Microbial Genomics

Exploring the mobilome and resistome of *Enterococcus faecium* in a One Health context across two continents

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Abstract

Enterococcus faecium is a ubiquitous opportunistic pathogen that is exhibiting increasing levels of antimicrobial resistance (AMR). Many of the genes that confer resistance and pathogenic functions are localized on mobile genetic elements (MGEs), which facilitate their transfer between lineages. Here, features including resistance determinants, virulence factors, and MGEs were profiled in a set of 1273 E. faecium genomes from two disparate geographic locations (in the UK and Canada) from a range of agricultural, clinical, and associated habitats. Neither lineages of E. faecium nor MGEs are constrained by geographic proximity, but our results show evidence of a strong association of many profiled genes and MGEs with habitat. Many features were associated with a group of clinical and municipal wastewater genomes that are likely forming a new human-associated ecotype. The evolutionary

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dynamics of *E. faecium* make it a highly versatile emerging pathogen, and its ability to acquire, transmit, and lose features presents a high risk for the emergence of new pathogenic variants and novel resistance combinations. This study provides a workflow for MGE-centric surveillance of AMR in *Enterococcus* that can be adapted to other pathogens.

1 Introduction

Enterococcus faecium is a ubiquitous, gram-positive, facultative anaerobic microorganism often isolated from a variety of natural environments including soil and water, and host-associated environments including the intestinal tract of humans and animals [1-3]. The presence of E. faecium in the intestinal tract of healthy subjects led to the belief that this microbe was an innocuous commensal, with an occasional role in opportunistic infections [4]. However, following a 1986 outbreak of vancomycin-resistant strains of Enterococcus faecalis and E. faecium in a London, UK, hospital [5], it became clear that this bacterium could cause grave illness in humans and, readily acquired antimicrobial resistance (AMR) genes through lateral gene transfer (LGT). Enterococci have the ability to share genes within an extended pool of mobile genetic elements (MGEs) [6], allowing them to serve as hubs for the transmission of AMR determinants to both gram-positive and gram-negative species [7]. Antimicrobial-resistant 10 enterococci are now a leading cause of hospital-acquired bloodstream and urinary tract infections [4]. According to 11 the World Health Organization, E. faecium is a member of a group of nosocomial pathogens called "ESKAPE" [8] 12 that have been given priority status on the list of pathogens for which new antimicrobials are urgently needed [9]. Up to the early 1990s, most nosocomial enterococcal infections were caused by E. faecalis, while E. faecium was 14 the causative agent of only about 10% of cases [10]. Over the past two decades, E. faecium infections have been constantly on the rise in the United States [11–13] and in Europe [14–18]. In 2021, Dadashi et al. reported that 16 the global prevalence of E. faecium among enterococci isolates from clinical infections was 40.6%, with 43.6% in 17 Asia, 38.0% in Europe, and 36.8% in America [19]. 18 Lebreton et al. [20, 21] described how E. faecalis and E. faecium have emerged independently through separate 19 events of LGT driven largely by MGEs. Specifically, in E. faecium, there is a deep split (about 3,000 years ago) 20 between strains commonly present in the microbiota of non-human animals (Clade A), which are the ancestors of 21 most of the current clinical, and human-adapted commensal strains (Clade B). This split coincides with the loss of 22 many genes related to the catabolism of dietary carbohydrates from Clade A strains and the MGE-mediated acquisition of genes encoding amino-carbohydrates typically involved in the glycocalyx formation during colonization of intestinal epithelial cells [20, 21]. The authors of these studies hypothesize that this difference in tropism is a re-25 flection of the preferred habitats between these two clades, with Clade B mostly community-associated and Clade A mainly with hospital-associated enterococci [22]. In studies from the United Kingdom (UK) by Gouliouris et 27 al. [23] and Alberta, Canada (AB) by Zaheer et al. [24], isolates from agricultural environments clearly separated from clinical ones constituting two distinct clades, supporting the hypothesis that they are specialized to distinct 29 ecological niches. This adaptation also reflects the nature of antimicrobials, heavy metals, and other selective pres-30 sures present in each niche. Gouliouris et al. also included a clear split between Clade A subclades, A1 and A2, 31 although we do not investigate these subclades in this study [23]. E. faecium is extremely apt at acquiring genes carried by MGEs including plasmids, genomic islands (GIs), and

prophages. In fact, the genome plasticity that renders this microorganism a formidable public-health threat relies

E. faecium mobilome & resistome

mainly on a large number of multifunctional accessory genes that can be laterally transferred between distantly related strains [6]. Plasmids are generally considered the main AMR gene-carrying MGEs in enterococci [25, 26]. Arredondo-Alonso et al. proposed that plasmids could be used to ascertain the niche specificity of E. faecium [27]. 37 However, AMR, heavy metal resistance (HMR), and virulence factor (VF) genes have also been detected in GIs 38 [28] and prophages [29]. Understanding the relative importance of habitat and geography in shaping the genome and corresponding resistance of E. faecium is vital for guiding antimicrobial use and AMR mitigation strategies. A "One Health" perspective that considers the emergence, dissemination, and transmission of resistance among human, agricultural, and environmental isolates is necessary to implement effective AMR surveillance and interventions. Existing surveil-43 lance systems rely heavily on phenotypic data and will benefit from whole-genome sequencing and analysis. The tools employed in this genomic study can connect phylogenetic, habitat, and geographic data to the prevalence of the mobilome and associated resistance and virulence genes, improving our knowledge of AMR dynamics in Enterococcus and other pathogens. Here we examine a combined set of 1273 E. faecium genomes from the United Kingdom and Alberta, Canada isolated from multiple habitats in order to determine the relationship between habitat, geography, and the occurrence and distribution of the mobilome and resistome of this opportunistic pathogen.

2 Methods

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2.1 Genome Assembly and Classification

The dataset for this study encompassed 1766 genomes: 334 *E. faecium* genomes from Alberta, Canada [24] and 1432 from the United Kingdom [23]. These genomes originated from isolates collected from five different sources: Clinical (CLIN), Agriculture (AGRI), Municipal Wastewater (WW-MUN), Agricultural Wastewater (WW-AGR), and Natural Water sources (NWS). FASTQ files for the AB genomes were retrieved from the Sequence Read Archive (SRA) of the National Center for Biotechnology Information (BioProject PRJNA604849), and UK genomes from the European Nucleotide Archive (Table S1). The quality of the FASTQ files was determined using FastQC v0.11.8 [30]; reads were trimmed using fastp v0.23.2 and assembled with Unicycler v0.4.8 [31] using default parameters. The quality of the assemblies was assessed using Quast v5.0.2 [32] (Figure 1A). An NG50 cutoff of 30,000 bp was used to remove low-quality genomes from subsequent analysis.

[Figure 1 about here.]

2.2 Pangenome and Generation and Visualization of Core-Genome Phylogenies

We constructed a core-genome phylogenetic tree for all of the *E. faecium* genomes. To do so, genomes were annotated using Prokka version 1.14.6 [33] followed by Roary v3.13.0 to construct a core-genome alignment [34]. As a reference, the genome of *E. faecium* DO ASM17439v2 was included and *E. hirae* ATCC9790 ASM27140v2 was used as an outgroup genome in the alignment. Using SNP-sites v2.5.1 a single-nucleotide polymorphism (SNP) alignment was produced and unambiguous nucleotide frequencies were counted [35]. The resultant SNP alignment and core-genome base frequencies were then used to generate a maximum-likelihood phylogenetic tree using IQTree v.2.1.4-beta [36] with the general time reversible model with invariable site plus discrete gamma model (Figure 1C). One thousand ultrafast bootstrap replicates and 1000 Shimodaira Hasegawa-like approximate likelihood ratio tests (SH-aLRT) were performed [37]. The phylogeny was then visualized using GrapeTree v.1.5.0 [38].

