1 No evidence for a common blood microbiome based on a

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population study of 9,770 healthy humans

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18 Abstract

19 Human blood is conventionally considered sterile. Recent studies have challenged this, 20 suggesting the presence of a blood microbiome in healthy humans. We present the 21 largest investigation to date of microbes in blood, based on shotgun sequencing 22 libraries from 9,770 healthy subjects. Leveraging the availability of data from multiple 23 cohorts, we stringently filtered for laboratory contaminants to identify 117 microbial 24 species detected in the blood of sampled individuals, some of which had signatures of 25 DNA replication. These primarily comprise of commensals associated with human body 26 sites such as the qut (n=40), mouth (n=32), and genitourinary tract (n=18), which are 27 species that are distinct from common pathogens detected in clinical blood cultures 28 based on more than a decade of records from a tertiary hospital. Contrary to the 29 expectations of a shared blood microbiome, no species were detected in 84% of 30 individuals, while only a median of one microbial species per individual was detected in 31 the remaining 16%. Futhermore, microbes of the same species were detected in <5% of 32 individuals, no co-occurrence patterns similar to microbiomes in other body sites was 33 observed, and no associations between host phenotypes (e.g. demographics and blood 34 parameters) and microbial species could be established. Overall, these results do not 35 support the hypothesis of a consistent core microbiome endogenous to human blood. 36 Rather, our findings support the transient and sporadic translocation of commensal 37 microbes, or their DNA, from other body sites into the bloodstream.

38 Introduction

39 In recent years, there has been considerable interest regarding the existence of a microbiome in the blood of healthy individuals, and its links to health and disease. 40 41 Human blood is traditionally considered a sterile environment (i.e., devoid of viable 42 microbes), where the occasional entry and proliferation of pathogens in blood can 43 trigger a dysregulated host response, resulting in severe clinical sequelae such as 44 sepsis, septic shock or death¹. Asymptomatic transient bacteraemia (i.e., bacterial 45 presence in blood) in blood donors is also known to be a major cause of transfusionrelated sepsis². Recent studies have suggested the presence of a blood microbiome, 46 47 providing evidence for microbes circulating in human blood for healthy individuals³⁻⁷ (reviewed in Castillo et al⁸). However, most of these studies were either done in 48 49 relatively small cohorts or lacked rigorous checks to distinguish true biological measurements from different sources of contamination⁸. As such, the concept of a 50 51 microbial community present in the blood of healthy individuals remains controversial 52 and is an area of active research. In this work, we analysed blood DNA sequencing data 53 from a population study of healthy individuals, comprising of multiple cohorts processed 54 by different laboratories with varied sequencing kits. By leveraging the large dataset (n=9,770) complete with batch information in our systematic differential analyses for 55 56 potential contaminants, our aim was to determine whether a blood microbiome truly 57 exists in the general population.

For meaningful discourse, it is useful to formalise what the presence of a hypothetical 58 'blood microbiome' entails. Berg et al.⁹ concluded that the term microbiome should refer 59 60 to a community of microbes that interact with each other and with the environment in 61 their ecological niche, which in our context is human blood. Therefore in a blood 62 microbiome, the presence of microbial cells in blood from healthy individuals should 63 exhibit community structures indicated by co-occurrence or mutual exclusion of 64 species¹⁰ as seen in the microbiomes of other sites such as the gut¹¹ or mouth¹². Furthermore, we may expect the presence of core microbial species, which can be 65 defined as species that are frequently observed and shared across individuals^{13,14}, such 66 as Staphylococcus epidermidis on human skin¹⁵. More precisely, taxa that are found in 67

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a substantial fraction of samples from distinct individuals (i.e. with high prevalence) may be considered 'core'. Notably, the prevalence threshold for defining core taxa is arbitrary, with previous microbiome studies using values ranging from 30-100% and many of these studies opting for 100%¹⁴. Regardless, identifying core microbes in blood would form the basis for associating microbiome changes with human health.

73 Existing studies have provided evidence for the presence of microbes in the blood of healthy individuals using both culture-based^{3,4} and culture-independent^{5–7} approaches. 74 The former approach involves blood culture experiments while the latter involves one or 75 a combination of the following molecular methods: 16S ribosomal RNA (rRNA) 76 77 quantitative polymerase chain reaction (qPCR), 16S rRNA amplicon sequencing, and 78 shotgun sequencing of RNA or DNA. Depending on the study design, these results 79 should be interpreted with caution due to several methodological and technical 80 limitations which include small sample sizes, limited taxonomic resolution, difficulties in distinguishing cell-free microbial DNA from live microbial cells, and the ubiquity of 81 environmental contamination^{8,16–19}. In particular, contaminating DNA must be accounted 82 83 for in order to characterize the blood microbiome. The workflow of sample processing, 84 from skin puncture during phlebotomy, to microbial detection, is rife with opportunities for microbes or microbial nucleic acids to be introduced. Contaminating microbial cells 85 86 introduced due to poor aseptic technique or insufficient disinfection of the skin puncture site²⁰ affects both culture-dependent and culture-independent approaches. Sequencing-87 based approaches are especially sensitive to contaminant microbial DNA native to 88 laboratory reagent kits (i.e., the 'kitome')¹⁹, exacerbated by the low microbial biomasses 89 90 in blood, accompanied by high host background which increases the noise-to-signal 91 ratio¹⁷. Correspondingly, comprehensive profiling of the breadth and prevalence of 92 microbial species in blood after accounting for external sources of contamination has 93 not yet been done and several aspects of the 'blood microbiome' remain unclear. For 94 instance, are the detected microbes endogenous to blood or translocated from other 95 body sites? Is there a core set of microbes that circulates in human blood? Is there a 96 microbial community whose structure and function could influence host health?

To address these questions, we performed the largest scale analysis of a blood sequencing dataset to date, based on DNA libraries for 9,770 healthy individuals from

99 six distinct cohorts (Supplementary Table 1). We applied various bioinformatic 100 techniques to differentiate DNA signatures of microbes in blood from potential reagent 101 contaminants and sequence analysis artefacts, leveraging the differences in reagent 102 kits used to process each cohort. We detected 117 microbial species in the blood of 103 these healthy individuals, most of which are commensals associated with the 104 microbiomes of other body sites. Additionally, we identified DNA signatures of replicating bacteria in blood using coverage-based peak-to-trough ratio analyses^{21,22}. 105 106 providing a culture-independent survey that has not been achieved previously. Despite 107 this, we found no evidence for microbial co-occurrence relationships, core species, or 108 associations with host phenotypes. These findings challenge the paradigm of a 'blood 109 microbiome' and instead support a model whereby microbes from other body sites (e.g. 110 gut, oral) sporadically translocate into the bloodstream of healthy individuals, albeit 111 more commonly than previously assumed. Overall, our observations serve to establish 112 a much needed baseline for the use of clinical metagenomics in investigating 113 bloodstream infections.

