Genomics of vancomycin-resistant *Enterococcus faecium*

Claire Gorrie†, Charlie Higgs†, Glen Carter2,3, Timothy P. Stinear2,3 and Benjamin Howden1,3,4,*

**Abstract**

Vancomycin-resistant *Enterococcus faecium* (VREfm) is a globally significant public health threat and was listed on the World Health Organization’s 2017 list of high-priority pathogens for which new treatments are urgently needed. Treatment options for invasive VREfm infections are very limited, and outcomes are often poor. Whole-genome sequencing is providing important new insights into VREfm evolution, drug resistance and hospital adaptation, and is increasingly being used to track VREfm transmission within hospitals to detect outbreaks and inform infection control practices. This mini-review provides an overview of recent data on the use of genomics to understand and respond to the global problem of VREfm.

**INTRODUCTION**

*Enterococcus faecium*, a species of Gram-positive cocci, is recognized as a globally important opportunistic pathogen that is capable of causing a range of human infections associated with high mortality rates, particularly in hospitalized individuals [1, 2]. Enterococci are inherently resistant to a number of antimicrobial classes, and over recent decades there has been a significant increase in the rates of acquired antimicrobial resistance (AMR) in *E. faecium*, including vancomycin-resistant *E. faecium* (VREfm) [3, 4]. Since the first reports in the late 1980s, VREfm have come to represent a globally significant public health threat, such that in some regions (e.g. the USA and Australia) up to 50% or more of all blood culture isolates of *E. faecium* are vancomycin-resistant (Fig. 1) [5].

The World Health Organization (WHO) recently published their list of priority bacterial pathogens for which new antibiotics are urgently needed, and VREfm is listed in the high-priority category [6]. As well as being resistant to vancomycin, VREfm usually exhibit resistance to a wide range of other antibiotics, including intrinsic resistance to cephalosporins, lincosamides, aminoglycosides and trimethoprim/sulfamethoxazole, almost universal resistance to penicillins, and increasing reports of acquired resistance to last-line agents such as linezolid and daptomycin [1, 5, 7, 8].

Genomic data can be used to understand the evolution of VREfm, determine resistance mechanisms and investigate the potential sources of human VREfm infection (Table 1).

Clones of VREfm associated with invasive human infection are recognized as primarily healthcare-associated pathogens. Approaches to molecular typing of VREfm are therefore important for informing molecular epidemiology and methods of infection prevention in the healthcare setting. Approaches such as multilocus sequence typing (MLST) have been used to classify *E. faecium* [9], but this approach has a number of limitations, including the lack of fine resolution necessary for inferring putative transmission. Phylogenetic analyses based on whole-genome sequencing (WGS) data are increasingly being used in combination with – or in place of – MLST in order to overcome this limitation and identify transmission events [10, 11].

This mini-review will describe how genomic data have been used to understand the mechanisms of vancomycin resistance in VREfm, the clonal structure and evolution of *E. faecium*.

**Abbreviations**: AMR, antimicrobial resistance; CC, clonal complex; cgMLST, core genome multilocus sequence typing; Efm, *Enterococcus faecium*/*E. faecium*; MLST, multilocus sequence typing; SNP, single nucleotide polymorphism; ST, sequence type; VRE, vancomycin-resistant Enterococcus; VREfm, vancomycin-resistant *Enterococcus faecium*; VSE, vancomycin-susceptible Enterococcus; VSEfm, vancomycin-susceptible *Enterococcus faecium*; WGS, whole genome sequencing; WHO, World Health Organization.

†These authors contributed equally to this work

*Correspondence:* Benjamin Howden, bhowden@unimelb.edu.au

**Keywords:** *Enterococcus faecium*; antibiotic resistance; whole-genome sequencing.

**Received 07 May 2019; Accepted 02 July 2019; Published 22 July 2019**

**Author affiliations:** 1Microbiological Diagnostic Unit Public Health Laboratory, Department of Microbiology and Immunology, The University of Melbourne at the Peter Doherty Institute for Infection and Immunity, Melbourne, Australia; 2Department of Microbiology and Immunology, The University of Melbourne at the Peter Doherty Institute for Infection and Immunity, Melbourne, Australia; 3Doherty Applied Microbial Genomics, Department of Microbiology and Immunology, The University of Melbourne at the Peter Doherty Institute for Infection and Immunity, Melbourne, Australia; 4Department of Infectious Diseases, Austin Health, Heidelberg, Australia.

**Keywords:** *Enterococcus faecium*; antibiotic resistance; whole-genome sequencing.

**Abbreviations:** AMR, antimicrobial resistance; CC, clonal complex; cgMLST, core genome multilocus sequence typing; Efm, *Enterococcus faecium*/*E. faecium*; MLST, multilocus sequence typing; SNP, single nucleotide polymorphism; ST, sequence type; VRE, vancomycin-resistant Enterococcus; VREfm, vancomycin-resistant *Enterococcus faecium*; VSE, vancomycin-susceptible Enterococcus; VSEfm, vancomycin-susceptible *Enterococcus faecium*; WGS, whole genome sequencing; WHO, World Health Organization.