Genomes were assigned to "Clade A" and "Clade B" based on groEL gene sequences as described in Hung et al. 73 [39]. Briefly, groEL sequences were extracted and sequences corresponding to "Clade A" strains E. faecium strain 74 V68, accession MH109129 and "Clade B" E. faecium strain 81, accession MH109127 were added as references. 75 Sequences were aligned using MAFFT v7.490 with default parameters and a maximum-likelihood tree was created 76 with IQTree v2.1.4 using as a model unequal purine/pyrimidine rates with empirical base frequencies and a pro-77 portion of invariable sites (TN+F+I). Our assignment of the genomes to the two categories was then guided by the topology of this tree. Habitats, countries of origin, and "Clades" were all mapped onto the core genes-based reference tree. "Clade B" was paraphyletic in the reconstructed tree, as a consequence we refer to these two categories as "Type A" for "Clade A" and "Type B" for "Clade B". 81

Prediction of Resistance Genes and Virulence Factors 82

The assembled contigs were used to detect AMR genes, HMR genes, and VFs (collectively referred to as target 83 genes) using specific databases for each gene type. AMR genes were detected with the Resistance Gene Identifier 84 (RGI) (v5.1.0) for the prediction of AMR genes based on homology and SNP models from the Comprehensive Antimicrobial Resistance Database (CARD) v3.1.0 [40]. RGI stratifies matches into three categories using a 86 curated BLAST bitscore cutoff that reflects known variation within a gene. Matches that score below a designated, target-specific cutoff are assigned to the "Loose" category, while matches above this cutoff are assigned to the 88 "Strict" category. Exact matches are assigned to the "Perfect" category. We limited our analysis to the Strict and 89 Perfect matches. To identify HMR genes and VFs, open reading frames (ORFs) present in the assemblies were first 90 annotated using Prodigal v2.6.3 [41]. Subsequently, homology search with an initial E-value threshold of 10^{-20} 91 against two databases, VFDB [42] and BacMet v2.0 [43] was conducted using DIAMOND-BLASTX v0.9.36 [44]. 92 Results were then filtered by percent identity (60%) and match coverage (60%). Clusters of highly similar genes 93 (>95%) were identified using vsearch v2.17.1 [45] (Figure 1B).

2.4 Prediction of Mobile Genetic Elements

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To predict the presence of plasmids in our short-read assemblies, the genomes were analyzed with MOB-suite 96 3.0.0 [46] with the v2020-05-05 database. The MOB-suite pipeline scans input assemblies for contigs containing 97 plasmid-related genes (e.g. relaxases and replicases) and repetitive regions, thereby identifying putative plasmid scaffolds. These scaffolds were compared against a database of mobility clusters (MOB-clusters) comprising preclustered reference plasmids. The putative plasmids were assigned to MOB-clusters by identifying the minimum 100 Mash distance [47] to a reference plasmid. The output consisted of a single contig sequence per MOB-cluster and 101 an annotation of their host-range predictions, mobility predictions, and assignment to a replicase (rep) gene cluster. 102 Contigs larger than 1 kb were examined for the presence of GIs with IslandCompare [48] using the reference 103 genome E. faecium DO ASM17439v2. IslandCompare uses the reference genome to generate an alignment-based 104 concatenated genome from each submitted draft genome. GIs are predicted by two underlying tools, IslandPath-105 DIMOB [49] and Sigi-HMM [50] that identify regions of the genome with anomalous dinucleotide bias (and at 106 least one mobility gene) and differential codon usage, respectively. IslandCompare also incorporates additional 107 functionality for ensuring the consistency of GI predictions across genomes in multi-genome datasets and clusters 108 the predicted GIs. Following analysis, any GI predictions that corresponded to the region of the genome that could 109 not be aligned to the reference genome were excluded. For GIs present in a relatively large proportion of genomes 110 (>10%), a manual assessment was performed for genes annotated in those GIs. One GI that consisted mainly of genes involved in replication was present in nearly all genomes (1208/1273) and excluded from the analysis. Concatenated genome files generated by IslandCompare were also used for prophage prediction via DBSCAN-113 SWA [51]. DBSCAN-SWA combines density-based spatial clustering of applications with noise (DBSCAN) and

a sliding window algorithm (SWA) for prophage detection (Figure 1B).

The taxonomic distribution of predicted genes and MGEs was assessed through a homology search. A reference database of predicted proteins was constructed from 20,100 complete bacterial genomes downloaded from RefSeq on December 24, 2021. DNA sequences of plasmid-associated contigs, GIs, and prophages were compared to this database using DIAMOND-BLASTX version 2.0.13 with a maximum e-value of 10^{-50} , 90% percent identity or greater, and minimum subject coverage of 90%. Only matches with a score of 95% or greater relative to the best match were retained. Final filtering of results used a minimum percentage identity threshold of 99%. The taxonomic distribution of matches was extracted from the resulting set of hits.

Sets of target genes that mapped to a given MGE were considered to be co-localized to that MGE. Co-localizations between predicted AMR, VF, and HMR genes were identified using python and summarized by gene cluster. Genes that did not localize to any MGE were treated as chromosomal.

2.5 Analysis of Feature Abundance

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We tested the hypothesis that E. faecium isolates from different sampling environments have differential abun-127 dances of AMR and HMR determinants, VFs, and MGEs (hereafter referred to collectively as "features"). A 128 three-way factorial ANOVA was performed to determine the extent that features were associated with habitat, 129 geographic location, type, and the interactions of these categories. 130 To perform ANOVA for each feature, the mean number of unique features of each type per genome was calculated. 131 Genomes originating from NWS and WW-AGR isolates were available only in the AB isolates so these genomes 132 were omitted. This resulted in 1203 genomes divided over 12 treatment groups (3 habitats, 2 geographic locations, 133 2 types). To account for the unequal number of isolates in each treatment group, we used unweighted marginal 134 mean sum of squares (SS), or type III SS, to calculate our ANOVA statistics [52]. To investigate feature frequency 135 variance in the omitted environments, a two-way ANOVA was performed using the 303 AB genomes with 10 treatment groups (5 habitats and 2 types). Where categories were found to be significant at $\alpha = 0.05$, pairwise 137 Tukey's HSD post-hoc significance testing was performed on within-category group means (Table S2).

2.6 Coevolutionary Associations of Target Genes and Mobile Genetic Elements

We investigated the correlation across and within features using phylogenetic profiles. A phylogenetically informed maximum-likelihood approach was used to predict pairs of features with coordinated patterns of gain and loss, 141 using the BayesTraits version 3.0 [53]. To characterize the associations of predicted genes and MGEs across the 142 tree, BayesTraits constructs two continuous-time Markov models for each gene/gene, gene/MGE, and MGE/MGE 143 pair based on their patterns of presence and absence: one model expresses the likelihood that the pair evolves in a correlated way, and the other the likelihood that they are gained and lost independently [54, 55]. The ratio of 145 these two likelihoods was used to generate a p-value that reflects the statistical significance of their association. 146 Only pairs in which both features occurred in at least 3 genomes were considered. Because directionality of the 147 association is not determined by BayesTraits, we infer this by referring to the distribution of the genes across the 148 phylogenetic tree, habitat, clade and geography using the presence and absence of the features. 149 The likelihood ratios and p-values corresponding to the gene-gene and gene-environment relationships were rep-

2 3 Results

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resented as network diagrams using Cytoscape (v3.8.2).

