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# **ORIGINAL RESEARCH**

# Metagenomic Study of the MESA: Detection of *Gemella Morbillorum* and Association With Coronary Heart Disease

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**BACKGROUND:** Inflammation is a feature of coronary heart disease (CHD), but the role of proinflammatory microbial infection in CHD remains understudied.

METHODS AND RESULTS: CHD was defined in the MESA (Multi-Ethnic Study of Atherosclerosis) as myocardial infarction (251 participants), resuscitated arrest (2 participants), and CHD death (80 participants). We analyzed sequencing reads from 4421 MESA participants in the National Heart, Lung, and Blood Institute Trans-Omics for Precision Medicine program using the PathSeq workflow of the Genome Analysis Tool Kit and a 65-gigabase microbial reference. Paired reads aligning to 840 microbes were detected in >1% of participants. The association of the presence of microbe reads with incident CHD (follow-up, ~18 years) was examined. First, important variables were ascertained using a single regularized Cox proportional hazard model, examining change of risk as a function of presence of microbe with age, sex, education level, Life's Simple 7, and inflammation. For variables of importance, the hazard ratio (HR) was estimated in separate (unregularized) Cox proportional hazard models including the same covariates (significance threshold Bonferroni corrected *P*<6×10<sup>-5</sup>, 0.05/840). Reads from 2 microbes were significantly associated with CHD: *Gemella morbillorum* (HR, 3.14 [95% CI, 1.92–5.12]; *P*=4.86×10<sup>-6</sup>) and *Pseudomonas* species NFACC19-2 (HR, 3.22 [95% CI, 2.03–5.41]; *P*=1.58×10<sup>-6</sup>).

**CONCLUSIONS:** Metagenomics of whole-genome sequence reads opens a possible frontier for detection of pathogens for chronic diseases. The association of *G morbillorum* and *Pseudomonas* species reads with CHD raises the possibilities that microbes may drive atherosclerotic inflammation and that treatments for specific pathogens may provide clinical utility for CHD reduction.

Key Words: cardiovascular heart disease ■ Gemella morbillorum ■ MESA ■ metagenomics ■ Pseudomonas

nflammation is a well-established feature of cardiovascular disease.<sup>1,2</sup> The hypothesis that pathogens contribute to this inflammation has seen acceptance and rejection since the demonstration in 1979 that an avian herpesvirus caused atherosclerotic-like lesions in chicken arteries and cholesterol accumulation in chicken smooth muscle cells.<sup>3</sup> The subsequent observation that *Chlamydia* infection increased atherosclerosis in mouse models led to clinical trials of antibiotics targeting *Chlamydia pneumoniae*, but results were uniformly negative for reduction of cardiovascular risk. Enthusiasm for the hypothesis therefore waned as equivocal results were obtained from studies of "candidate pathogens," including *Streptococcus* species, *Human papilloma virus*, *Helicobacter pylori*, and the viruses Epstein-Barr, hepatitis A, *Herpes* species, and influenza (for review, see previous studies<sup>4,5</sup>).

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# RESEARCH PERSPECTIVE

#### What Is New?

- We conducted a metagenomic analysis of whole-genome sequencing data from the MESA (Multi-Ethnic Study of Atherosclerosis). Sequencing reads that do not align with the human reference were aligned with a microbial reference with bacterial, fungal, unicellular eukaryote, and viral sequences.
- We found an association between the presence of Gemella morbillorum and Pseudomonas reads and subsequent cardiovascular heart disease.
- These organisms were also detected in samples from atheroclerotic plaque.

# What Questions Should Be Addressed Next?

- Because the detection of microbes by reanalysis of sequencing data from human blood samples is currently controversial, can these findings be confirmed by other approaches?
- Can these findings be confirmed by reanalysis of sequencing data in other cohorts?