114 **Results**

Robust inference of microbial DNA signatures in blood based on multi-cohortanalysis

117 Blood samples from healthy individuals typically contain low microbial biomass accompanied by high host DNA background¹⁷, making it difficult to discriminate between 118 119 biologically relevant signals from artefactual ones. We first addressed artefacts arising 120 during bioinformatic sequence analysis by performing stringent quality control on 121 samples (Figure 1a), comprising of read quality trimming and filtering, removal of low 122 complexity sequences that are of ambiguous taxonomic origin, exclusion of reads that 123 likely originate from human DNA (**Methods**), and removal of samples with low number 124 of reads (<100 read pairs) of microbial origin after taxonomic classification with 125 *Kraken2*²³. This provided a species-level characterisation of microbial DNA signatures in 126 blood for most (n=8,892) samples. To minimise noise due to false positive taxonomic 127 assigments, we applied an abundance-cutoff based filter to discriminate between 128 species that are likely present from those that could be misclassification artefacts

129 (**Methods**). Additionally, we validated the reliability of the microbial species detected via 130 Kraken2 by comparison to read alignment analysis using reference genomes, where 131 recovery of large fractions of a microbial reference covered uniformly by mapped reads 132 improves our confidence that they are true positives as opposed to sequencing or analysis artefacts^{24,25}. We validated 96% of the microbial species that had sufficient 133 134 read coverage using this mapping-based approach. We further observed an excellent 135 linear relationship between the number of *Kraken2*-assigned read pairs and the number of aligned read pairs on the log10 scale (slope=1.15; F=154, d.f.=1, p<0.001; 136 137 **Supplementary Figure 1**), suggesting that *Kraken2* taxonomic assignments are a 138 reliable proxy for the more precise and stringent read alignment approach. These 139 findings collectively provide confidence that the microbial species detected in our blood 140 sequencing libraries are not likely sequence analysis artefacts.

141 To address artefactual signals arising due to reagent and handling contamination during 142 sample processing, we used a series of stringent decontamination filters (Figure 1a). 143 These filters are based on the idea that contamination artefacts will lead to false positive 144 detections that are often correlated with each other (within-batch consistency) and 145 biased towards specific laboratory batches (between-batch variability; **Supplementary Figure 2**)²⁶, and such analysis was found to be highly effective for *in silico* 146 decontamination in previous studies²⁷⁻²⁹ (Methods). Additionally, the identification of 147 148 batch-specific contaminants in this study was greatly aided by the availability of multiple 149 large cohorts of healthy individuals (**Supplementary Table 1**), and corresponding rich 150 batch information, including reagent kit types and lot numbers. Application of reagent 151 and handling contamination filters resulted in a final list of 117 microbial species that 152 were detected in the whole blood samples of 8,892 individuals (Supplementary Table 153 2). The list of 117 confidently detected microbial species spanned 56 genera, and 154 comprised of 110 bacteria, 5 viruses and 2 fungi.

To estimate the effectiveness of our filtering strategy in improving biological signal while reducing contamination noise, we examined the types of microbial species detected in our dataset before (870 species) and after (117 species) all filters were applied (**Figure 1b-d**). Firstly, the microbial species were cross-referenced against a published list of common genera seen as contaminants in sequencing data as curated by Poore *et al*³⁰

and derived from the list published by Salter et al^{19} . In this list, genera were either 160 classified as likely contaminants, mixed-evidence (i.e., both a pathogen and common 161 162 contaminant), or potential pathogens/commensals. Following decontamination, the 163 proportion of detected species that are classified as contaminants decreased from 21% 164 to 10% (Figure 1b). Next, the microbial species were compared against human blood 165 culture records spanning more than a decade (2011-2021) from a tertiary hospital 166 (Figure 1c). These blood cultures were typically ordered if clinical indications of 167 bacteraemia were present, and therefore represent the range of microbial species that 168 are known to cause symptomatic infection as detected in a clinical setting. The 169 proportion of species that have been cultured from blood increased from 12% to 27% 170 after decontamination, suggesting that our filtering procedures enriched for microbial 171 species which are capable of invading the bloodstream. Finally, we compared the 172 proportion of human-associated microbes before and after decontamination using a host-pathogen association database describing the host range of pathogens³¹ (Figure 173 174 1d). For species that were not found in this database, a systematic PubMed search 175 (Methods) was performed to determine if there was at least one past report of human 176 infection. The proportion of human-associated species increased from 40% to 78% after 177 decontamination, indicating that they are more likely to be biologically relevant. Finally, 178 we tested our results against the null hypotheses that the 117 microbial species 179 retained after decontamination produced the same proportions of species classified as 180 likely contaminants, human-associated, or that were detected in blood culture compared 181 to species picked at random (**Methods**). This analysis suggests that our 182 decontamination filters significantly decreased the proportions of likely contaminants, 183 while increasing the proportions of human-associated species and those detected in 184 blood cultures (p<0.005; Supplementary Figure 3). These results collectively suggest 185 that by using a set of contaminant-identification heuristics, our filters are sensitive and 186 specific in retaining a higher proportion of biologically relevant taxa while removing likely 187 contaminants.

188 Blood microbial signatures from healthy individuals reflect sporadic translocation

189 of DNA from commensals

190 We next determined the fraction of distinct, healthy individuals for which microbes could 191 be detected (i.e., prevalence). Notably, the most prevalent microbial species, C. acnes, 192 was observed in 4.7% of individuals (Figure 2a), suggesting that none of the 117 193 microbes can be considered 'core' species that are consistently detected across most 194 healthy individuals. Additionally, we did not detect any microbial species in most (82%) 195 of the samples after decontamination (Figure 2b), whereas the remaining 18% of 196 samples had a median of only one microbial species per sample. This low number of 197 species detected per sample was not due to insufficient sequencing depth since there 198 was a weak negative correlation between the number of confidently detected species 199 per sample and the microbial read depth (Spearman's ρ =-0.232, ρ <0.001). Furthermore, 200 some samples containing no microbial species had a microbial read count of up to ~2.1 201 million (median=6,187 reads; distribution shown in **Supplementary Figure 4**). That is, 202 even though a considerable number of reads were classified as microbial, they were all 203 assigned to contaminant species. These results suggest that the presence of microbes 204 in the blood of healthy and apparently asymptomatic individuals, as estimated by our 205 detection methods, is infrequent and sporadic.

Given past reports of bacterial translocation from the mouth³² or gut³³ into blood, we 206 207 asked if the microbes we detected could have originated from various body sites. To do 208 so, we assigned potential body site origins to the 117 microbial species detected in 209 blood based on microbe-to-body-site mappings extracted from the Disbiome 210 database³⁴. We found that many (n=59; 50%) of these confidently detected species are 211 indeed human commensals that are present at various human body sites (Figure 2c). 212 While some of these species may be contaminants that have survived our stringent 213 decontamination filters, this observation, together with their low prevalence, suggests 214 that the microbial DNA of many of these species may have transiently translocated from 215 other locations in the body rather than being endogenous to blood. We further 216 categorised the microbial species based on their growth environments (Figure 2d). A 217 significant portion (n=42; 36%) of the species were obligate anaerobes or obligate 218 intracellular microbes, atypical of skin-associated microbes that may be introduced

during phlebotomy², indicating that they are not likely to be sampling artefacts. All in all, the diverse origins of the microbes detected in blood, together with their low prevalence across a healthy population, is consistent with sporadic translocation of commensals, or their DNA, into the bloodstream.