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Genome Assembly and Distribution 3.1

Out of 1766 sequenced genome initially selected for this study, 1273 genome assemblies (i.e., 303 from AB and 154 970 from UK) that passed quality-control measures (NG50 \geq 30,000) were selected for further analysis. The 155 NG50 values of accepted genomes ranged from 30,088 to 467,170 bp, with a mean of 148 contigs (range: 18 to 156 304). Assembly sizes varied from 2,373,576 bp to 3,301,308 bp with a mean value of 2,830,264 bp (Figure 2A) 157 and with a mean GC content of 37.8% (range from 37.25% to 38.49%). The pangenome of the 1273 E. faecium 158 genomes consisted of 26,246 genes (Table S3), including 1101 (4.2%) present in 99-100% of the genomes (the 159 core genome); 212 (0.8%) in 95-99% of genomes (the "soft core" genome); 2207 genes (8.4%) in the "shell" 160 genome (15-95% of genomes), and 22,726 genes (86.6%) in the "cloud" (less than 15% of genomes). A total of 161 382 genes belonging to the AMR (n = 82), HMR (n = 32), and/or VF (n = 268) classes were identified at different 162 frequencies throughout the analyzed genomes (Figure 2B-H). Our analysis of the predicted mobilome identified 1131 sets of MGEs, all of which were part of the accessory 164 genome except for Streptococcus phages predicted by DBSCAN-SWA in 1272 genomes. MOB-suite assigned plasmid-associated contigs to 263 different clusters. Of these, 88 were assigned to reference plasmid clusters and 166 175 were categorized as novel with 144 (83.2%) of these associated with UK isolates. Given that many of these novel clusters may be misclassified chromosomal fragments, we did not include these in our subsequent analysis. 168 The 7805 GIs were predicted to constitute 824 groups of GIs with only 15 GI clusters present in more than 10% of 169 the isolates, 477 of them being unique to single genomes. 170

[Figure 2 about here.]

Distribution of the Resistome, Virulence Factors, and MGEs by Type Assignment, Geographical Origin, and Habitat

The factorial three-way ANOVA compared the main effects of habitat, geography, and type as well as their interaction on the frequency of target features (Table 1). For all features, marginal effects at a threshold of $\alpha = 0.05$ 175 were observed for at least one of the categories (geography, habitat, and type). However, in each case at least 176 one significant interaction effect was also observed which suggests conditionally dependent patterns of association and the need to interpret main effects with caution. Habitat showed the strongest marginal effects for all features 178 except HMR, with $p < 10^{-7}$ in all other cases. Geography was associated with HMR, VFs, and plasmid clusters, but post-hoc significance testing strongly suggested that the HMR relationship $(p = 4.16 \times 10^{-5})$ is an artefact 180 of interaction effects driven by Type A agricultural isolates from the UK (Table S2B). AMR and phage showed 181 significant associations with the Type A / Type B split, although the latter $(p = 4.69 \times 10^{-2})$ was not supported 182 by post-hoc testing. In the two-way ANOVA using Alberta genomes, results were consistent with the three-way 183 ANOVA with the exception of HMR determinant frequencies, which were found to be significantly affected by 184 habitat at p<0.05 (Table 2). Inspection of the HMR frequency post-hoc significance tests indicate that this effect is 185 likely caused by habitat × type interactions, with NW and CLIN isolates showing significant differences between 186 Type A and Type B HMR determinant frequencies (Table S2B). 187 A total of 19,599 AMR genes was predicted in the 1273 isolates: removal of duplicates yielded 16,898 genes with 188 a mean occurrence of 13.27 per genome. We observed a large discrepancy between types A (14.3±5.3 predicted 189 AMR genes per genome) and B (5.9±1.6 AMR genes per genome) (Table S4; Figure 3). Plasmids and GIs showed discrepancies as well, with 6.2±3.0 mean occurrences of distinct plasmid clusters in Type A and 2.0±1.3 in Type 191 B. GIs were found 7802 times, with 6.5±2.8 of mean occurrences in Type A and 3.7±1.6 in Type B. Conversely, HMRs, VFs, and phages showed similar distributions between types even when genomes were partitioned by 193 geography and habitat.

For AMR genes, plasmids, and GIs, the difference in distribution between types A and B was still observed 195 after considering the geographical origin of the samples (Table S4). However, we detected a large variation in 196 the distribution of the AMR genes, plasmids, and GIs between UK and AB samples isolated from municipal 197 wastewater. Specifically, the UK Type A genomes had 14.3±6.4 AMR genes, 5.5±3.0 plasmids, and 6.3±2.3 GIs 198 compared to the AB Type A genomes with 6.44±1.9 AMR genes, 2.7±1.9 plasmids, and 3.2±1.5 GIs (Table S4). 199 Differences in the distribution of AMR genes, plasmids, and GIs between types A and B were also detected across 200 habitats. Specifically, Type A CLIN samples from UK and AB had the highest mean of AMR genes per genome 201 of all the isolates (15.9±4.5 and 15.0±4.0 respectively) with corresponding variation in the relative distributions of 202 plasmids and GIs. 203

[Table 1 about here.]

Table 2 about here.]

[Figure 3 about here.]

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208 3.3 Phylogenetic associations of target genes and MGEs

or type (Figure S2B).

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The phylogenetic tree inferred from the 1273 genomes separates isolates by type (Figure 4). Over half of Type A 209 was comprised of almost entirely CLIN and WW-MUN isolates (mainly consisting of UK isolates), but two additional subclades possessed isolates from all habitats. Type B encompassed isolates collected from both countries 211 and all five habitats, with environmental and geographical categories constituting monophyletic groups. 212 Most features were irregularly distributed across the phylogenetic tree (Figures S1A-F). Relatively few AMR genes were present in Type B compared to Type A, particularly Type A CLIN. Only eatAv, a variant of eatA 214 that confers resistance to multiple antibiotics, was present in the majority of non-clinical subtrees and absent 215 from most CLIN and WW-MUN isolates. Both the vanA and vanB operons were restricted to the CLIN / WW-216 MUN subtree. Two sets of HMR genes showed strong negative associations, largely mapping onto the Type A / 217 Type B division; the corresponding genes had the same names (chtR, ruvB, copB, and chtS). These clusters may 218 have been divided because of sequence dissimilarity rather than functional differences. Although some VFs were 219 preferentially associated with Type A or Type B isolates, very few were exclusively confined to one or the other. 220 Some plasmids and GIs were over-represented in the CLIN / WW-MUN Type A, and less frequently associated 221 with the non-clinical isolates and Type B. 222 Significant (p-values < 0.01) associations were observed for AMR features with geography (18.0%), type (12.5%), 223 and habitat (11.5%; $p < 10^{-10}$) (Figure S2A). Overall, the strongest positive and negative associations were seen 224 for the CLIN and AGRI habitats, respectively. The vanA genes and genes conferring resistance to aminoglycosides, macrolides, tetracycline, trimethoprim, streptothricin all met this threshold (Table S4). As an example, vanA was 226 prevalent in CLIN genomes but almost never present in AGRI genomes. VFs exhibited associations with habitat, geography and type (Figure S2A, S2C). HMR genes associated most strongly with geographic origin than habitat 228