# Nonstandard Abbreviations and Acronyms

MESA Multi-Ethnic Study of Atherosclerosis

However, evidence continues to accumulate for the hypothesis that microbes contribute to inflammation in cardiovascular diseases. For example, in epidemiology studies, severe infection, defined as hospital admission for sepsis or pneumonia, was associated with increased risk for cardiovascular disease in the Swedish Military Conscription Register<sup>6</sup> (236739 subjects), OptumLabs Data Warehouse (2258464 hospitalizations),7 and UK BioBank (331683 for discovery, 271329 for replication).8 In the laboratory, treatment of monocytes with microbes results in macrophages with an atherosclerotic phenotype as defined by cytokines and foam cell formation.<sup>9,10</sup> In biomarker studies, cell-free DNA in plasma has been associated with cardiovascular disease. 11,12 In addition, clinical trials of anti-inflammatory treatments, such as colchicine and anti-interleukin-1ß (canakinumab), have been successful at reducing the recurrence of cardiovascular events after myocardial infarction. 13-15 These observations revive interest in the hypothesis that microbes contribute to cardiovascular disease and suggest that success with a clinical trial of antimicrobial therapy awaits identification of organisms more pathogenic for the cardiovascular system.

Although the blood has traditionally been considered to be free of microbes when healthy, microbes have been detected, 16 possible sources are the gut microbiome, particularly in dysbiosis, and the oral microbiome, particularly in periodontitis. One method for detection of microbial organisms is a "metagenomic analysis," the examination of short sequence reads from next-generation sequencing that do not align with the human reference and so are "left over" from a sequencing experiment. 17 Alignment of these "left overs" with a microbial reference file may identify known microbes, 18 and contig assembly of overlapping short reads may identify hitherto unculturable and unknown microbes. 19

To detect microbes and test for association with incident coronary heart disease (CHD), we report a metagenomic study of short-read, whole-genome sequence data from 4424 participants of the MESA (Multi-Ethnic Study of Atherosclerosis), generated as part of the National Heart, Lung, and Blood Institute Trans-Omics of Precision Medicine program.<sup>20</sup>

### **METHODS**

#### **Data Access**

We used data publicly available data from the National Center for Biotechnology Information with the following accessions: MESA, PRJNA396088; PRJNA991655<sup>21</sup>; and PRJNA242791.<sup>22</sup> We used PathSeq and the Genome Analysis Tool Kit as implemented in the BioData Catalyst without modification, except that we updated the human reference in the Genome Analysis Tool Kit bundle to GRCh38.p14 (see Metagenomics analysis, below).

### Study Participants: MESA

MESA is a prospective, community-based cohort study of 6814 men and women aged 45 to 84 years, free of clinical cardiovascular disease at the time of enrollment in 2000 to 2002. Participants were recruited from 6 US regions: Baltimore, MD; Chicago, IL; Los Angeles, CA; New York NY; St. Paul, MN; and Winston-Salem, NC, and at baseline had the following sex and ethnicity (selfreported) distributions: 53% female, and 38% non-Hispanic White (self-report "White or Caucasian" and not "Spanish, Hispanic, or Latino" in 2000-2002), 28% African American, 12% Chinese, and 22% Hispanic (both Caribbean and Mexico/Central America). All participants provided written informed consent, including for genetic study. MESA has been approved by the institutional review boards of each field center, the Data Coordinating Center at the University of Washington (Seattle, WA), the Central Laboratory at the University of Vermont (Burlington, VT); and the Genetic Analysis Center and CT Reading Center at The Lundquist Institute (Torrance, CA). The Institutional Review Board at The Lundquist Institute for Biomedical Innovation approved this specific project. Extensive details of MESA have been described. 23,24 At the baseline examination, clinical characteristics and anthropometric measurements were obtained by trained personnel using standardized protocols. A fasting blood sample was also drawn (after a minimum 8-hour fast) and stored at -80 ° C until DNA isolation. Questionnaires were administered to collect self-reported demographic data, including age, sex, race, dietary information, and health behaviors. Subsequently, participants participated in follow-up telephone calls at yearly intervals for the ascertainment of cardiovascular events, and in clinical examinations at ≈18-month intervals.

# **Coronary Heart Disease**

CHD events were adjudicated from yearly telephone follow-up, medical records, and the National Death Index. CHD was defined by myocardial infarction, resuscitated cardiac arrest, and CHD death ("hard" CHD in the MESA protocols). Agatston score was determined by computed tomography.