223 Microbial presence in blood (i.e., bacteraemia) is typically associated with a range of 224 clinical sequelae from mild fevers to sepsis. As such, we asked if the common microbes 225 identified in patients with disease-associated bacteraemia are different from those 226 detected in our cohorts of healthy individuals. To do so, we compared the prevalence of 227 microbes detected in the sequenced blood samples against observations from 11 years 228 of hospital blood culture records. The prevalence of microbial genera detected in the 229 hospital blood culture records clearly differed from that in our sequenced blood 230 samples, despite the overlap in detected taxa (Figure 2e). For example, while 231 Staphylococcus, Escherichia and Klebisiella were the predominant genera identified in blood cultures, they were rarely detected in our blood sequencing libraries. We 232 performed a similar comparison with a previous study³⁵ which sequenced blood 233 234 microbial signatures in sepsis patients and found a similar difference in prevalence 235 compared to our dataset (**Supplementary Figure 5**), confirming that our observations 236 are not due to differences in the detection methods (sequencing vs. culture-based) 237 used. If the species detected through sequencing were genuine, and represent 238 microbial cells, these findings may be explained by the potentially higher virulence of 239 pathogens detected in the clinic, which are more likely to cause clinical symptoms in 240 individuals that would result in exclusion during our recruitment process. Conversely, 241 under the same assumptions, our findings suggest that the microbes detected in the blood of healthy individuals are potentially better tolerated by the immune system (e.g. 242 Bifidobacterium spp.³⁶ and Faecalibacterium prausnitizil³⁷ with immunomodulatory 243 properties as gut commensals; Figure 2a). 244

Evidence for replicating microbial cells but without community structure or hostassociations

To better characterise the microbial DNA signatures detected in blood, we asked if they reflect the presence of viable microbial cells as opposed to circulating cell-free DNA. 249 This is because the former would allow for complex microbe-microbe or microbe-host 250 interactions that would be of greater and more direct clinical relevance. In contrast to previous approaches that used microbial cultures^{3,38}, we looked for more broad-based 251 evidence of live bacterial growth in by applying replication rate analyses^{21,22} on our 252 253 sequenced blood samples. This approach is based on the principle that DNA 254 sequencing of replicating bacteria would yield an increased read coverage (i.e., peak) nearer to the origin of replication (Ori) and decreased coverage (i.e., trough) nearer to 255 the terminus (Ter)²². A coverage peak-to-trough ratio (PTR) greater than one is 256 257 indicative of bacterial replication. Through this analysis, we found evidence for 258 replication of 11 bacterial species out of the 20 that were sufficiently abundant to do this 259 analysis (Figure 3a). The median-smoothed coverage plots of the replicating species all 260 exhibited the sinusoidal coverage pattern (in black; Figure 3b) characteristic of 261 replicating bacterial cells²². This contrasts with the even coverage patterns of three 262 contaminants representative identified during the decontamination steps: 263 Achromobacter xylosoxidans, Pseudomonas mendocina and Alcaligenes faecalis (Figure 3c). The Ori and Ter positions determined using coverage biases largely 264 corresponded with an orthogonal method based on the GC-skew³⁹ of bacterial 265 266 genomes, suggesting that the replication rate analyses are reliable. Additionally, all but one of these replicating species are present in hospital blood culture records and in 267 previous reports of bacteraemia⁴⁰⁻⁴⁹ (Figure 3a), indicating their ability to replicate in 268 269 human blood. Overall, beyond the detection of microbial DNA, we present the first 270 culture-independent molecular signatures for microbial replication from blood.

271 Given the presence of live bacteria, we investigated if the microbial species detected showed patterns of microbe-microbe interactions as would be expected from a microbial 272 community. To do so, we computed pairwise *SparCC* correlations⁵⁰ between species, 273 274 where positive and negative values indicate co-occurrence and mutual-exclusion, 275 respectively. SparCC correlation is a reliable metric for assessing co-occurrence since it 276 accounts for the sparse and compositional nature of microbial taxonomic profiles that 277 confound standard correlation inference techniques⁵⁰. We visualised SparCC 278 correlations of the 117 microbial species confidently detected in blood sequencing 279 libraries using network graphs, where each node is a species and each edge represents

280 the co-occurrence/exclusion associations between two species (Figure 4a). We could 281 not detect strong community co-occurrence/exclusion patterns, with most associations 282 being weak (SparCC correlation<0.05), and only 19 pairwise associations exceeding a 283 correlation value of 0.2, with four exceeding a value of 0.3 (Figure 4a). To determine if 284 this result is a function of our stringent decontamination filters, we generated 285 independent network graphs for the five adult cohorts before decontamination filtering 286 and examined the co-occurrence/exclusion associations shared across cohorts. With an 287 already lenient SparCC correlation threshold of 0.2, we identified no associations 288 common to all the network graphs (Figure 4b), indicating that there were no consistent 289 detectable microbial community associations in blood typical of microbiomes in various 290 human body sites.

291 Previous studies have demonstrated the use of blood microbial DNA as a biomarker for disease, demonstrating associations with cancer³⁰, type II diabetes⁵¹ and periodontal 292 293 disease⁵². In a similar vein, we investigated if the presence of microbes was associated 294 with host phenotypes in our dataset. We first examined if microbes were detected more 295 frequently in infants relative to adults. Given that the still-developing immune systems of infants puts them at greater risk of infection relative to healthy adults⁵³, we reasoned 296 297 that the prevalence of microbes in blood may differ within a birth cohort (GUSTO) 298 relative to adult cohorts. Indeed, samples from GUSTO appeared to have a higher 299 prevalence of microbes associated with most human body sites (Supplementary 300 Figure 6a). This was in part, driven by genitourinary tract-associated microbes, 301 Fannyhessea vaginae, Lactobacillus jensenii, Lactobacillus crispatus, Lactobacillus 302 iners, and Gardnerella vaginalis (Supplementary Figure 6b). Similarly, we found 303 enrichment of gut-associated bacteria such as Bifidobacterium spp. in GUSTO 304 (**Supplementary Figure 6c**). These findings suggest that bacterial translocation may be 305 more frequent in infants relative to adults, though differences in sample collection 306 (umbilical cord versus venipuncture) could also explain them. A future study controlling 307 for differences in sampling methods would be useful for further exploration of this 308 observation.