Physical location of the resistome genes and virulence factors in the genome 3.4

We next examined the localization of the 72,786 predicted AMR, HMR, and VF genes in the 1273 genomes. A 231 total of 18,518 (25.4%) predicted genes mapped to one or more MGEs, with 20.6% mapping to plasmids, 5.1% to GIs, and 2.2% to prophages. There were a total of 102 MGEs with colocalized AMR and VF genes and a total of 233 7 MGEs with both AMR and HMR genes (Table S5). The dominant plasmid clusters AB369, AC731, and AB173 were identified in 400, 678, and 187 genomes, respec-235 tively. Common AMR genes in these plasmids included the vanA operon, sat-4, ermB, and the aminoglycoside 236 resistance genes aac(6')-Ie-aph(2'')-Ia, aad(6), and aph(3')-IIIa. Six VFs associated with the PilA pilus struc-237 ture were also frequently found on these plasmids. However, the relative numbers of these genes differed among 238 predicted plasmid clusters, with some containing solely AMR or VF genes. AB756, the fourth most-abundant plasmid cluster, also had substantial numbers of the three aminoglycoside resistance genes mentioned above, ermB, 240 and sat-4, along with lsaE and the tetracycline resistance genes tetL, tetM, and tet(W/N/W). AH273, the seventh most-abundant plasmid cluster, contained the HMR genes encoding the UDP-glucose 4-epimerase galE and the 242 copper-translocating ATPase *copA*. 243 The most common GIs mainly housed tetracycline- and vancomycin-resistance genes. GI 14 (identified in 186 244 genomes) was associated with dfrG, tet(W/N/W), and tetM; GI 8 (identified in 310 genomes) with dfrF; and GI 34 245 (identified in 51 genomes) with the vanB suite of genes. GI 69 was found less frequently than the predominant GIs, 246 being present in 18 genomes and containing ant(9)-Ia, efrA, and ermA. Other predicted GIs had aminoglycoside-247 resistance genes, ermB (a macrolide resistance gene), and sat-4. Predicted VFs in GIs included bsh (VFC36), a bile salt hydrolase, ssaB (VFC39), a Manganese/Zinc ABC transporter substrate-binding lipoprotein precursor, 249 fss3 (VFC42) a fibrinogen binding protein, ecbA (VFC84 and VFC86) a collagen binding protein, and multiple genes involved in capsule formation (epsE (VFC48), gmd (VCF51), cps2K (VCF52)). 251 Most prophage-associated genes mapped to either annotated "Streptococcus phage" (1366 / 1578) or "Enterococ-252 cus phage" (100 / 1578). Genes that mapped to the predicted Streptococcus phages were similar to those observed 253 in the plasmids and GIs, including those associated with aminoglycoside, erythromycin (ermB), streptothricin 254 (sat-4), and tetracycline resistance. While some common VFs were unique (eg. lap (VFC14), an alcohol dehydro-255 genase involved in adhesion to the host cells), others were similar to those found to be localized to GIs including 256 bsh (VFC22 and VFC36), fss3 (VCF42), and ecbA (VFC84). Predicted Enterococcus phages had several instances of dfrA42, bsh (VFC22). 258 Many of the genes noted above showed biased associations with the corresponding MGEs. For example, over 93% 259 of all vanA and tetracycline-resistance genes mapped to predicted plasmids, as were over 80% of the macrolide 260 and streptothricin resistance genes ermB and sat-4, respectively. However, the gene-centric view also identified 261 rare genes with strong biases including catA8, lnuB, ermT, and chloramphenicol acetyltransferases. Over 75% of 262 vanB and dfrF genes were associated with GIs; other AMR genes with strong biases included optrA (65%) and 263 lnuG (61.9%). The genes most strongly associated with prophages were the collagen-binding MSCRAMM gene (86.5% of genes), tet(W/N/W) and tetM (61.7% and 40.4% respectively) and dfrG (33.5%). However, all of these 265 genes were also strongly associated with plasmids, GIs, or both. No prophage-specific genes were identified.

3.5 **Phylogenetic Distribution of MGEs** 267

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We examined the predicted host-range distributions of all features via direct homology search and the host-range prediction feature of MOB-suite, MOB-suite predicted a total of 7232 putative plasmids which grouped into 88 269 unique MOB-clusters. A total of 4470 plasmid-associated contigs were predicted to be non-mobilizable, while 1782 were predicted to be mobilizable and 980 were predicted to be conjugative. The majority of plasmidassociated contigs (n=5652) were predicted to be specific to Enterococcus. Relatively few plasmid clusters had

very narrow or very wide distributions outside of *Enterococcus*: a total of 358 clusters were associated with some other single genus, while 347 clusters were predicted to occur in phyla other than Firmicutes.

Although plasmid host range was often narrow, individual plasmid-associated genes were often strikingly similar

(>99% identity over at least 90% of the subject sequence) to genes from other taxonomic groups, even at the phylum level. Of the contigs annotated as cluster AC731 across 678 genomes (36 mobilizable), 206 had at least one aminoglycoside-resistance gene with a high-stringency match outside of *Enterococcus*, frequently to *Staphylococcus*, *Streptococcus*, *Campylobacter*, and members of the family Enterobacteriaceae, while 112 matched at least one gene in the *vanA* group, often across multiple phyla. Not all plasmids showed evidence of recent LGT. All non-hypothetical genes in plasmid cluster AH273, including a range of metal-associated transporters, had no stringent matches outside of *Enterococcus*. All the annotated members of this plasmid cluster were predicted by MOB-suite to be non-mobilizable.

Most GIs were dominated by integrases and other signatures of MGEs, and poorly annotated genes with products that include general ABC transporters. The most common GI was found in 484 genomes; over 90% of these GIs had a suite of genes found in multiple phyla and included toxin-antitoxin and pilin genes, peptidases, and annotated ABC transporter permeases. GI 26, found in 88 genomes, had a very high incidence of multiphylum tetM genes. The vanB genes found in GI 34 were nearly identical to those in other Firmicutes such as Staphylococcus, and occasionally in members of Enterobacteriaceae such as Klebsiella. Similarly, prophage genes with stringent matches to groups outside Enterococcus were predominantly associated with mobility and included endonucleases, integrases, and transposases. However, over 500 genomes had genes annotated as ermB, with stringent matches to other phyla. A similar number of genomes had at least one aminoglycoside-resistance gene, the most common being ant(6)-Ia.. Other common genes involved in transcription included transcription factors and 500 instances of the σ^{70} subunit of RNA polymerase.

3.6 Distribution and Associations of Vancomycin-Resistance Genes

Both the *vanA* and *vanB* gene clusters showed a strong association with CLIN and WW-MUN but variable distribution and association with MGEs (Figure 5). The *vanA* gene clusters were found to be disproportionately associated with plasmids in both the AB and UK datasets (Table S5). Overall, 458/474 *vanA* genes were found to colocalize to and associate with several plasmids of which AB369 and AC731 were the most abundant. The *vanA* genes were primarily identified in CLIN (100/474 AB and 270/474 UK) and WW-MUN samples from the UK (103/474), with only 1 isolate from UK AGRI sources.

Conversely, all of the vanB gene clusters were found in UK genomes and, in 51/57 genomes, these genes colocalized to GIs (Figure 5). For the remaining 6/57 genomes, the vanB was in the unaligned portion of the genome that was not included in the GI analysis. All of the vanB genes were predicted to fall on a single GI cluster (except for one representative that has a large insertion in the middle of the GI) and contained a Tn916 transposase. The positive association of vanB to GI cluster 34 was also supported ($p < 10^{-16}$). The majority (39/57) of vanB genes were in CLIN isolates with WW-MUN isolates composing the remaining 18/57 instances.

[Figure 5 about here.]