## **Covariates**

Age, sex, household income, highest education levels, and smoking status were obtained through in-person interviews with trained assessors. Participants' smoking information was categorized into current smokers compared with former/never smokers. Physical activity was assessed using a detailed, semiquantitative questionnaire adapted from the Cross-Cultural Activity Participation Study. Habitual dietary intake was assessed via a food frequency questionnaire that asked about the frequency of intake, and typical portion size, of 120 foods (including mixed dishes, such as chow mein) over the past 12 months. Height and weight were measured in duplicate by trained study staff. A mean of both measurements was used to calculate body mass index as weight in kilograms (kg) divided by height in meters (m) squared (kg/m²). Interleukin-6 was measured via ultrasensitive ELISA (Quantikine HS Human IL-6 Immunoassay; R&D Systems, Minneapolis, MN). Health behavior and clinical information at baseline was summarized into the American Heart Association's Life's Simple 7 score; this score reflects proximity to "ideal cardiovascular health" by aggregating (1) smoking status; (2) body mass index; (3) physical activity; (4) healthy diet; (5) total cholesterol; (6) blood pressure; and (7) fasting plasma glucose.<sup>25,26</sup>

# **Metagenomic Analysis**

The scoring of the presence/absence of microbes was ascertained via metagenomic analysis of whole-genome

sequence data from MESA as part of the Trans-Omics for Precision Medicine program of the National Heart, Lung, and Blood Institute. Sequencing was by the Broad Institute Genomics Platform (Stacey Gabriel, principal investigator), reads transferred to the Trans-Omics of Precision Medicine Informatics Research Center (University of Michigan), and aligned to the human reference with stringent machine-learning-based quality metrics.<sup>20</sup> Average read length was 151 nucleotides. Quality of the sequencing data can also be seen by the mean and median read depth of 38x and the high proportion of the human genome covered with a depth of >10× (mean, 0.979; median, 0.987). Sequence files ("Cram files") were transferred from the Sequence Read Archive to a Terra/BioDataCatalyst workspace via Gen3 (The National Heart, Lung, and Blood Institute BioData Catalyst, Zenodo.).<sup>27</sup> The PathSeq workflow in the Genome Analysis Tool Kit, and as developed for BioData Catalyst, 28-31 was applied in 3 steps: first, to remove human reads with low complexity, low quality, and duplication; second, to align the remaining short reads to a microbial reference, consisting of 65 gigabases of sequence from archaea, bacteria, fungi, protozoa, and viral genomes; then third, to perform a taxonomic classification, matching aligned reads with known microbes. A microbial-aligned read was considered "detected" if the alignment had an identity score > 90% with both read pairs aligning to the same microbe reference fasta file in the microbial reference; these are considered "unambiguous" by the PathSeq workflow. Because of zero inflation for the population distribution of transcript read frequency for several microbes, metagenomic reads for each were converted to indicate the presence or absence of each microbial organism in the sample.

# Statistical Analysis Sample Characteristics

Demographic information and health characteristics, stratified by quartiles of total number of microbes detected via transcript reads, were calculated as total number (N) and percentage (%). Crude differences between the quartiles (ie, without controlling for covariates or correcting for multiple testing) were conducted using linear regression on transformed outcomes for continuous variables (P trend interpreted) and a  $\chi^2$  test for binary or other categorical traits.

# Associations Between the Detected Reads and Incident CHD

All microbes with presence or absence of at least 1 read in >1% of participants were included in tests of association with incidence of CHD. Although HIV was included in our microbial reference, there were no

groups of patients with HIV in this study. Furthermore, MESA ancillary studies have indicated that <1% of participants are positive for either hepatitis B or hepatitis C, and so these microbes were not included in our analyses.