309 Next, we systematically tested for pairwise associations between eight host phenotypes 310 that were documented on the day of blood collection and the presence of each of the 311 117 microbial species detected in blood. These host phenotypes attributes were: sex, 312 ancestry, age, body mass index (BMI), blood total cholesterol (TC), blood triglycerides 313 (TG), systolic and diastolic blood pressure (SBP and DBP). Given the multiple large 314 independent cohorts, we could perform statistical tests on each cohort separately, which 315 allowed us to assess the consistency of identified association patterns across the 316 different cohorts. Since these cohorts were sampled from a homogenous population, 317 true association patterns are expected to be detected repeatedly regardless of cohort. 318 Using this statistical testing approach, we found only five significant microbe-phenotype 319 associations (p<0.05; **Supplementary Table 3**) after adjusting for multiple comparisons. 320 Notably, all but one of the significant associations were present in only one cohort. The 321 exception was C. acnes, which was significantly associated with ancestry in two 322 cohorts. However, while C. acnes was more prevalent in individuals of Malay ancestry 323 within the SEED cohort, it was more prevalent in Chinese individuals within the MEC 324 cohort (Supplementary Figure 7). These cohort specific differences could be due to 325 other demographic variables that were not recorded in this study, or perhaps from C. 326 acnes subspecies differences. To ensure that we did not miss any associations due to 327 the possible non-linearity of host-phenotype and microbial relationships, we also derived 328 categorical phenotypes based on the recorded phenotypic information. These include 329 being elderly (age>=65), and other measures of 'poorer health', such as being obese 330 (BMI>30), having high blood triglycerides (TG>2.3 mmol/L), high total cholesterol 331 (TC>=6.3 mmol/L), or high blood pressure (SBP>=130 and DBP>=80). We then tested 332 for pairwise associations between these derived phenotypes and the presence of any 333 bacteria but found no significant associations (p>0.05; Supplementary Table 4). 334 Collectively, these results suggest no consistent associations between the presence of 335 microbes in blood and the host phenotypes tested within a healthy population of 336 individuals.

337 **Discussion**

We present the largest scale analysis, to date, of microbial signatures in human blood with rigorous accounting for computational and contamination artefacts and found no evidence for a common blood microbiome in a healthy population. Instead, we observed

341 mostly sporadic instances of blood harbouring DNA from single microbial species of 342 diverse bodily origins, some of which might be actively replicating. Our findings hint at 343 the possibility that the bloodstream represents a route for movement of microbes 344 between different body sites in healthy individuals. However, the low prevalence of the 345 detected species suggest that this movement is likely to be infrequent and transient. 346 Unresolved questions remain about how interconnected the microbiomes at various 347 body sites are, and whether these processes are altered during disease or throughout a 348 person's lifetime. Can perturbations to the microbial community at one body site affect 349 that at another site, and how does the host immune system asymptomatically regulate 350 microbial presence in blood? Our study lays the groundwork for future investigations 351 into these questions, which may pave the way for a systemic understanding of the 352 human microbiome across body sites in relation to human health and disease.

353 We employed a series of decontamination filters to differentiate microbial signatures in 354 blood from artefactual signals associated with reagent and handling contamination, on 355 the basis that the latter display strong batch-specific biases (**Supplementary Figure 2**; 356 see **Methods**). Although our approach substantially improved the signal-to-noise ratio 357 (Fig. 1b-d), it is still likely not fully effective in removing contaminants, evidenced from 358 the fact that 10% of the 117 microbial species remaining after decontamination were still 359 flagged as being of environmental or non-human origin (Fig. 1b, "likely contaminant"). 360 Hence, we recommend that any decontamination procedures should include further 361 comparisons to various microbiome databases (Fig. 1b-d) to prioritise species for 362 validation in future studies. For example, one might prioritise species that are not 363 common contaminants, detected in blood cultures, and that are human associated 364 (Supplementary Table 2) for follow-up experiments. Nevertheless, it is important to 365 note that we could not detect a common blood microbiome despite the likely presence 366 of residual contamination artefacts.

We observed signatures of replicating DNA from putatively genuine microbial species in blood by applying an existing PTR-based replication analysis approach. However, we cannot distinguish signals arising from replicating microbes in blood from those derived from microbial cells (intact or otherwise) which were recently replicating at other body sites before entering the bloodstream. Interestingly, while we could detect replication 372 signatures in blood associated with 11 out of 20 species with sufficient coverage across 373 their genomes, we could not detect any amongst the 20 most prevalent contaminant 374 species identified by our decontamination filters, including species from the genera 375 Alcaligenes, Caulobacter, Bradyrhizobium and Sphingomonas, suggesting that the 376 replication signatures detected in our dataset are not likely to be due to 'kitome' 377 contamination. Furthermore, this observation highlights the potential use of replication 378 analyses for discriminating between putatively genuine taxa from 'kitome' contaminants 379 in future metagenomic studies.

380 We found no core species in human blood on the basis of low prevalence across 381 individuals in our population-level dataset. The prevalence estimates provided in this 382 study are contingent on the sensitivity of detecting microbes through sequencing. 383 Previous studies have shown that untargeted shotgun sequencing is highly sensitive for 384 the detection of microbes in blood at a total sequencing depth of 20-30 million reads per sample^{35,54,55}, perhaps even more so than culture-based methods^{56,57}. In contrast, a 385 386 median of 373 million reads was generated per sample for our sequencing libraries. 387 suggesting that our methods do not lack sensitivity. Our prevalence estimates are also 388 affected by the abundance thresholds used to determine whether a species is present in 389 a single sample (i.e., abundance filter; **Figure 1a**). We defined these thresholds in terms 390 of both absolute read count and relative abundance, which were determined based on 391 simulation experiments (see Methods). Overly stringent abundance thresholds would 392 lead to the erroneous masking of genuine signals, leading to an underestimation of 393 microbial prevalence. However, even when relaxing the threshold to just a relative 394 abundance of 0.001, none of the species, whether flagged as a contaminant or not, had 395 more than 52% prevalence (Supplementary Table 5). Furthermore, the 20 most 396 prevalent species at this threshold are all environmental microbes, and mostly comprise 397 of Sphingomonas and Bradyrhizobium species, which are known to be common sequencing-associated contaminants¹⁹. This suggests that independent of our 398 399 decontamination filters, none of the species detected qualify as core members.

In addition to not being able to detect any core species, we could not detect any strong
 co-occurrence or mutual exclusion associations between species regardless of whether
 our decontamination filters were applied. These associations generally reflect

cooperation or competition between species, respectively⁵⁸. Indeed, within a microbial 403 404 community, metabolic dependencies of species and the ability of different species to 405 complement these dependencies have been shown to be a key driver of microbial cooccurrence⁵⁹. On the other hand, competitive behaviours such as nutrient sequestration 406 407 to deprive potential competitors of nutrients or producing adhesins to bind and occupy favourable sites in an environment⁶⁰ can lead to mutual exclusion between species. The 408 409 fact that we could not detect any strong associations therefore points to the absence of 410 an interacting microbial community in healthy humans. Of note, since our dataset was 411 derived from circulating venous blood, we are, in principle, not able to detect microbial 412 interactions that may be occurring at other sites of the bloodstream such as the inner 413 endothelial lining of blood vessels. Experiments investigating the adherence of bacteria 414 to blood vessel linings may provide further insight into this.