3.7 Other Notable AMR Gene Classes

In addition to *msrC* (Figure 6), a species-specific gene of *E. faecium* that confers low-level intrinsic resistance to macrolide and streptogramin B compounds [56], multiple macrolide-resistance genes were identified. The most abundant were *ermB*, (835/1271; 66%), *ermT* (14.7%), and *ermA* (6.7%) (Figure S3). Some of these genes showed a bias for the CLIN (*ermT*) or AGRI and WW-AGR (*ermA*) environments, while *ermB* was prevalent in all

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environments except NWS and WW-MUN from AB. The majority of these genes were localized on plasmids, with ermA identified on AB369 (44/47;94%) ermB mostly associated with AC731 (146/662; 22%) and AB369 (130/662; 20%). In AGRI genomes, ermB was commonly associated with AC730 (37/76; 49%) and AB756 (15/76; 20%). Plasmid clusters AC731 and AB369 were also associated with vanA and ermB. The ermA gene, was exclusively associated with Type A and the AGRI isolates, while ermB was positively associated with Type A, as well as with UK, AGRI, NWS, CLIN and WW-MUN isolates. The ermT gene demonstrated a positive association with Type A, UK, and CLIN isolates and a negative association with AGRI and NWS isolates.

[Figure 6 about here.]

Tetracycline-resistance genes were common, often plasmid-associated, in the analyzed genomes (Figure S4). tetM

(783/1273; 61.5%) was most prevalent, followed by tet(45) (455/1273; 35.7%) in CLIN genomes from the UK. 323 Other tetracycline genes including tet(W/N/W) (236/1273; 18.5%) and tetU (168/1273; 13.2%) were found at 324 higher rates in WW-MUN and CLIN genomes from the UK; tetL (90/1273; 7.1%) was found mostly in agriculture 325 and WW-AGR with higher levels in AGRI genomes in the UK; tetS (37/1273; 2.9%) was found primarily in 326 WW-MUN and NWS. tet(40) and tetO were both found in UK WW-MUN. 327 The aminoglycoside resistance gene aac6-li, responsible for intrinsic aminoglycoside resistance in this species 328 [57], was found in the majority of genomes (1271/1273; 99.8%) (Figure 6). Interestingly, a three-gene lo-329 cus aad(6)-sat4-aph(3')-IIIa (595/1273; 46.7%) conferring resistance to aminoglycosides and streptothricin, was 330 present in AGRI, CLIN, and WW-MUN isolates at higher prevalence in UK than AB. A bi-functional protein-331 coding gene aac(6')-le-aph(2")-la (468/1273; 36.8%) was also found at higher abundance in the UK CLIN and 332 WW-MUN isolates compared to AB. Both ant(6)-Ia (166/1273; 13.0%) and ant(9)-Ia (82/1273; 6.4%) exhibited 333 a higher prevalence in AGRI isolates than isolates from other habitats. A small number of CLIN and WW-MUN isolates from UK harboured ant(9)-Ia, while corresponding Alberta isolates lacked this gene. Other rarely detected 335 aminoglycoside-resistance genes included apmA (11/156 UK AGRI genomes), aph2-IVa (2/270 UK WW-MUN genomes), aac(6)-Iak (2/544 UK CLIN and 2/270 UK WW-MUN genomes), aac(6)-II (1/544 UK CLIN genomes), 337 aph2-Ie (1/270 UK WW-MUN genomes), and ant(4)-Ib (2/51 AB AGRI genomes).

3.8 Heavy Metal and Biocide Resistance Genes

Genes related to copper resistance and transport comprised a large portion of the predicted HMR genes comprising 12/32 gene clusters and 2542/7914 predicted genes. The most common were two *copB* clusters (BacMet clusters 3 and 9) and a *copA* cluster (BacMet cluster 5). While cluster 3 *copB* were spread across habitats and countries, the cluster 9 *copB* were most predominant in AB WW-MUN, AGRI, NWS, and WW-AGR isolates and were underrepresented in CLIN and all UK isolates (Figure S5). The *copB* genes chromsomal except for 2/1155 cluster 3 and 3/137 cluster 9 *copB* representatives associated with GIs and plasmids, respectively. The *copA* cluster 5 was also prevalent in all environments, although more so in CLIN isolates and UK WW-MUN. Most *copA* (549/917) were also predicted to be associated with the chromosome while 366/917 cases were predicted to be localized to plasmid AH273. Another cluster of copper-resistance genes primarily associated with the agricultural environment in the UK included the genes *mco* (BacMet cluster 14), *tcrB* (BacMet cluster 15), *tcrA* (BacMet cluster 16), and *copY/tcrY* (BacMet cluster 17). These genes all showed strong association with one another as well as the plasmid AC726. There were five instances identified where copper genes and mercury-resistance genes were colocalized on a single MGE, with plasmid cluster AD908 involved in three instances. All of these cases were identified in AGRI genomes from the UK.

3.9 Virulence Factors

Both the ssaB and fss3 genes have been shown to play a role in adhesion. These genes were primarily identified 355 in CLIN genomes from both countries and WW-MUN genomes from the UK. ssaB was either localized on the 356 chromosome (364/627; 42%) or on GIs (263/627; 58%) (Figure S6). In particular, 59% (215/263) of the genes were associated with a single GI. An additional 36% (95/263) were identified on GI 23, which also carried fss3 358 in 85% (81/95) of cases. GI 23 was primarily identified in the UK dataset (93/95). fss3 was found in 63% of UK CLIN genomes but only 14% of AB CLIN genomes. Among the remaining fss3 genes not present on GI 23, 37% 360 (181/483) were present on the chromosome, 27% (132/483) on other GIs, 18% (88/483) on regions predicted to 361 be both a GI and a prophage, and one predicted to be on a non-GI-associated prophage. Both ssaB and fss3 were 362 strongly correlated to each other, clinical-related AMR genes, and MGEs. A total of 25 VFDB gene clusters were predicted to be pilin genes common to all habitats. The CLIN genomes 364 had the highest prevalence of these genes and the proportion of UK AGRI isolates with each of these genes was 365 higher than the AB AGRI isolates (Figure S7). Plasmids were the most common localization site of pilA (929/958; 97.0%), pilE (1023/1060; 96.5%), and pilF (881/909; 96.9%), with the most common colocalized plasmids being 367 AD907 and AC731. The chromosomal pilB gene had a similar prevalence across all datasets and was found on the chromosome. 369

70 4 Discussion

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E. faecium, commonly a minor harmless component of the enteric microbiota, has become a leading causative agent of healthcare-associated infections since the early 1980s [58, 59]. The combination of approaches we applied here can augment phenotype-based "One Health" genomic-surveillance workflows for *E. faecium* and other bacterial pathogens. Using the whole-genome approach the potential for gene transfer and the distributions of genes within and between habitats can be defined.

4.1 Genomic Epidemiology Suggests Strong Habitat Associations But Few Barriers to Transmission

The AB and UK genomes are distributed on the core genome-based phylogenetic tree independent of type or 378 habitat. This indicates that geographic separation has very little impact on the population structure of E. faecium. 379 There was no significant difference in the abundance of MGEs between types or geographic origin (Table 1, 2) 380 which appears to contradict the findings of previous studies that detected significantly higher numbers of MGEs 381 in Clade A than in Clade B [21, 27, 60]. The phylogenetic approach of BayesTraits suggested that the observed 382 differences were driven by increased MGE abundance in CLIN and WW-MUN isolates (Figure 3). The lack of 383 observed association with type suggests that MGEs can move between phylogenetically distant E. faecium isolates and that MGEs within populations are similar across continental barriers. 385 The global patterns of association seen among AMR and VF genes mirrored those of MGEs, with habitat gener-386 ating the smallest p-values in the ANOVA tests (Tables 1 and 2). Geographic origin showed strong associations 387 only in interactions with habitat for AMR genes and MGEs, likely as a result of intensive use of antimicrobials 388 which can result in the emergence of multidrug-resistant E. faecium. E. faecium may acquire a multi-drug resistance plasmid in the clinic but lose it upon introduction to another habitat [61]. This process can occur rapidly and 390 repeatedly, with habitat serving only as an ecological filter rather than a barrier to transmission. Analysis of the composition of features by groupings of type, habitat, and geography supports the division of features based on 392 type and habitat.

