Microbiome and Antimicrobial Resistance Gene Dynamics in International Travelers

Charles Langelier, Michael Graves,¹
Katrina Kalantar,¹ Saharai Caldera,
Robert Durrant, Mark Fisher, Richard Backman,
Windy Tanner, Joseph L. DeRisi, Daniel T. Leung

We used metagenomic next-generation sequencing to longitudinally assess the gut microbiota and antimicrobial resistomes of international travelers to clarify global exchange of resistant organisms. Travel resulted in an increase in antimicrobial resistance genes and a greater proportion of *Escherichia* species within gut microbial communities without impacting diversity.

International travel is a known contributor to the emergence of organisms with antimicrobial resistance (AMR) (1-4). Colonization with resistant microbes acquired during travel can persist asymptomatically for extended periods and result in transmission into the environment and susceptible populations (5). The mechanisms underlying acquisition of AMR bacteria during travel are incompletely understood, although changes in the intestinal microbiota are hypothesized to play a role (6). To clarify AMR exchange during global travel, we used metagenomic nextgeneration sequencing (mNGS) to assess gut microbiota composition and the antimicrobial resistome.

The Study

During March 2016–2018, we recruited adults with planned travel to Asia or Africa for healthcare-related work. Participants introduced 1 tablespoon of stool into vials with either RNAprotect (QIAGEN, http://www.qiagen.com) or Cary-Blair (CB) media and then submitted samples and surveys pretravel, posttravel, 30 days posttravel, and 6 months posttravel. Upon receipt, we stored RNAprotect samples at -80°C and CB samples at 4°C until inoculation onto chromogenic agar plates selective for extended-spectrum

Author affiliations: University of California, San Francisco, California, USA (C. Langelier, K. Kalantar, S. Caldera, J.L. DeRisi); Chan Zuckerberg Biohub, San Francisco (C. Langelier, S. Caldera, J.L. DeRisi); University of Utah School of Medicine, Salt Lake City, Utah, USA (M. Graves, R. Durrant, M. Fisher, R. Backman, W. Tanner, D.T. Leung); ARUP Laboratories, Salt Lake City (R. Durrant, M. Fisher)

DOI: https://doi.org/10.3201/eid2507.181492

β-lactamase (ESBL)–producing bacteria (CHROMagar ESBL) and incubation overnight at 37°C. We then inoculated single colonies into LB broth and incubated overnight at 37°C. If multiple morphotypes were identified, we conducted separate subcultures. We performed speciation by using matrix-assisted laser desorption/ionization time-of-flight mass spectrometry.

DNA and RNA extracted using the QIAGEN Powerfecal kit underwent metagenomic sequencing as previously described (7). Raw data are available publicly (Bioproject PRJNA509512). We detected enteric microbiota using a recently developed bioinformatics pipeline (7). We aggregated microbial alignments at the genus level before downstream analyses. To control for background environmental and reagent contaminants, we incorporated no-template water control samples alongside extracted nucleic acid and carried them forward throughout library preparation and sequencing. We then performed direct subtraction of total reads aligning to each microbial genus present in controls from each study sample before downstream analyses.

We used the SRST2 computational tool and AR-Gannot_r2 database (https://github.com/katholt/srst2) to identify AMR genes with allele coverage of >20% (8). Although a precise definition of ESBL has not been established, we used a working definition of Ambler class A–D β -lactamases with known or predicted ability to confer resistance to first- through third-generation cephalosporins (9,10). We required detection of chromosomally encoded Ambler class C β -lactamases (i.e., AmpC) by both DNA-Seq and RNA-Seq to capture actively expressed genes.

Nine of 10 participants were culture-positive for ESBL-producing *Escherichia coli* (ESBL-PE) upon return, including 8 persons who traveled to Nepal and 1 who went to Nigeria. One traveler was found to be colonized before departure (traveler 3 [T3]); 3 travelers had persistent carriage at the 30-day visit (T2, T3, and T5) and 2 at 6 months (T3 and T5) (Table 1). Although 4 participants experienced diarrheal symptoms during travel, only 1 (T5) had persistent diarrheal symptoms at 6 months. Diarrheal symptoms were not associated with persistent ESBL-PE colonization at any point, and no travelers reported antibiotic use or receipt of healthcare in an inpatient setting while traveling. All travelers were exposed to inpatient healthcare facilities, and 3 travelers reported street food consumption.