415 The availability of 11 years of blood culture records from the same country of origin as 416 our blood samples enabled a reliable comparison of the prevalence of microbes in the 417 healthy population and in the clinic. This is because the frequency of infections caused by different microbial species is known to differ from country to country⁶¹. Despite this, 418 419 we expect that some of the variation in prevalence estimates may be due to the 420 differences in detection methods. That said, previous studies have shown a strong concordance between culture and sequencing-based detection^{35,54,56,57}. suggesting that 421 422 the distinction between the prevalence of microbes found in healthy individuals and in 423 the clinic is not due to the differences in detection methods. Our results support the 424 conclusion that microbial presence in blood (i.e., bacteraemia) does not always lead to 425 disease. These results are consistent with our other observation that microbial DNA 426 detected in our cohorts of asymptomatic individuals tend to be from commensals, which 427 may inherently be less virulent and better tolerated by the host compared to disease-428 causing pathogens. Indeed, the long-standing co-evolution of humans and colonizing 429 microbes, places a selective pressure against high virulence phenotypes in these microbes to maintain host viability⁶². Simultaneously, there is a selective pressure for 430 431 immunomodulatory phenotypes in commensals to improve their fitness, evidenced by the wealth of immunomodulatory activities found in the gut microbiome⁶³. This agrees 432 433 with previous findings that colonisation by commensals modulate early development of

the immune system⁶⁴, which would allow a measured and regulated response against 434 435 translocated commensals. By extension, the immunomodulatory properties of bacteria 436 and their links to host tolerance to bacteraemia may be key factors in determining 437 clinical outcomes. Perhaps, the presence (or lack) of these properties may determine 438 whether an individual with bacteraemia is asymptomatic or septic. For example, 439 abundant gut bacterial species such as *Bacteroides* spp. were not commonly detected 440 in blood. Further exploration into the immunomodulatory activities of commensals vis-à-441 vis common blood culture pathogens may be the key to design therapeutics to manage 442 or prevent the dysregulated host response that defines sepsis¹.

443 We found no convincing associations between both measured (e.g. TC, SBP) and 444 derived (e.g. obesity) host phenotypes with microbial presence that were consistent 445 across the different cohorts. This suggests that the risk of transient microbial 446 translocation, at least across our cohorts of healthy adults, is fairly consistent. In 447 contrast, this risk may increase in individuals with more severe disease. In fact, variable 448 microbial DNA profiles in blood have been used to delineate health and disease states. This has most prominently been shown for sepsis^{35,54–57,65}, where the presence of viable 449 microbes is expected, but also for cancer³⁰, periodontal disease⁵², and chronic kidney 450 451 disease⁶⁶, which are unrelated to bloodstream infections. These studies highlight the 452 promise of metagenomic sequencing of blood for developing diagnostic, prognostic, or 453 therapeutic tools. Our characterisation of the species breadth in healthy individuals 454 forms a crucial baseline for comparison with that in diseased individuals. Indeed, our 455 findings open new doors to understanding why and how blood microbial profiles 456 correlate with health status. One possible hypothesis is that mucosal integrity is 457 compromised in a disease state, leading to higher translocation rates of microbes into 458 the bloodstream. This is consistent with findings of increased intestinal permeability (i.e., 'leaky gut') in disease or even during physiological stress⁶⁷. Future studies testing this 459 460 hypothesis may consider a focus on the gut-associated bacteria that were detected in 461 our study (e.g. Bifidobacterium adolescentis, Faecalibacterium prausnitzii). Further 462 experimental investigations into the mechanisms of microbial translocation and the 463 modulatory effects of the microbiomes present at other body sites may shed light on the 464 relationship between microbial presence in blood and health status.

If we take the definition of a 'microbiome' as a microbial community whose member 465 species interact amongst themselves and with their ecological niche⁹, our findings lead 466 467 to the conclusion that there is no consistent circulating blood microbiome in healthy 468 individuals. Sporadic and transient translocation of commensals from other body sites 469 into the bloodstream (Figure 5) is the more parsimonious explanation for the 470 observation that most of the microbes detected are commensals from other body sites. 471 Furthermore, the relatively low prevalence of microbes in blood suggests rapid 472 clearance of translocated microbes rather than prolonged colonisation in blood. Based 473 on these findings, we advocate against the use of the term 'blood microbiome' or 474 'circulating microbiome', which are potentially misleading, when referring to the 475 detection of microbial DNA or of microbial cells in blood due to transient translocation 476 events.

477 Methods

478 Datasets

479 Our sequencing dataset. also known the SG10K_Health dataset as 480 (https://www.npm.sg/collaborate/partners/sg10k/), comprises of shotgun sequencing 481 libraries of DNA extracted from the whole blood or umbilical cord blood of 9,770 healthy Singaporean individuals⁶⁸ who were recruited as part of six independent cohorts. 482 483 Individuals were deemed to be healthy if they do not have any personal history of major 484 disorders such as stroke, cardiovascular diseases, cancer, diabetes and renal failure. 485 Oral health information was not collected and therefore not part of the exclusion criteria. 486 Whole blood for sequencing was collected via venipuncture only from the five adult 487 cohorts (median age=49; interguartile range=16): Health for Life in Singapore (HELIOS; 488 n=2,286), SingHealth Duke-NUS Institute of Precision Medicine (PRISM, n=1,257), Tan 489 Tock Seng Hospital Personalised Medicine Normal Controls (TTSH, n=920), Singapore Epidemiology of Eye Diseases (SEED, n=1,436)^{69,70}, and the Multi-Ethnic Cohort (MEC, 490 n=2.902⁷¹. Additionally, cord blood was collected only for the birth cohort Growing Up 491 in Singapore Towards healthy Outcomes (GUSTO; n=969)⁷². Measurement of host 492 493 phenotypes was performed on the day of blood collection, except for the GUSTO cohort 494 where measurements were taken at a later timepoint when the children were at a 495 median age of 6.1 (interquartile range=0.1). Using nearest neighbor approaches to reference genotypes⁷³, individuals were broadly categorised into four ethnic categories 496 representing distinct genetic ancestries: Chinese (59%), Malays (19%), Indians (21%) 497 498 and Others (1%). All individuals were deemed healthy at the point of recruitment if they 499 did not include any self-reported diseases in the recruitment questionnaires. All cohort 500 studies were approved by relevant institutional ethics review boards. A summary of the 501 cohort demographics and the ethics review approval reference numbers are provided in 502 Supplementary Table 1.