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Antimicrobial use may account for some of the variance in the distribution of AMR genes. The few differences we observe may be due to different host origins in the two locations: the UK AGRI genomes include isolates from chicken, turkey, pig, beef, and dairy cattle while the AB AGRI samples only originated from beef cattle production sources. Second, antimicrobial use in agriculture differs between Alberta and the UK. For example, the strongest associations of AMR genes with location were among the aminoglycoside family (e.g., aad(6)), which are used in the UK but not Alberta [23, 24]. However, our information about antibiotic use is incomplete, especially for chicken and turkey [62, 63].

4.2 A Highly Diversified and Dynamic Accessory Genome Allows the Rapid Acquisition of Resistance and Other Traits

Although some of the 88 plasmid clusters and 824 GI sets identified are likely very similar, they are nonetheless different enough in sequence and gene content to be differentiated. The number of unique prophages is likely an underestimate due to the grouping of some phages by name (e.g., 'Streptococcus phage') rather than by homology. Key resistance genes were observed in association with many predicted plasmid clusters. For example, 14 observed clusters had at least one vanA gene. Similarly, multiple clusters showed associations with tetracycline, aminoglycoside, and macrolide-resistance genes. The dispersion of genes across multiple clusters likely diminishes the strength of observed associations, such that we may not identify all MGEs that associate with specific genes. Additionally, aggregation of plasmids and GIs into broader clusters (such as plasmid incompatibility groups) and consequently fewer classes might improve our ability to detect important associations. The increased prevalence of many classes of resistance genes and MGEs in the clinical environment suggests that this may be the key focal point of plasmid evolution, with novel combinations forming through recombination events. The lack of geographic barriers, and the apparent ability of *E. faecium* to move between habitats, suggests that new MGEs will not be limited in their ability to disperse. The clear ability of *E. faecium* to acquire and disseminate new genes from distantly related species creates additional risks for the emergence of new combinations of AMR determinants.

The groEL-based clade-mapping on our reference tree supports a monophyletic Clade A, as described and proposed

4.3 An Emerging Clade of Pathogenic E. faecium

by Palmer et al. [64], but a paraphyletic "Clade B", which has led to our designation of these two groups as 419 "types" rather than clades. Earlier work proposed a division of Clade A into a pathogenic subclade A1, and a commensal group specific to non-human animals capable of causing sporadic infections, subclade A2 [21]. 421 However, consistent with recent observations by others, we observed a large group comprised of CLIN and WW-MUN isolates that branched within the larger grouping that included genomes isolated from all habitats (Type A); 423 this tree topology has been referred to as a clonal expansion [65, 66]. Importantly, our phylogenetic tree is based on the core genome of our isolates (n=1273) and, therefore, its topology should be less affected by LGT events 425 than a gene-focused or whole-genome tree and thus should better reflect the structure and evolutionary trajectory 426 of *E. faecium* populations. 427 Our core-genome tree suggests that there may be a sub-population adapting to the clinical niche beyond the simple 428 and more plastic advantage provided by MGEs. This is consistent with the observation from Leclerque et al. 429 that members of this clinical expansion are out-competed by other E. faecium clones in natural environments [67] 430 and by Montealegre et al. that strains from Type B have higher fitness than Type A in the absence of antibiotics [68]. These findings, together with our results, support the hypothesis proposed by Prieto et al. that this clinical 432 clonal expansion is so specialized to its environment that its strains are unfit to populate other environments and niches. This population could get more isolated and drift away from the rest of the species [66]. If the clinical-434 associated group we detected in our dataset has some level of diversification, it may satisfy the Cohan & Perry

(2007) definition of an ecotype, with lineage cohesiveness conferred by genetic similarities and distinguished by 436 unique adaptations (e.g., AMR genes) and ecological capabilities. Cohan & Perry hypothesized that periodic 437 selective sweeps reduce genetic variation between the genomes of organisms specialized to certain ecological 438 niches, increasing the differences between these ecotypes and the rest of the named species [69]. However, other 439 authors emphasize the role of recombination in bacterial divergence which allows for gene-specific sweeps [70]. 440 Other pathogenic species can give some insight into the driving forces shaping the E. faecium population structure 441 we observed. For example, in the last 30 years a new multidrug resistant genotype, H58, of Salmonella enterica 442 sp. enterica sv. Typhi (S. Typhi) emerged as a clonal expansion and has rapidly spread globally [71]. This clade, 443 like E. faecium, has rapidly differentiated into major antimicrobial-resistant lineages [72], but, unlike E. faecium, 444 the differentiation has been driven by strong geographical selection [71]. Interestingly and contrary to E. faecium, 445 H58 has a similar fitness to other Salmonella genotypes in absence of antimicrobials and local genetic drift rather than niche specialization is responsible for its diversification. This difference suggests that the E. faecium clinical 447 ecotype is locked into clinical associated environments as competitive exclusion from other strains prevent its expansion [67, 68]. However, relying solely on this niche restriction to guide our surveillance efforts would be a 449 mistake, as recombinatory events are common in E. faecium [73] and several circulating non-clinical Type A strains are likely recombinants between Type A and Type B [74]. Therefore, it is possible for neglected non-pathogenic 451 strains to acquire the traits necessary to expand to the clinical environment while retaining their cosmopolitan lifestyle. 453

4.4 Towards Monitoring of Evolving Threats

that can reshuffle according to environmental pressures and opportunities, with geographic distance, phylogenetic 456 distance, and habitat boundaries as no obstacle. Although the analytical pipeline we apply here was effective in detecting environmental and genetic connections, improvements in sampling, sequencing, and analysis will be 458 needed to realize the full potential of genomic monitoring. While it makes sense to focus efforts on the sampling of clinical isolates, isolates from other environments need to be collected with appropriate metadata such as local 460 antimicrobial usage and connectivity patterns with other sampling locations. The limitations of short-read sequencing are well documented, and MGEs are generally more difficult to recover 462 due to the increased abundance of repeat regions [75–77]. Hybrid long-read / short-read assemblies can provide 463 complete or near-complete information about MGE gene content, and enrich reference databases to serve as references for short-read assemblies. Future work should also include refinements of the statistical methods used 465 and techniques to identify key genes. For example, contextual information such as gene order can enhance the differentiation of true AMR genes from highly similar false positives. In our analysis, we found that the filtering 467 parameters applied to the analytical outputs are of paramount importance. In fact, after curation, some of the more rare genes we identified proved to be artifacts generated by thresholds that were not stringent enough to root out 469 low levels of sequence contamination or short but unreliable matches to databases.

As in many other pathogens, the genomic plasticity of E. faecium effectively creates a reservoir of genes and MGEs

5 Conclusion

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Our examination of the evolution of *E. faecium* revealed a great capacity to acquire traits such as antimicrobial resistance, fueled by a large repertoire of MGEs. Misuse of antimicrobials has driven *E. faecium*'s transformation to a major contributor to morbidity and mortality worldwide. The datasets we consider here, when combined with other large collections, provide a robust snapshot of the current population structure and evolutionary trends of *E. faecium* across the One Health continuum providing crucial insight to target surveillance, design public health

- policies, and inform interventions. The work we present here can support the careful stewardship, comprehensive
- monitoring, and measurable outcomes that will be necessary to manage E. faecium and other evolving pathogens.