#### Variable Selection

Because of the prevalence of detected reads being as low as 1% for some microbes, we first selected variables of importance using a single penalized Cox proportional hazard ("survival") model. Our rationale was that penalized models can minimize the increased likelihood of false positives seen with sparse data, particularly when there are few events.32-34 Elastic net was used to apply a penalty function to the Cox proportional hazards models, to account for the sparse nature of the data structure for some microbes. Optimal penalty parameters for the penalty value (mixing percentage;  $\alpha$ ) and the strength of the penalty (regularization penalty; λ) were ascertained via the R package "caret" using cross validation. Briefly, data in the full data set were randomly assigned to training (2/3 of participants) and test (1/3 of participants) data sets. Parameter selection was conducted via bootstrapped estimates (25 repeats) of models for all values of  $\lambda$  between 0 and 1 (inclusive) at intervals of 0.05. Optimization was reached via feature-wise normalization change in successive coordinate descent iterations.<sup>35</sup> Model performance was judged on the basis of root mean square error of approximation, with  $\alpha$  and  $\lambda$  parameters giving rise to the minimum mean cross-validated error used to generate new coefficients for the association of plasma microbe transcripts with incident events.

#### **Effect Size Estimation**

As penalized models do not provide confidence estimates around associations, for all microbes selected as variables of importance in the penalized models, the association was also parameterized using standard (nonpenalized) Cox proportional hazard models, modeling how the risk of experiencing a CHD event changes as a function of the presence of each microbe over time. Each microbe selected as a variable of importance in the regularized models was included in a separate model, and all models adjusted for were age, sex, self-reported race and ethnicity, highest education (as a proxy for socioeconomic status), Life's Simple 7 score, and interleukin-6 levels (as a proxy for inflammation). Microbial transcriptome-wide significance for these models was set at a Bonferroni-corrected P<5.95×10<sup>-6</sup> (0.05/840 included microbes).

#### Sensitivity Analyses

For microbes detected in plaque, we repeated the detected reads-event analyses, after stratifying participants on baseline Agatston score=0 or >0, to probe for the possibility that the presence of microbes followed, or occurred simultaneously with, the onset of pathology given that Agatston score provides a general adjustment for the observed confounding of coronary artery calcium.

# **RESULTS**

# **Sample Characteristics**

After data merge, the total number of participants in the complete data set for this report was 4421. Over an average of 15.61±3.45 follow-up person years (total person years=69007), we observed 333 CHD events (251 myocardial infarctions, 2 resuscitated cardiac arrests, and 80 CHD deaths). The mean number of nonhuman microbes detected by metagenomic sequencing in each participant was 79.15±20.05 (range, 30-262; Table 1). The number of microbes detected differed between groups defined by self-reported race and ethnicity ( $X_2$ =30.39, df=9, P=3.8×10<sup>-4</sup>, Table 1), and those who went on to develop CHD ( $X_2=9$ , df=3, P=0.02). No association was found between the average number of microbes detected and baseline age. sex, total cholesterol, systolic blood pressure, highdensity lipoprotein cholesterol, current smoking status, or diabetes status (all P>0.05; Table 1).

A total of 5335 (of 5379 or 99.2%) MESA "cram" files were successfully processed by the PathSeq workflow. Mean number of nonhuman reads per participant was ~350000 of a mean of ~860 million total reads. The mean percentage of these that subsequently aligned on the microbial reference was 2.3%. The number of microbes detected by at least 1 read was 4460, and the number of microbes detected in >1% of the participants was 840.

#### Association With Incident CHD

Our rationale for beginning with a regularized Cox proportional hazard model is that this approach is useful for the sparse and high dimensional data in this report. 32,33 In regularized Cox proportional hazard models that included the presence/absence of transcripts for 840 microbes, along with age, sex, self-reported race and ethnicity, highest education level, Life's Simple 7 score, and interleukin-6 values, the presence of gene transcripts for 10 microbes were selected as variables of importance for predicting the likelihood of experiencing a subsequent incident CHD event (Table 2). After a Bonferroni correction for multiple testing, 2 microbes met were significant for association with incident

Table 1. Sample Characteristics at Baseline, Stratified by Quartile of Number of Microbial Species Detected