Additionally, we retrieved anonymised blood culture records from Singapore General Hospital, the largest tertiary hospital in Singapore. These records span the years 2011-2021 and include aerobic, anaerobic and fungal blood cultures taken from 282,576 unique patients. These blood cultures were ordered as part of routine clinical

507 management, that is, when clinically indicated for the investigation of bacteremia or 508 fungemia. Blood cultures were performed and analysed as per hospital standard 509 operating procedures. In brief, blood samples were collected aseptically and inoculated into BD[™] BACTEC[™] bottles at the bedside (BD[™] BACTEC[™] Plus Aerobic/F Culture 510 vials Plastic [catalogue number 442023] for aerobic blood culture, BD[™] BACTEC[™] 511 512 Plus Anaerobic/F Culture vials Plastic [catalogue number 442022] for anaerobic blood 513 culture and Myco/F Lytic [catalogue number 42288] for fungal blood culture). The 514 inoculated bottles were transported to the diagnostic laboratory at ambient temperature and incubated in the BD[™] BACTEC[™] FX Blood Culture System on arrival. Aerobic and 515 516 anaerobic blood culture bottles were incubated for a maximum of five days, and fungal 517 blood culture bottles were incubated for a maximum of 28 days. Blood culture bottles that were flagged positive by the BD[™] BACTEC[™] FX Blood Culture System were 518 519 inoculated onto solid media, and the resultant colonies were identified using a 520 combination of biochemical tests and matrix assisted laser desorption ionization-time of flight mass spectrometry (MALDI-TOF MS) (Bruker[®] microflex LRF). 521

522 Sample preparation and batch metadata

523 DNA from whole blood was extracted using one of six different DNA extraction kits. 524 Paired-end 151bp sequencing with an insert size of 350bp was performed up to 15-fold 525 or 30-fold coverage of the human genome. Library preparation was performed using 526 one of three library preparation kits. Sequencing was performed on the Illumina HiSeg X 527 platform with HiSeq PE Cluster Kits and HiSeq SBS Kits. The type of extraction kits and 528 library preparation kits used, and lot numbers for the SBS Kits, PE Cluster Kits, and 529 sequencing flow cells used are provided as batch metadata. All reagent kits used, the 530 number of batches and the number of samples processed per batch are provided in 531 Supplementary Table 6.

532 Data pre-processing and quality control

The bioinformatic processing steps applied to the sequencing libraries are summarised in **Figure 1a**. Read alignment of sequencing reads to the GRCh38 human reference genome was already performed as part of a separate study⁶⁸ using *BWA-MEM* $v0.7.17^{74}$. We retrieved read pairs where both members of the pair did not map to the human genome. Following which, we performed quality control of the sequencing reads. We trimmed low quality bases at the ends of reads with quality <Q10 (base quality trimming) and discarded reads with average read quality less than Q10 (read quality filter). We also discarded low complexity sequences with an average entropy less than 0.6, with a sliding window of 50 and k-mer length of five (low complexity read filter). All basic quality control steps were performed using *bbduk* from the *BBTools suite* v37.62 (sourceforge.net/projects/bbmap/).

544 Taxonomic classification of blood sequencing libraries

Taxonomic classification of non-human reads was done using *Kraken2* v2.1.2²³ with the 545 '-paired' flag. We used the *PlusPF* database (17th May 2021 release) maintained by 546 547 Langmead (https://genome-Ben 548 idx.s3.amazonaws.com/kraken/k2_pluspf_20210517.tar.gz), which includes archaeal, 549 bacterial, viral, protozoan, and fungal references. Of all non-human read pairs, 72% 550 were classified as microbial at the species level, yielding 8,890 species. Samples with 551 less than 100 microbial read pairs were removed, resulting in a final dataset comprising 552 8,892 samples, with a median microbial read-pair count of 6187.

553 To minimise noise in the taxonomic assignments, we defined a set of abundance 554 thresholds whereby species with abundance values less than or equal to these 555 thresholds (i.e., relative abundance ≤ 0.005 , read pairs assigned ≤ 10) were counted as 556 absent (set to zero read counts). We performed simulations to systematically determine 557 a relative abundance threshold that minimizes false positive species assignments. Sequencing reads were simulated using InSilicoSeg v1.5.4⁷⁵ with error models trained 558 559 on the SG10K Health sequencing libraries and processed using the same bioinformatic 560 steps as per the SG10K_Health dataset to obtain microbial taxonomic profiles. We 561 simulated 373 million reads equivalent to the median library read count of all samples, comprising reads from the GRCh38 human reference and ten microbial genomes 562 563 (Yersinia enterocolitica, Leclercia adecarboxylata, Moraxella osloensis, Streptococcus 564 pneumoniae, Pasteurella multocida, Staphylococcus epidermidis, Actinomyces 565 viscosus, Torque teno virus, Human betaherpesvirus 6A, Candida albicans) at various 566 proportions. Due to read misclassification, some of the simulated reads were

567 erroneously assigned to another species and produced false positives. A final relative 568 abundance threshold of 0.005 that delineated these false positive assignments from 569 true positives was selected (**Supplementary Figure 8**). Following the application of 570 these thresholds, the relative abundance distribution of microbial taxa classified as 571 present were distinct from the distribution for those classified as absent 572 (Supplementary Figure 9). Furthermore, the distribution of abundances for microbe-573 negative samples is centred around a relative abundance of 0.0001, i.e. at least tenfold 574 below the typical relative abundance thresholds used to determine if a taxon is present or absent (0.001-0.045¹⁴). Relative abundances were calculated by dividing the 575 576 microbial read count in a sample by the total number of microbial reads assigned to that 577 sample.

578 Decontamination filters

579 After application of the presence/absence filter, we identified and removed putative contaminants using established decontamination heuristics²⁶ that have been validated 580 in previous studies^{27,28}, prior to our downstream analyses. These rules were applied 581 582 using eight types of batch information: source cohort, DNA extraction kit type, library preparation kit type, and lot numbers for sequencing-by-synthesis kit (box 1, box 2), 583 584 paired-end cluster kit (box 1, box 2) and sequencing flow cell used. Other batch 585 information such as the pipettes and consumables used, or storage location and 586 duration were not recorded and could potentially contribute to some level of batch-587 specific contamination. However, these batches are expected to be correlated with the 588 other types of batch information available, and so the resultant contaminants could in 589 theory be accounted for using our filters. We describe the four decontamination filters 590 used, as shown in Figure 1a, in sequential order:

(1) Prevalence filter. A microbial species is considered a contaminant specific to a
 batch if it is present at greater than 25% prevalence in that batch and has greater
 than a two-fold higher prevalence than that for any other batch. Batches with less
 than 100 samples were excluded from this analysis. This filter is based on the
 principle that species which are highly prevalent in some batches but lowly

596 prevalent or absent in others are likely contaminants²⁶. We illustrate this for an 597 example species in **Supplementary Figure 10a**.

- 598 (2) Correlation filter. A microbial species is considered a contaminant if it is highly 599 correlated (Spearman's ρ >0.7) with any contaminant within the same batch, as 600 identified by the prevalence filter. This filter is based on the principle that contaminants are highly correlated within the same batch²⁶. Spearman's ρ was 601 calculated using centred log-ratio (CLR) transformed⁷⁶ microbial relative 602 abundances. CLR transformations and Spearman's p were calculated using the 603 clr function as part of the compositions package⁷⁷ and cor.test function in R. We 604 605 illustrate this within-batch correlation for an example species in **Supplementary** 606 Figure 10b.
- 607 (3) Batch filter. A non-contaminant microbial species must be detected in samples 608 processed by at least two reagent kit batches or reagent types. That is, any 609 species that is only detected in a single batch for any of the reagent kits used 610 (Supplementary Table 6) are considered contaminants. This filter is based on 611 the principle that species that can be repeatedly observed across different reagent batches are more likely to reflect genuine non-contaminant signals²⁶. 612 613 Library preparation kit type was excluded from this analysis since only three kit 614 types were used, with 86% of samples processed using one of the kits.
- (4) *Read count filter.* A microbial species is considered a sequencing or analysis
 artefact if it is not assigned at least 100 reads in at least one sample. This filter is
 based on the principle that species that are always assigned a low number of
 read pairs, never exceeding the background noise within sequencing libraries,
 are more likely to be artefactual rather than genuine signals. An example of an
 artefactual species is *Candidatus Nitrosocosmicus franklandus*, which was
 assigned at most 22 read pairs by *Kraken2* across 21 sequenced samples.
- To demonstrate the effectiveness of our decontamination filters, we additionally tested our results against the null hypothesis that the 117 microbial species retained after decontamination produced the same proportions of species classified as likely contaminants, human-associated, or that were detected in blood culture compared to if

we picked these species at random. In this analysis, we generated 1000 sets of 117 microbial species that were randomly selected from the list of species before decontamination and compared the species to the three databases as per **Figure 1b-d**. P-values were calculated by taking the proportion of random iterations that generated proportions of species classified as likely contaminants, detected in blood, or humanassociated that were as extreme or more extreme than those observed for the 117 species retained by our decontamination filters.