479 **6** Author statements

6.1 Authors and contributions

- 481 H.S., K.G., F.S.L.B., R.C.F., R.Z., and R.G.B. conceptualized the study. H.S., K.G., A.M., F.M., A.K., C.L.,
- 482 C.N.R., J.H.E.N., J.R., K.B., M.O., B.P.A., A.R.R., A.G.M., F.S.L.B., R.C.F., and R.G.B. contributed one or more
- of experimental design and execution, development and validation of software, and data analysis. H.S., F.M.,
- T.A.M., S.J.P., K.E.R., T.G., and R.G.B. contributed datasets and/or performed curation of datasets. H.S., K.G.,
- 485 A.M., A.K., R.C.F., R.Z., and R.G.B. prepared the initial draft of the manuscript. All authors edited and approved
- the final version of the manuscript.

487 6.2 Conflict of interest statement

The author(s) declare that there are no conflicts of interest.

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497 **6.4 Data summary**

- The code and data to reproduce the results are available at GitHub (https://github.com/beiko-lab/efaecium-niche).
- Supplementary Data are available at [WHERE]

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Overview of data processing workflow. (A) Short reads were trimmed using fastp, with quality checks by FastQC before and after trimming. The reads were then assembled using Unicycler and quality checked using QUAST. (B) The quality-checked assemblies were annotated with Prokka, MOB-suite, RGI, and DIAMOND. DIAMOND annotation was performed using the VFDB and BacMet databases. The Prokka annotations were passed to IslandCompare to infer probable GIs. The outputs from IslandCompare were passed to DBSCAN-SWA to infer probable phages. (C) Annotated contigs from Prokka were passed to Roary for pangenome calculation and core-genome alignment. The core genome alignment's static sites were removed using SNP-sites and the resultant alignment was passed to IQ-Tree to calculate a maximum likelihood phylogenetic tree. (D) All annotated target genes and MGEs were tabulated and passed to BayesTraits for co-evolutionary analysis, performing hypothesis testing for correlated evolution between pairs of features. 23 Size and count distributions of genomic features. First row: Genome size (A) and Pan-genome distribution predicted by Roary (B). Second row: frequency distribution of AMR genes (C), HMR genes (D), and VFs (E). Third row: frequency distribution of plasmids (F), GIs (G), and phages (H). Multiple occurrences of a feature in a given genome were counted only once. For clarity, only features detected in at least five genomes were plotted. Plot annotations indicate the number of 24 3 Abundance of features by habitat type and geographic location. "AB" indicates genomes sampled from Alberta, Canada. "UK" indicates genomes sampled from the United Kingdom. Counts indicate the number of unique features of a given category found per genome. Bars indicate quartiles. Points/diamonds are considered to be outliers if they fall outside $1.5 \times$ the interquartile range. 25 Maximum-likelihood core-genome phylogenetic tree of 1273 E. faecium genomes with E. hirae ATCC9790 as the outgroup, and E. faecium DO ASM17439v2 reference genome. The tree was constructed with 1,854,991 nucleotide sites, 79,440 of which were parsimony informative, using the general time reversible substitution model with invariant sites and four Gamma rate categories. Branch lengths are log-transformed and scaled down to 13% length for improved readability. Nodes are colored by sampling location, with hue indicating habitat and saturation indicating 26 Statistical associations and physical localization of vanA (A-D) and vanB (E-H) genes. (A,E) Phylogenetic distribution of van genes and other features with an associated likelihood ratio \geq 100. (B,F) Statistical association network of vanA/vanB genes with other features. Gene and MGE colours are consistent with those in Figure 2. (C,G) Example of gene order on an annotated plasmid (C) and GI (G) Green genes correspond to "Perfect" matches with reference genes in the CARD database, yellow genes are "Strict" hits. (D-H) Distribution of genes by habitat. Bar colors 27 Heatmap showing the presence of AMR determinant genes detected in 1273 E. faecium genomes analyzed in this study. The y-axis indicates genomes (color coded by habitat, geography and type) sorted by topology of the core genome maximum likelihood tree. AMR determinants (x-axis) are

sorted by drug class. * denotes variant versions of intrinsic genes conferring AMR.

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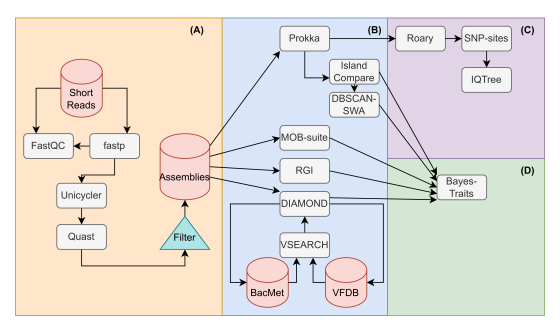


Fig. 1. Overview of data processing workflow. (A) Short reads were trimmed using fastp, with quality checks by FastQC before and after trimming. The reads were then assembled using Unicycler and quality checked using QUAST. (B) The quality-checked assemblies were annotated with Prokka, MOB-suite, RGI, and DIAMOND. DIAMOND annotation was performed using the VFDB and BacMet databases. The Prokka annotations were passed to IslandCompare to infer probable GIs. The outputs from IslandCompare were passed to DBSCAN-SWA to infer probable phages. (C) Annotated contigs from Prokka were passed to Roary for pangenome calculation and core-genome alignment. The core genome alignment's static sites were removed using SNP-sites and the resultant alignment was passed to IQ-Tree to calculate a maximum likelihood phylogenetic tree. (D) All annotated target genes and MGEs were tabulated and passed to BayesTraits for co-evolutionary analysis, performing hypothesis testing for correlated evolution between pairs of features.

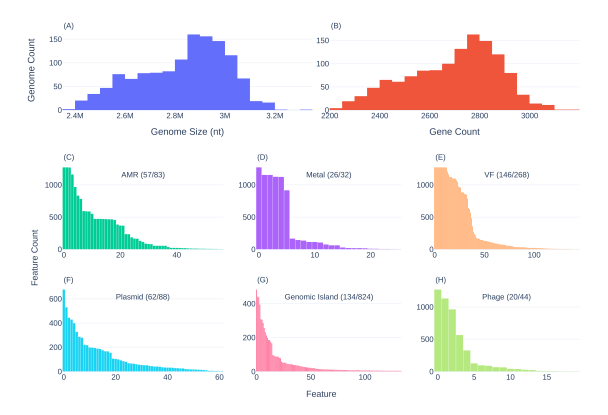


Fig. 2. Size and count distributions of genomic features. First row: Genome size (A) and Pan-genome distribution predicted by Roary (B). Second row: frequency distribution of AMR genes (C), HMR genes (D), and VFs (E). Third row: frequency distribution of plasmids (F), GIs (G), and phages (H). Multiple occurrences of a feature in a given genome were counted only once. For clarity, only features detected in at least five genomes were plotted. Plot annotations indicate the number of features plotted and the number of total features detected.

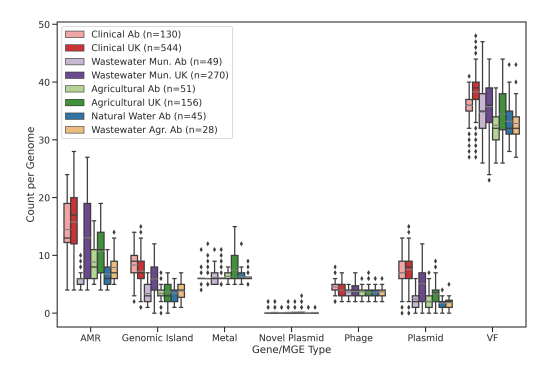


Fig. 3. Abundance of features by habitat type and geographic location. "AB" indicates genomes sampled from Alberta, Canada. "UK" indicates genomes sampled from the United Kingdom. Counts indicate the number of unique features of a given category found per genome. Bars indicate quartiles. Points/diamonds are considered to be outliers if they fall outside $1.5 \times$ the interquartile range. Grey bars indicate mean values.