	Quartile						
Characteristic	1 (N=1172)	2 (N=1057)	3 (N=1148)	4 (N=1044)			
Demographic characteristics							
Age, mean±SD, y	60.92±9.80	61.01±9.84	61.25±9.89	60.77±9.82			
Sex, N (%) male	626 (53.41)	566 (53.55)	580 (50.52)	513 (49.14)			
Self-reported race and ethnicity, N (%)*							
Non-Hispanic White	442 (37.71)	456 (43.15)	453 (39.46)	440 (42.15)			
Chinese	201 (17.15)	128 (12.11)	141 (12.28)	113 (10.83)			
African American	257 (21.93)	239 (22.61)	295 (16.99)	262 (25.10)			
Hispanic	272 (23.21)	234 (22.14)	259 (22.56)	229 (21.93)			
Metagenomic sequencing							
Microbial species detected, mean±SD, N	56.48±7.82	73.33±3.11	84.01±3.45	105.16±17.2			
Clinical/health characteristics							
Life's Simple 7 score, mean±SD	8.79±2.07	8.61±2.13	8.60±2.05	8.68±2.06			
Outcomes							
CHD event, N (%)	78 (6.66)	63 (5.96)	99 (8.62)	93 (8.91)*			

<sup>\*</sup>P<0.05 in tests of difference.

CHD: Pseudomonas (hazard ratio [HR]= $3.32\pm0.34$ ,  $P=1.58\times10^{-5}$ ; penalized HR=1.5; Table 2), and Gemella morbillorum (HR= $3.14\pm0.25$ ,  $P=4.86\times10^{-6}$ ; penalized HR=2.40; Table 2).

Subsequently, we considered the role of overlapping infections in incident CHD. Because we detected both *Pseudomonas* and *G morbillorum* in only 1 individual,

we did not estimate the association of joint infection with these microbes on incident CHD. However, there was overlap in infections across the 10 microbes selected as variables of importance in the penalized Cox proportional hazards model. We detected none of these 10 microbes in 56% of the sample, 1 of the 10 microbes in 36%, and >1 in 8%. Thus, the number of

Table 2. Frequencies and Percentages of Individuals With the Presence of Transcripts in the Plasma at Baseline, Stratified by Those Who Did/Did Not Experience an Incident CHD Event, for All Microbial Species Selected as Variables of Importance in Regularized Models for Incident CHD, and Parameter Estimates From Nonregularized Survival Models

	Frequencies	Parameter estimates		
Species	did not develop incident CHD (N=4088)	Developed incident CHD (N=333)	HR (±95% CIs)	P value
Mesorhizobium species LNJC395A00	1390 (34)	127 (38)	1.23 (0.94–1.54)	0.07
Pseudomonas species NFACC19-2	58 (1.42)*	17 (5.11)*	3.32 (2.03–5.41)*	1.58×10 <sup>-6*</sup>
Porphyromonadaceae KA00676	44 (1.08)	10 (3.00)	3.04 (1.56–5.91)	0.001
Streptococcus species HMSC071D03	42 (1.03)	9 (2.70)	3.14 (1.61–6.11)	7.56×10 <sup>-4</sup>
Gemella morbillorum	70 (1.71)*	18 (4.41)*	3.14 (1.92–5.12)*	4.86×10 <sup>-6*</sup>
Laccaria bicolor	63 (1.54)	12 (3.60)	2.16 (1.18–3.94)	0.01
Citrobacter braakii	48 (1.17)	12 (3.60)	2.24 (1.25–4.02)	0.007
Streptomyces griseorubens	238 (5.82)	28 (8.41)	1.39 (0.94–2.04)	0.10
Prevotella species oral taxon_306	45 (1.00)	9 (2.70)	2.49 (1.28–4.84)	0.007
Buttiauxella gaviniae	52 (1.27)	11 (3.30)	1.89 (1.03–3.46)	0.04
Anaerococcus provenciensis	45 (1.00)	10 (3.00)	2.84 (1.51–5.34)	0.001

All models controlled for age, sex, highest education level, income, Lifes's Simple 7 score, and interleukin-6. CHD indicates coronary heart disease; and HR, hazard ratio.