¹These authors contributed equally to this article.

Table 1. Selected characteristics of 10 travelers and assessment of ESBL-producing Enterobacteraciae*

Traveler	Destination	Duration, d	Diarrhea	Pretravel	Posttravel	30 d posttravel	6 mo posttravel
T1	Nepal	30	N	_	AmpC	OXA-209	OXA-209
T2	Nepal	30	N	_	AmpC, CTX-M1	AmpC	_
T3	Nepal	30	N	OXA-209	AmpC, CTX-M1, OXA-209	OXA-209	OXA-209
T4	Nepal	16	N	_	AmpC	_	NR
T5	Nepal	30	Υ	_	AmpC, SHV-12	AmpC	AmpC, CTX-M-1
T6	Nepal	15	Υ	_	AmpC, CTX-M-1	_	_
T7	Nepal	18	N	_	AmpC, CTX-M-1	NR	NR
T8	Uganda	14	N	_	- (AmpC)†	_	_
T9	Nigeria	60	Υ	_	AmpC	_	_
T10	Nepal	30	Υ	_	AmpC, CTX-M-1	-	NR

*Participants submitted samples pretravel (within 1 week before departure), posttravel (within 1 week after return), 30 days posttravel, and 6 months posttravel. ESBL, extended-spectrum β-lactamase; N, no; NR, no sample received; Y, yes; –, ESBL-negative by culture. †T8 was the only participant who was phenotypically ESBL-negative after travel.

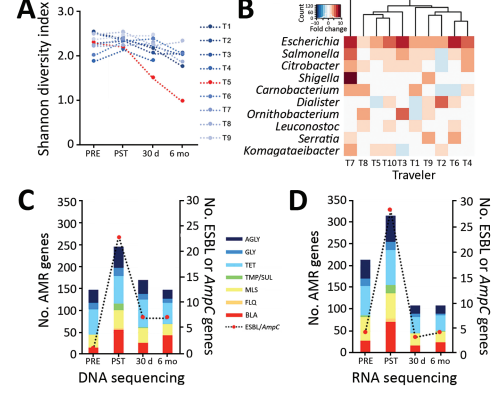
We first examined changes in gut microbiome α diversity after international travel (Figure, panel A) and found that the Shannon diversity index (SDI) did not significantly change upon return (p = 0.674 by Wilcoxon rank-sum test) or at day 30 posttravel (p = 0.250 by Wilcoxon rank-sum test). We then assessed whether microbial community composition differed across all participants at their post-travel versus pretravel visit but found no difference (Bray Curtis Index p = 0.23 by permutational multivariate analysis of variance). Although global composition and diversity of gut microbiota did not significantly change after travel, we observed significant differences in the abundance of

discrete genera. Across all participants, *Enterobacteriaceae* demonstrated the greatest fold change in abundance posttravel; *Escherichia* was the genus most differentially increased (p<0.001 by Wilcoxon rank-sum test) (Figure, panel B; Appendix Figures 1, 2, https://wwwnc.cdc.gov/EID/article/25/7/18-1492-App1.pdf).

Analysis of the antimicrobial resistome revealed an increase in AMR genes and transcripts after return from travel (p<0.01 for DNA and p = 0.03 for RNA sequencing, both by Wilcoxon rank-sum test) (Table 2; Figure, panels C–D; Appendix Figure 3). ESBL-encoding genes, AmpC-encoding genes, or both were identified in 100%

Figure. Microbiome and AMR gene dynamics in international travelers. A) Longitudinal profile of traveler aut microbiome diversity measured by Shannon diversity Index. Traveler 5 (T5) had a Shannon diversity index >3 SDs below the mean when measured at 30 days and 6 months posttravel. B) Microbes, by genus, demonstrating the greatest fold change in abundance after travel on the basis of DNA sequencing nucleotide alignments. C) Total number of AMR genes identified with ≥20% allele coverage by DNA sequencing. D) Total number of AMR genes identified with ≥20% allele coverage by RNA sequencing. AGLY, aminoglycoside; AMR, antimicrobial resistance; BLA, β-lactamase; FLQ, fluoroquinolone; MLS, macrolide, lincosamide,

streptogramin; PRE,



pretravel (within 1 week before departure); PST, posttravel (within 1 week after return); TMP/SUL, trimethoprim/sulfamethoxazole; TET, tetracycline; ESBL, extended-spectrum β -lactamase.