633 Characterisation of microbial species

634 We classified microbial species as human-associated or not based on a published hostpathogen association database⁷⁸. In this database, host-pathogen associations are 635 636 defined by the presence of at least one documented infection of the host by the pathogen³¹. For species that were not found in this database, we performed a 637 638 systematic PubMed search using the search terms: (microbial species name) AND 639 (human) AND ((infection) OR (commensal)). Similarly, species that had at least one 640 published report of human colonisation/infection were considered human-associated. 641 Additionally, we classified the potential body site origins for each microbial species 642 using the Disbiome database, which collects data and metadata of published microbiome studies in a standardised way³⁴. We extracted the information for all 643 644 microbiome database URL: experiments in the using the 'https://disbiome.ugent.be:8080/experiment' (accessed 26th April 2022). We first 645 646 extracted microbe-to-sample type mappings from this information (e.g. C. acnes \rightarrow skin 647 swab). We then manually classified each sample type into different body sites (e.g. skin 648 swab \rightarrow skin). This allowed us to generate microbe-to-body site mappings. Sample types 649 with ambiguous body site origins (e.g. abscess pus) were excluded. The range of 650 sample types within the Disbiome database used to derive the microbe-body-site mappings are provided in **Supplementary Table 7**. Finally, we classified microbial 651 652 species based on their growth requirements, with reference to a clinical microbiology 653 textbook⁷⁹. Viruses were classified as obligate intracellular. The microbiological 654 classifications for each species are provided in Supplementary Table 2.

655 Estimating coverage breadth and bacterial replication rates

656 We performed read alignment of sequencing libraries to microbial reference genomes using *Bowtie v2.4.5*⁸⁰ with default parameters. In total, we used references for 28 of the 657 117 microbial species detected in blood, comprising all bacterial species with at least 658 659 1000 Kraken2-assigned read pairs in a single sample and all viral species (n=5). For 660 each species, we aligned the microbial reads of five sample libraries with the most 661 reads assigned to that species, to the reference genome of that species. For each sample and microbial genome, the genome coverage per position was computed using 662 the *pileup* function as part of the Rsamtools v2.8.0 package⁸¹ in R. In principle, 663 recovery of a larger fraction of a microbial genome provides a higher confidence that it 664 is truly present in a sample^{24,25}. We could recover at least 10% of the microbial 665 genomes for 27/28 (96%) of the species. Since it is difficult to assess coverage breadth 666 667 for a species covered by a low number of reads, we only performed this analysis on all 668 viruses (n=5), and all bacterial species with at least 1000 Kraken2-assigned read pairs 669 (n=23), which corresponds to ~10% coverage over a typical 3Mbp bacterial genome 670 (assuming non-overlapping reads). For the replication rate analyses, PTR values were calculated using the *bPTR* function in *iRep* $v1.1.0^{21}$, which is based on the method 671 proposed by Korem et al.²². The Ori and Ter positions were determined based on the 672 673 coverage peaks and troughs (in red and blue, respectively; Figure 3). Ori and Ter 674 positions were also calculated using a cumulative GC-skew line, which is expected to be in anti-phase with the sinusoidal coverage pattern across the genome³⁹ (in green: 675 676 Figure 3).

677 Microbial networks

Microbial co-occurrence/mutual exclusion associations were computed using the SparCC algorithm⁵⁰, implemented in the SpiecEasi v1.1.2 package⁸² in *R* and the microbial networks were visualized using *Igraph* v1.2.9⁸³. We excluded the birth cohort GUSTO since it is of a different demographic that may possess a distinct set of microbial associations.

683 Detecting associations between microbial taxonomic profiles and host phenotypes

684 We tested for microbe-host phenotype associations within individual cohorts separately. 685 For the two categorical host phenotypes, genetic sex and ancestry, we tested for differences in the prevalence of each microbial species between the different categories 686 687 using a two-sided Fisher's exact test (*fisher.test* function in R). For the continuous 688 variables (age, BMI, TC, TG, SBP and DBP) we used a two-sided Mann-Whitney U test 689 (*wilcox.test* function in *R*) to test for differences in the distributions of the variables when 690 a species was present or absent. Benjamini-Hochberg multiple-testing correction was 691 applied only after consolidating the p-values from both tests and for all cohorts using the 692 *p.adjust* function in *R*. Statistical tests were only performed if a species was present in 693 at least 50 samples in total. Separately, for derived phenotypes (i.e., being elderly or 694 measures of 'poorer health'), we used the Fisher's exact test before applying Benjamini-695 Hochberg multiple-testing correction. In all cases, samples with missing host 696 phenotypes were excluded.

697 Data analysis and visualisation

All data analyses were performed using *R* v4.1.0 or using *Python* v3.9.12. Visualisations

- were performed using *ggplot* $v3.3.5^{84}$. **Figure 5** was created using BioRender.com
- 700 under an academic subscription.

701 **Data availability**

Requests for the sequencing data used in this study should be made through theNational

Precision Medicine (NPM) Programme Data Access Committee (<u>contact_npco@gis.a-</u> star.edu.sg). All other data used in our analyses are hosted on Zenodo (https://doi.org/10.5281/zenodo.7368262). The accession numbers for all genome references used are provided in **Supplementary Table 8**.

708 **Code availability**

All custom code used to perform the analyses reported here are hosted on GitHub
 (<u>https://github.com/cednotsed/blood_microbial_signatures.git</u>).

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(1) The Health for Life in Singapore (HELIOS) study at the Lee Kong Chian School of
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725 (2) The Growing up in Singapore Towards Healthy Outcomes (GUSTO) study, which is 726 jointly hosted by the National University Hospital (NUH), KK Women's and Children's 727 Hospital (KKH), the National University of Singapore (NUS) and the Singapore 728 Institute for Clinical Sciences (SICS), Agency for Science Technology and Research 729 (A*STAR) (supported by the Singapore National Research Foundation under its 730 Translational and Clinical Research (TCR) Flagship Programme and administered 731 by the Singapore Ministry of Health's National Medical Research Council (NMRC), 732 Singapore - NMRC/TCR/004-NUS/2008: NMRC/TCR/012-NUHS/2014. Additional 733 funding was provided by SICS and IAF-PP H17/01/a0/005);

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possible.