<sup>\*</sup>Significant associations (P<1.5×10<sup>-5</sup> after a Bonferoni correction).

microbes present, from all microbes selected as variables of importance to incident CHD, was associated with incident CHD (HR=1.51 $\pm$ 1.05, P=2.07×10<sup>-9</sup>), with each additional microbe increasing the risk of incident CHD by 1.5 times. In comparison, when considering all microbes present in at least 1% of the sample, total microbial burden (ie, the number of microbes detected) was not significantly associated with incident CHD (HR=1.00 $\pm$ 1.00, P=0.04).

# **Detection in Plaque Samples**

To find additional support for the presence of *G morbil*lorum and Pseudomonas species NFACC19-2, we applied PathSeq to data available in the Sequence Read Archive. Our rationale for repeating an analysis of these data was that the original articles did not present results at the microbe level and that we wanted to increase our chance of detection by selecting data sets with CHD-related characteristics. First, we applied PathSeq to 12 files in the Sequence Read Archive (SRA040611), composed of DNA from carotid atherosclerotic plaque obtained from endarterectomy of 7 patients after cerebral ischemia or stroke and from autopsy of 5 patients with cause of death unrelated to cardiovascular disease.<sup>23</sup> G morbillorum reads were detected in 4 of 5 of the autopsy and 2 of 7 of the endarterectomy samples. Notably, 50 Gemella reads were detected in 1 of the autopsy samples, a number 5 times the highest number observed in MESA participants. Pseudomonas species NFACC19-2 was detected in 1 autopsy sample. Second, we applied PathSeg to 27 RNA-sequencing files generated by SMART sequencing<sup>24</sup> (BioProject PRJNA991655; GSE236610), composed of RNA samples retrieved directly from balloons in percutaneous coronary interventions; G morbillorum reads were detected in 6 of 13 patients with stable CAD and 10 of 14 with acute coronary syndrome and reads from Pseudomonas strain NFACC19-2 were observed in 2 of 13 patients with stable CAD and 9 of 14 patients with acute coronary syndrome.

# **Sensitivity Analyses**

We stratified analyses by Agatston score, a measure of calcium deposit in the coronary artery, 36 to probe for

the possibility that atherosclerotic processes predispose individuals to infection of the putative microbes, in part driving the link between baseline infection and incident CHD. Of those with an Agaston score of 0, N=29 experienced an incident CHD event, of whom 2 showed the presence of bacteria at baseline for each of *G morbillorum* or *Pseudomonas* species NFACC19-2. This precluded an analysis of these microbes within Agaston strata individually. Of those with an Agaston score of 0 at baseline, who subsequently experienced incident CHD, N=18 showed the presence of at least 1 microbe selected as a variable of importance to incident CHD. Analysis of these data did not suggest modification of the associations (Table 3).

### DISCUSSION

We conducted a metagenomic study of the short whole-genome sequencing reads from the blood samples of 4421 participants in the MESA and report a significant association between subsequent CHD events and detection of sequence reads from 2 bacteria, *G morbillorum* and *Pseudomonas* strain NFACC19-2. With the same workflow, we detected these 2 bacteria in publicly available data from endarterectomy samples and from RNA retrieved from balloons from percutaneous coronary interventions.

There are numerous reports of infection attributed to *G morbillorum*, for example, endocarditis,<sup>37–39</sup> osteoarticular infection,<sup>40</sup> necrotizing soft tissue infection,<sup>29,30</sup> and septic shock in the immunocompromised.<sup>41</sup> *G morbillorum* has been detected in 95% of oral samples (Human Oral Microbiome Database V3.1), and 1% of gut samples (the Human Microbiome Project I; microbiomedb.org); this observation suggests that the oral microbiome is the source of *G morbillorum*. The fact that the genus *Gemella* harbors virulence factors from multiple *Gemella* strains may explain inconsistencies in the designation of microbes associated with infection.<sup>42</sup>

The reference sequence for *Pseudomonas* species NFACC19-2 (GCF\_900119125.1) has not been further assigned to a species as of this report. The lack of assignment points to some of the problems with microbial studies: (a) large numbers of microbial species are poorly characterized, (b) some have not been cultured,