Table 2. Fold change in abundance of AMR genes found in *Enterobacteraciae* with ≥20% allele coverage compared with pretravel, by

resistance	anana	α r	drug	clace*

Toolotanios genie en anagenaes	Fold change compared with pretravel							
	Posttravel		30 d		6 mo			
Resistance gene or drug class	DNA	RNA	DNA	RNA	DNA	RNA		
β-lactam AMR genes								
AmpC	>100	>100	>100	1	>100	2		
AmpH	>100	>100	>100	2	>100	1		
CTX	>100	>100	>100	>100	>100	>100		
MrdA	50	>100	0	>100	5	>100		
OXA	2	1	1	0	1	0		
SHV	>100	>100	>100	>100	>100	>100		
TEM	>100	64	54	0	15	1		
Other antibiotic classes								
Aminoglycoside	2	3	1	0	0	0		
Fluoroquinolone	>100	>100	>100	>100	>100	>100		
Glycopeptide	0	1	0	0	0	0		
Macrolide, lincosamide, streptogramin	2	3	1	3	1	1		
Sulfa	22	29	1	0	0	0		
Tetracycline	1	1	1	0	1	0		
Trimethoprim	>100	88	>100	0	>100	1		

^{*}Among 10 participants who submitted samples pretravel (within 1 week before departure), posttravel (within 1 week after return), 30 days posttravel, and 6 month posttravel. AMR, antimicrobial resistance.

of samples with an ESBL culture-determined phenotype and in 14% of samples without, including 1 participant (T8) who was phenotypically ESBL-negative after travel (Table 1). β -lactam-resistance genes increased posttravel, including AmpC, CTX-M, OXA, and SHV gene families known or predicted to confer ESBL production, as well as diverse additional β - lactamase genes (Appendix Figure 3). Travel also resulted in an increase in qnr plasmid-mediated quinolone-resistance genes, as well as trimethoprim (dfr)-, sulfa-, macrolide-, and aminoglycoside-resistance genes (Table 2). Genes conferring resistance to tetracyclines and aminoglycosides were most abundant in travelers at baseline and remained stable or decreased during travel (Table 2).

We found no significant differences in SDI between persistent carriers and those who lost carriage at 30 days (n = 3; p = 0.56 by t-test) or at 6 months (n = 2; p = 0.27 by t-test) posttravel. T5, who was colonized at both time points and who was the only traveler with persistent diarrheal symptoms, had an SDI >3 SDs below the mean at 6 months (Figure, panel A). Bray-Curtis distance measured pretravel or posttravel did not differ between travelers who were ESBL-PE positive by culture at 30 days or 6 months posttravel (p = 0.32 by permutational multivariate analysis of variance). No individual taxa were associated with posttravel ESBL positivity on the basis of an adjusted p value <0.05 (t-test) at 30 days or 6 months posttravel.

Conclusions

International travel is a well-recognized contributor to the global spread of emerging infectious diseases, including AMR bacteria (1,4). We analyzed the enteric microbiota and resistomes of returned travelers and found a marked increase in AMR genes that was associated with an increased

proportion of *Escherichia* bacteria. At baseline, few participants had evidence of ESBL colonization; however, after travel, ESBL and actively transcribed AmpC genes were notably increased, consistent with previous reports (4,11). Both mNGS and culture-based methods found evidence of persistent ESBL colonization after 6 months, suggesting that travel can induce long-term changes in the antimicrobial resistome (5). In addition to β -lactamase genes, mNGS identified a diversity of other AMR gene classes that increased in abundance after travel. For example, 80% of participants acquired horizontally transferable qnr genes after travel, reflecting the limited utility of quinolones for treatment of traveler's diarrhea (12). Although we did not detect genes known to encode carbapenemases, participants might have harbored carbapenem-resistant Enterobacteraciae, given that a combination of an ESBL or AmpC gene with a porin mutation or efflux pump can lead to carbapenem resistance (13).