749 **Ethics declaration**

All individuals in the participating cohorts were recruited with signed informed consent from the participating individual or parent/guardian in the case of minors. All studies were approved by relevant institutional ethics review boards detailed in **Supplementary Table 1**.

754 Figure legends

Figure 1: Decontamination results. (a) Summary of pre-processing steps and filters applied to taxonomic profiles (*n*=9,770 individuals) and the number of species retained after each filter. Pie charts showing the proportion of microbial species that are (b) human-associated, (c) common sequencing contaminants, and (d) detected in blood culture records, before and after applying the decontamination filters.

760 Figure 2: Microbial signatures in human blood from healthy individuals. (a) Bar 761 chart showing the prevalence of the top 50 confidently detected microbial species in all 762 8,892 blood sequencing libraries. (b) Histogram of the number of microbial species per 763 sample. (c) Bar chart of the human body sites that the 117 confidently detected species are associated with, as determined using the Disbiome database³⁴. Species are 764 765 classified as 'multiple' if they are associated with more than one body site and classified 766 otherwise if they are only associated with a single body site. (d) Piechart showing the 767 microbiological classification of the 117 confidently detected species. (e) Bar chart 768 showing prevalence of genera in blood culture records and in the blood sequencing 769 libraries before and after decontamination.

770 Figure 3: Evidence for replicating bacteria in blood samples from healthy 771 individuals. (a) Summary statistics for samples where bacterial species were deemed to be replicating using *iRep*²¹ (i.e., peak-to-trough ratio (PTR)>1). The number of reads 772 assigned to the species by Kraken2²³, the possible body sites the species are 773 774 associated with, whether they were previously reported in published studies of 775 bacteraemia, the overall prevalence of the species across all 8,892 individuals in our 776 study and the calculated PTR values, are indicated. Coverage plots of (b) three 777 representative confidently detected species and (c) three representative contaminant 778 species, showing the expected patterns of Ori to Ter coverage skew only where 779 expected i.e. confidently detected species.

Figure 4: Microbial co-occurrence networks. (a) *SparCC*⁵⁰ co-occurrence networks computed from all samples with at least two microbial species following decontamination at different *SparCC* correlation thresholds (0.05, 0.2, 0.3). Only associations with a magnitude of *SparCC* correlation greater than the respective 784 thresholds are retained. (b) SparCC networks for individual cohorts at a correlation 785 threshold of 0.2. No co-occurrence associations were retained after taking the 786 intersection of edges between all cohort networks. For (a) and (b), each node 787 represents a single microbial species, and each edge a single association between a 788 pair of microbial species. Edge thickness is scaled by the magnitude of correlation. The 789 number of samples used to compute each network and the correlation thresholds used 790 are annotated. Positive and negative SparCC correlations are indicated in green and 791 blue respectively.

Figure 5: Potential models for microbes in blood. Our findings suggest that there is no consistent circulating blood microbiome (i.e., the blood microbiome model). The more likely model is where microbes from other body sites transiently and sporadically translocate into blood. Created with BioRender.com under an academic subscription.

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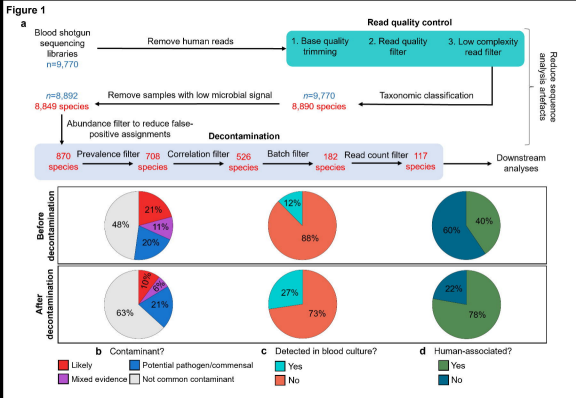
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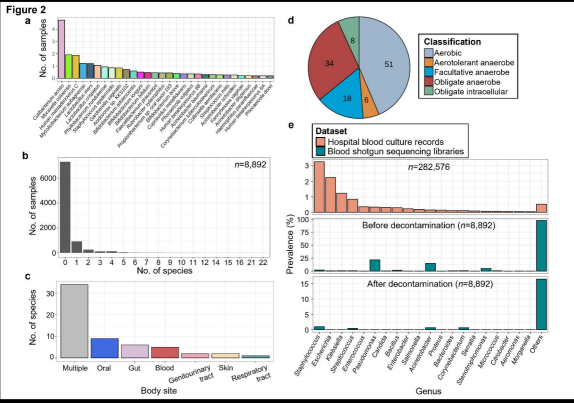
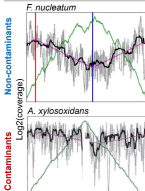


Figure 3

Sample	Species	Body site(s)	Reported in blood	No. of read pairs assigned	Overall prevalence (%)	PTR
WHB4594	Fusobacterium nucleatum	genitourinary tract, gut, mouth	4	194199	0.11	1.68
WHB9179	Neisseria subflava	gut, mouth	~	15385	0.16	1.51
WHB9179	Haemophilus parainfluenzae	gut, mouth, respiratory tract	*	12183	0.2	1.17
WHB4035	Fannyhessea vaginae	genitourinary tract	~	10395	0.24	1.88
WHB6459	Staphylococcus epidermidis	gut, mouth, respiratory tract, skin	*	9140	0.85	1.57
WHB10710	Lactobacillus crispatus	genitourinary tract, gut, mouth	~	7799	1.06	1.57
0116-0053	Acinetobacter baumannii	mouth	~	7673	0.31	1.9
WHB9179	Neisseria flavescens	mouth	×	3787	0.06	1.38
WHB9978	Rickettsia sp. Tillamook 23		×	2923	0.02	1.35
WHH1248	Moraxella osloensis	skin	~	2402	1.91	1.33
WHB9812	Corynebacterium imitans		~	1976	0.02	1.59

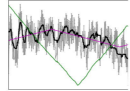
b

С



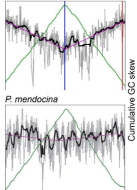
F. vaginae

A. faecalis



Genome position (complete genome)





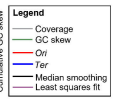


Figure 4

а

After decontamination (117 species)

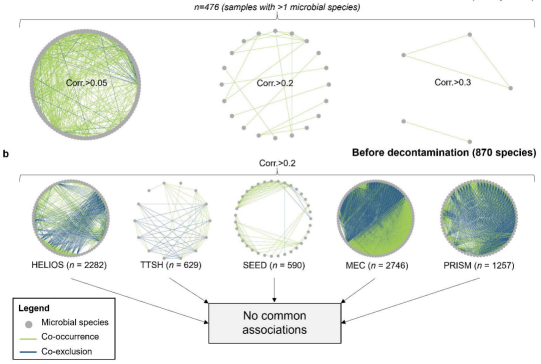


Figure 5