Table 3. Parameters From Cox Proportional Hazard Models for the Association of Microbial Transcripts (Present Versus Absent) With Incident CHD, Stratified by Baseline Agaston Score (0 Versus >0), for All Microbes Significantly Associated With Incident CHD in the Whole Population

	Agatston score=0 (N=1466)		Agatston score >0 (N=2828)	
Variable	HR (+/- 95% CIs)	P value	HR (+/- 95% CIs)	P value
Pseudomonas species NFACC19-2	3.27 (1.91–5.60)	1.67×10 <sup>-5</sup>	6.45 (1.50–27.74)	0.01
Gemella morbillorum	4.46 (1.09–19.99)	0.04	3.05 (1.81–5.15)	2.95×10 <sup>-5</sup>

All models controlled for age, sex, highest education level, income, Lifes's Simple 7 score, and interleukin-6. CHD indicates coronary heart disease; and HR, hazard ratio.

and (c) and species assignment changes, making comparisons to older literature difficult. Furthermore, earlier microbiome studies used sequencing of the ribosomal region and do not have the resolution of later studies. However, the related *Pseudomonas aeruginosa* is a model organism for *Pseudomonas* infection, an opportunistic pathogen well known for infections in immunocompromised hosts, for nosocomial infections, and for its ability to develop multidrug resistance.

The strengths of this study include the high quality of data available from both MESA and Trans-Omics of Precision Medicine for >4000 subjects with median follow-up of 17 years. Furthermore, we reduced false-positive results by using the PathSeq workflow, filtering to remove low-quality and duplicate reads, low-complexity sequence, human Y chromosome sequence, and residual human reads before alignment with the microbial reference. We confined our analysis to microbes detected in >1% of subjects and with a >90% alignment of both paired reads to the same reference (designated "unambiguous" by the software). The association of the Gemella and Pseudomonas remained significant after correction for the number of microbes (840 in >1% of participants). We further detected the same microbes in sequence reads from atherosclerosis-related tissues.

A major weakness of this study is that the microbes were not detected directly by microbiological culturing but indirectly by alignment of reads with a microbial reference. The inference of microbial viability in the blood from sequence reads without the presence of overt infection or confirmatory culturing is currently controversial, and the sensitivity and specificity of metagenomics to various microbes remains under investigation. A second concern of our study is that our analytical model was applied to a small number of CHD events with 6 adjustment variables and so parameter estimates may be unstable. Therefore, these results present an association warranting further investigation and replication, and should not been seen as clinically actionable. We think replication is required, with confirmation of these results to include: (a) additional metagenomic analyses of whole-genome sequencing files in major studies with cardiovascular phenotypes (a larger sample size); (b) identification of microbes in atherosclerotic plaque by means of microbiological methods (eg, culturing and biochemical testing); and (c) once greater support has been observed, testing the effect of identified microbes in animal models of cardiovascular disease (testing whether exposure to a given microbe causes or exacerbates CHD). The third major concern is the lack of longitudinal data on infection persistence, and on factors that could link infections to CHD, such as inflammation. Thus, we have refrained from either causal or mechanistic interpretations, as is appropriate for the observational nature of our findings,

and await the results from a variety of possible further studies.

Given the recent success of clinical trials of inflammation therapies in reducing cardiovascular disease. our results suggest that the metagenomic exploration of microbial diversity in well-characterized cardiovascular disease cohorts may identify pathogens and zoonoses contributing to heart disease. This exploration should include both alignment with known organisms as assembled in a microbial reference, as in this report, and assembly into contigs for detection of hitherto unknown or unculturable microbes. 43 We speculate that microbe identification and characterization by metagenomics may generate a "short list" of organisms suitable for the development of immunologic therapies for cardiovascular disease, perhaps even to clinical trials of a "vaccine," to reduce or eliminate the constant lowgrade blood infection fed from the oral microbiome.

#### ARTICLE INFORMATION

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#### **Disclosures**

None.

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