Changes in microbiome diversity were not associated with ESBL positivity at 30 days or 6 months posttravel, suggesting that disruption of the antimicrobial resistome can occur in the setting of a preserved microbial community structure. We observed a high rate of ESBL-PE acquisition in this cohort, most of whom traveled to the Indian subcontinent, consistent with previous studies of travelers returning from this region (1). Notably, none of the travelers in this cohort reported antibiotic use, suggesting that substantial ESBL-PE colonization can occur even in the absence of antibiotic-related disruption of commensal gut microbiota.

Because this study is limited by small sample size, relevant associations might have been missed. In addition, we could not assess the presence of carbapenem-resistant *Enterobacteriaceae* by using culture-based methods. mNGS

and phenotypic antimicrobial resistance need assessment in larger cohorts traveling to more destinations. Nonetheless, our findings highlight the pervasiveness of AMR microbe exchange during international travel and the promise of mNGS for assessing the global exchange of antimicrobial resistance.

Acknowledgment

We thank Theresa Sofarelli for inspiration and support for this study.

About the Author

Dr. Langelier is an assistant professor in the Division of Infectious Diseases at the University of California, San Francisco. His research interests involve using metagenomics and transcriptional profiling to investigate host–pathogen interactions and understand the causes of diagnostically challenging diseases.

References

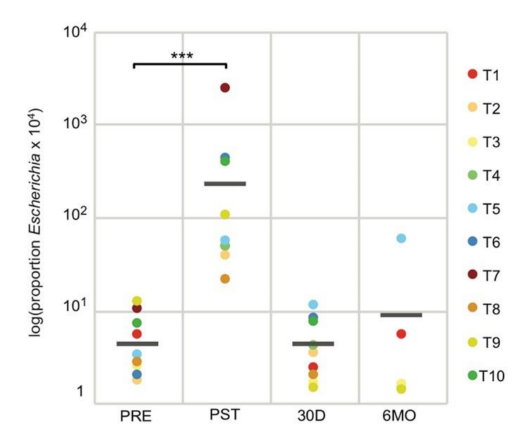
- Arcilla MS, van Hattem JM, Haverkate MR, Bootsma MCJ, van Genderen PJJ, Goorhuis A, et al. Import and spread of extended-spectrum β-lactamase-producing *Enterobacteriaceae* by international travellers (COMBAT study): a prospective, multicentre cohort study. Lancet Infect Dis. 2017;17:78–85. http://dx.doi.org/10.1016/S1473-3099(16)30319-X
- Kuenzli E, Jaeger VK, Frei R, Neumayr A, DeCrom S, Haller S, et al. High colonization rates of extended-spectrum β-lactamase (ESBL)-producing *Escherichia coli* in Swiss travellers to South Asia—a prospective observational multicentre cohort study looking at epidemiology, microbiology and risk factors. BMC Infect Dis. 2014;14:528. http://dx.doi.org/10.1186/1471-2334-14-528
- Hassing RJ, Alsma J, Arcilla MS, van Genderen PJ, Stricker BH, Verbon A. International travel and acquisition of multidrugresistant *Enterobacteriaceae*: a systematic review. Euro Surveill. 2015;20:20.
- von Wintersdorff CJH, Penders J, Stobberingh EE, Oude Lashof AML, Hoebe CJPA, Savelkoul PHM, et al. High rates of antimicrobial