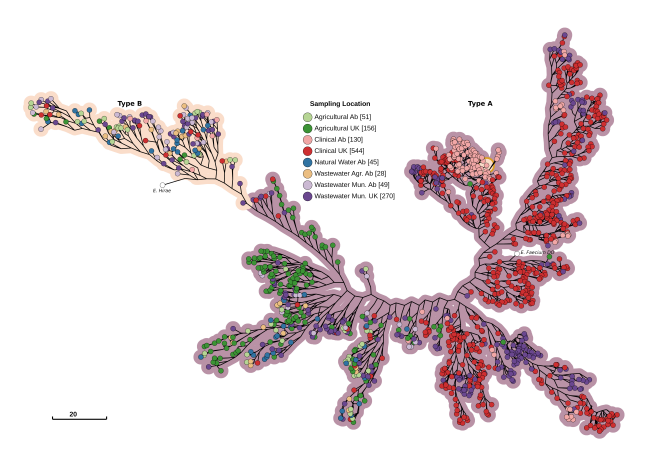


Fig. 4. Maximum-likelihood core-genome phylogenetic tree of 1273 *E. faecium* genomes with *E. hirae* ATCC9790 as the outgroup, and *E. faecium* DO ASM17439v2 reference genome. The tree was constructed with 1,854,991 nucleotide sites, 79,440 of which were parsimony informative, using the general time reversible substitution model with invariant sites and four Gamma rate categories. Branch lengths are log-transformed and scaled down to 13% length for improved readability. Nodes are colored by sampling location, with hue indicating habitat and saturation indicating geography.

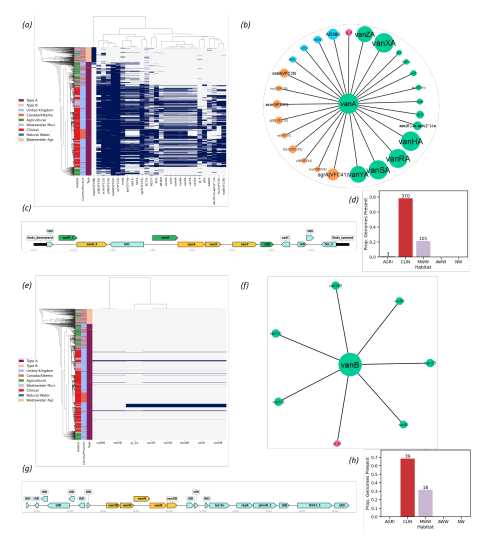


Fig. 5. Statistical associations and physical localization of vanA (A-D) and vanB (E-H) genes. (A,E) Phylogenetic distribution of van genes and other features with an associated likelihood ratio ≥ 100 . (B,F) Statistical association network of vanA/vanB genes with other features. Gene and MGE colours are consistent with those in Figure 2. (C,G) Example of gene order on an annotated plasmid (C) and GI (G) Green genes correspond to "Perfect" matches with reference genes in the CARD database, yellow genes are "Strict" hits. (D-H) Distribution of genes by habitat. Bar colors correspond to their habitats as per the legend in (A,E).

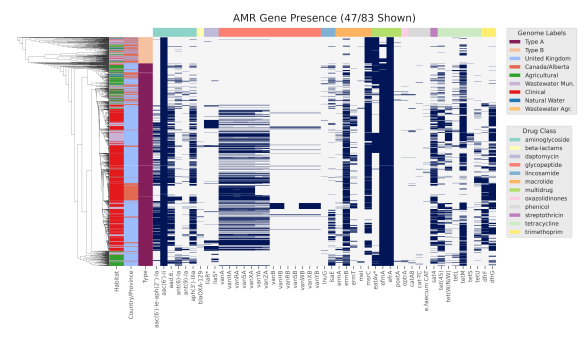


Fig. 6. Heatmap showing the presence of AMR determinant genes detected in 1273 *E. faecium* genomes analyzed in this study. The y-axis indicates genomes (color coded by habitat, geography and type) sorted by topology of the core genome maximum likelihood tree. AMR determinants (x-axis) are sorted by drug class. * denotes variant versions of intrinsic genes conferring AMR.

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Three-way ANOVA results for 1200 genomes in habitats that were sampled in both AB and UK. 803 Type III error correction was used to account for imbalanced classes. Columns indicate p-values 804 testing differences of mean unique features per genome. Factor1×Factor2 indicates interaction 805 30 Two-way ANOVA for 303 AB genome assemblies, performed separately to account for the WW-2 807 AGR and NWS habitats exclusive to AB. Type III error correction used to account for imbal-808 anced classes. Columns indicate significance values testing differences of mean unique features 809 per genome. "Habitat×Type" indicates the interaction effect of habitat and type categories. 31 810

Table 1. Three-way ANOVA results for 1200 genomes in habitats that were sampled in both AB and UK. Type III error correction was used to account for imbalanced classes. Columns indicate p-values testing differences of mean unique features per genome. Factor1×Factor2 indicates interaction effects among categories.

Variable	AMR	HMR	VF	Plasmid	GI	Phage
Type	6.42×10^{-04}	2.36×10^{-01}	8.92×10^{-02}	5.87×10^{-01}	9.59×10^{-01}	4.69×10^{-02}
Hab.	5.09×10^{-15}	7.26×10^{-02}	1.29×10^{-09}	1.74×10^{-31}	5.03×10^{-40}	1.14×10^{-08}
Geo.	8.34×10^{-01}	4.16×10^{-05}	2.28×10^{-04}	8.76×10^{-03}	6.46×10^{-01}	7.80×10^{-01}
Type×Hab.	6.31×10^{-04}	2.54×10^{-02}	1.46×10^{-04}	1.05×10^{-04}	5.26×10^{-07}	7.41×10^{-04}
Type×Geo.	3.04×10^{-01}	6.90×10^{-01}	7.12×10^{-01}	3.62×10^{-01}	4.42×10^{-01}	6.11×10^{-01}
Habitat×Geo.	2.20×10^{-08}	1.62×10^{-04}	6.58×10^{-01}	2.50×10^{-04}	1.76×10^{-13}	1.16×10^{-03}
Type \times Geo. \times Hab.	1.76×10^{-04}	4.62×10^{-01}	4.88×10^{-03}	1.74×10^{-01}	2.61×10^{-03}	7.04×10^{-02}

Table 2. Two-way ANOVA for 303 AB genome assemblies, performed separately to account for the WW-AGR and NWS habitats exclusive to AB. Type III error correction used to account for imbalanced classes. Columns indicate significance values testing differences of mean unique features per genome. "Habitat×Type" indicates the interaction effect of habitat and type categories.

Variable	AMR	HMR	VF	Plasmid	GI	Phage
Habitat		2.48×10^{-03}				
Type	2.34×10^{-07}	1.21×10^{-01}	7.09×10^{-02}	4.40×10^{-01}	9.50×10^{-01}	4.48×10^{-02}
Habitat×Type	4.71×10^{-08}	2.57×10^{-04}	2.32×10^{-05}	1.76×10^{-08}	1.16×10^{-09}	1.93×10^{-03}