; this was true for both richness and Shannon index. However, no difference was seen in microbiome composition between AGE groups upon ordination. As composition analyses only included the most abundant genera, the reduction in diversity metrics may be borne by only less abundant genera.

Our secretor status results support previous findings of no differences in microbiome composition between secretor status groups [16, 17]. Though previous studies found a relationship between secretor status, norovirus IgA titer (past norovirus infection), and microbiome composition, we did not see relationships between analogous variables in our data with a larger sample size examining recent infections [17]. Additional work incorporating pre- and postinfection stool samples, such as challenge studies [11], could better explore microbiome changes following infection.

Our study had at least 4 limitations, including lack of serial pre-infection stools, a limited sample size, potential nonrepresentativeness of the VA population compared with the general US population, and additional factors (e.g., unreported medication, dietary differences, norovirus genotype, or co-infecting pathogens) that were not explored here.

In summary, we did not detect an association between secretor status or norovirus infection and microbiome diversity or composition. Though findings suggest that there might not be a...
relationship between norovirus infection and the microbiome, longitudinal studies would be better suited to capture any causal effect of secretor status and norovirus infection on the microbiome.

Acknowledgments
We sincerely thank the collaborators and study teams at our SUPERNOVA sites for their continued time and efforts that made this study possible.

Financial support. This work was supported by the Foundation for Atlanta Veterans Education and Outreach (FAVER; formerly the Atlanta Research and Education Foundation or AREF) and the Centers for Disease Control and Prevention, Atlanta, Georgia.

Potential conflicts of interest. The authors have no conflicts of interest to disclose. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

Patient consent. This study does not include factors necessitating patient consent.

Figure 2. Non-metric multidimensional scaling ordination shows no difference in microbiome composition by secretor status and norovirus infection, with possible greater differences in composition among patients with AGE. Ordination using non-metric multidimensional scaling based on Bray-Curtis distances was used to compare overall microbiome composition between secretors and non-secretors (A and B), norovirus-positive patients and norovirus-negative patients (C and D), as well as patients with AGE and those without AGE (E and F). Included populations by panel: A, all individuals, with secretor points appearing slightly more dispersed. B Only patients without AGE, where microbiome compositions appear similar across both secretor status groups. C, all individuals, with points for norovirus-positive patients appearing slightly more dispersed. D, only patients with AGE, where points for norovirus-positive patients also appeared more dispersed than norovirus-negative patients. E, all individuals, with points for AGE-free individuals more clustered together compared to points for patients with AGE. F, only patients without norovirus and there is little apparent difference between the two groups. Ellipses in all panels are based on multivariate t-distribution. One individual (square point, norovirus-negative and AGE-free) with an inconclusive secretor status is shown on secretor status plots.

Disclaimer. The conclusions, findings, and opinions expressed by authors contributing to this journal do not necessarily reflect the official position of the US Department of Health and Human Services, the Public Health Service, the Centers for Disease Control and Prevention, or the authors’ affiliated institutions. Use of trade names is for identification only and does not imply endorsement by the Public Health Service or the US Department of Health and Human Services.

Data availability. Sequence data are available at the NBI SRA database under BioProject accession PRJNA741048.

References