- drug resistance gene acquisition after international travel, the Netherlands. Emerg Infect Dis. 2014;20:649–57. http://dx.doi.org/10.3201/eid2004.131718
- van Duijkeren E, Wielders CCH, Dierikx CM, van Hoek AHAM, Hengeveld P, Veenman C, et al. Long-term carriage of extendedspectrum β-lactamase–producing *Escherichia coli* and *Klebsiella pneumoniae* in the general population in the Netherlands. Clin Infect Dis. 2018;66:1368–76. http://dx.doi.org/10.1093/cid/cix1015
- David LA, Materna AC, Friedman J, Campos-Baptista MI, Blackburn MC, Perrotta A, et al. Host lifestyle affects human microbiota on daily timescales. Genome Biol. 2014;15:R89. http://dx.doi.org/10.1186/gb-2014-15-7-r89
- Langelier C, Kalantar KL, Moazed F, Wilson MR, Crawford ED, Deiss T, et al. Integrating host response and unbiased microbe detection for lower respiratory tract infection diagnosis in critically ill adults. Proc Natl Acad Sci U S A. 2018;115:E12353–62. http://dx.doi.org/10.1073/pnas.1809700115
- Inouye M, Dashnow H, Raven L-A, Schultz MB, Pope BJ, Tomita T, et al. SRST2: Rapid genomic surveillance for public health and hospital microbiology labs. Genome Med. 2014;6:90. http://dx.doi.org/10.1186/s13073-014-0090-6
- Bush K, Jacoby GA. Updated functional classification of β-lactamases. Antimicrob Agents Chemother. 2010;54:969–76. http://dx.doi.org/10.1128/AAC.01009-09
- Giske CG, Sundsfjord AS, Kahlmeter G, Woodford N, Nordmann P, Paterson DL, et al. Redefining extended-spectrum β-lactamases: balancing science and clinical need. J Antimicrob Chemother. 2009;63:1–4. http://dx.doi.org/10.1093/jac/dkn444
- Decker BK, Lau AF, Dekker JP, Spalding CD, Sinaii N, Conlan S, et al. Healthcare personnel intestinal colonization with multidrugresistant organisms. Clin Microbiol Infect. 2018;24:82.e1–4. http://dx.doi.org/10.1016/j.cmi.2017.05.010
- Riddle MS, Connor BA, Beeching NJ, DuPont HL, Hamer DH, Kozarsky P, et al. Guidelines for the prevention and treatment of travelers' diarrhea: a graded expert panel report. J Travel Med. 2017;24(Suppl 1):S57–74. https://doi.org/10.1093/jtm/tax026
- Logan LK, Weinstein RA. The epidemiology of carbapenemresistant *Enterobacteriaceae*: the impact and evolution of a global menace. J Infect Dis. 2017;215(Suppl_1):S28–36.

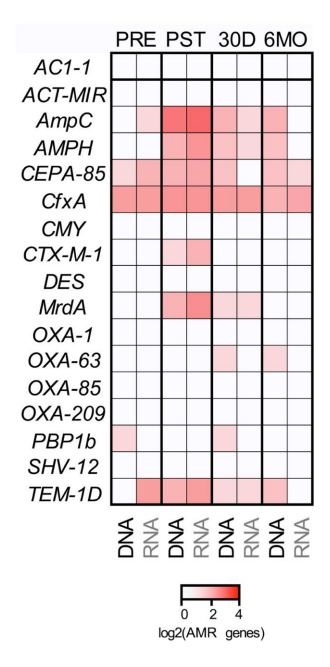
Address for correspondence: Charles Langelier, University of California San Francisco, Infectious Diseases, 513 Parnassus Ave, Room S-380, San Francisco, CA 94143, USA; email: chaz.langelier@ucsf.edu

Microbiome and Antimicrobial Resistance Gene Dynamics in International Travelers

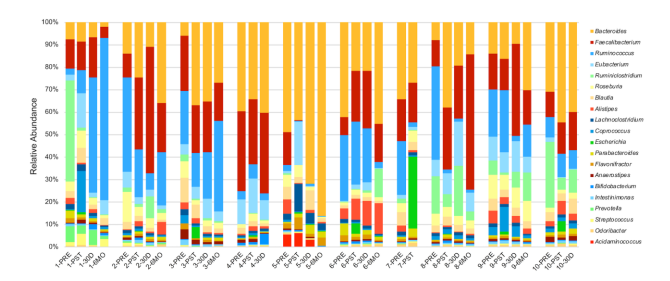
Appendix



Appendix Figure 1. Assessment of relative abundance of *Escherichia* over time, pre- and post-travel, revealed an increase (p<0.001 by Wilcoxon Rank Sum) post-travel.



Appendix Figure 2. Relative proportion of the 20 most abundant microbial genera present in the enteric microbiomes of international travelers, assessed longitudinally using shotgun metagenomic DNA sequencing. Participants submitted samples pre-travel (PRE, within 1 week before departure), post-travel (PST, within 1 week after return), 30 days post-travel (30D), and 6 months post-travel (6MO).



Appendix Figure 3. Beta-lactamase genes identified in the gut microbiomes of travelers over time, preand post-travel. Gradient shading corresponds to gene abundance (average allele sequencing depth per million reads mapped, AdpM).