†These authors contributed equally to this work

**Data statement:** All supporting data, code and protocols have been provided within the article or through supplementary data files.

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and VREfm clinical isolates, and emerging mechanisms of resistance to last-line antibiotics.

**Mechanisms of vancomycin resistance**

Vancomycin resistance in hospital-adapted *E. faecium* can be conferred by a number of van operon variants, though the most common, and therefore most clinically relevant, are the VanA and VanB operons (Table 2) [3, 5]. Both van operons encode a ligase that alters the target binding site of vancomycin, which is critical for antibiotic binding [25]. Globally, vanA VREfm is most common, while in some regions, including in Australia, vanB VREfm has been the major issue. However, this trend is now changing, with a recent rapid increase in vanA VREfm being seen in some parts of the Australia [26] and some parts of Europe, such as Germany, reporting a rapid increase in vanB VREfm [27]. Other van types are less commonly reported.

The VanA operon is carried on Tn1546-type transposons, which display a high degree of heterogeneity. Point mutations, deletions and various insertion sequences have all been associated with Tn1546-type transposons [46, 47]. Tn1546 is found on pRUM-like plasmids (predominant in the USA) and Inc18 plasmids (predominant in Europe) [5], however, much remains unknown and the molecular epidemiology of VanA-containing plasmids may differ significantly between regions. In addition to contributing to the spread of vancomycin resistance, Inc18 and pRUM-like plasmids can also harbour resistance genes for multiple other antibiotics [48].

The human gastrointestinal tract is a natural reservoir of non-enterococcal species containing the VanB operon [49], and the *in vivo* transfer of a VanB-containing transposon from a non-enterococcal species to vancomycin-susceptible *E. faecium* may explain the emergence of VREfm in some cases [20]. The human bowel has also been shown to be a potential reservoir for the VanD and VanG operons [50]. The natural reservoir of the VanA operon remains unknown [49, 51].

**Impact Statement**

The rise of vancomycin-resistant *Enterococcus faecium* (VREfm) poses a public health threat, especially within healthcare settings. To monitor and limit the spread of VREfm we need to understand how it evolves and acquires vancomycin resistance, how transmission networks are operating and how VREfm is developing resistance to last-line antibiotics. This mini-review provides an overview of how whole-genome sequencing is being used to address these questions.

**Classification and adaptative evolution of VRE clones associated with human colonization and infection**

**Strain classification methods and the role of WGS**

Historically, VREfm have been classified based on a number of typing methods, including MLST, which characterizes the diversity of isolates based on the allelic variation of seven housekeeping genes in order to assign a sequence type (ST) [9]. Classifying VREfm, and *E. faecium* more broadly, by ST has allowed the identification of persistent or dominant clones on both a local and a global scale; however, MLST has a
Table 1. Selected publications using WGS to investigate genomic features, evolution and transmission of *E. faecium*

| Category                  | Aims                                                                 | Main findings                                                                                                                                                                                                 |
|----------------------------|----------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Complete genomes           | Lam et al. [12] Published 3.0 MB complete genome of an ST17 isolate from Australia |
|                            | Qin et al. [13] Published a fully assembled, finished *E. faecium* genome |
|                            | Galloway-Peña et al. [14] Study the diversity of *E. faecium* and identify any hospital or community-associated clades. The sequences were used to define the core genome and to infer the role of recombination after isolation. |
|                            | de Been et al. [15] Understand the role of recombination within the core genome of *E. faecium* isolates, and how recombination contributes to the diversity within different clades. |
|                            | Lebreton et al. [16] Use WGS to understand the role of recombination within the core genome of *E. faecium* isolates, and how recombination contributes to the diversity within different clades. |
|                            | Raven et al. [17] Study the genomic variability within the *E. faecium* population of the United Kingdom and Ireland using WGS. The study aimed to understand the genetic diversity of hospital-associated and community-associated *E. faecium* isolates. |
|                            | Galloway-Peña et al. [18] Use WGS to investigate the diversity of *E. faecium* within a hospital network. |
|                            | de Been et al. [19] Understand the role of recombination within the core genome of *E. faecium* isolates, and how recombination contributes to the diversity within different clades. |
|                            | Lebreton et al. [20] Use WGS to understand the role of recombination within the core genome of *E. faecium* isolates, and how recombination contributes to the diversity within different clades. |
|                            | Raven et al. [21] Study the genomic variability within the *E. faecium* population of the United Kingdom and Ireland using WGS. The study aimed to understand the genetic diversity of hospital-associated and community-associated *E. faecium* isolates. |
|                            | Theodore et al. [22] Understand the role of recombination within the core genome of *E. faecium* isolates, and how recombination contributes to the diversity within different clades. |
|                            | Pinholt et al. [23] Study the epidemiology and clonal relatedness of *E. faecium* isolates to investigate the reasons for failure of infection control strategies. |
|                            | Raven et al. [24] Undertake a retrospective genomic based transmission investigation at a single hospital in the United Kingdom. |
|                            | Pinholt et al. [25] Investigate the epidemiology and clonal relatedness of *E. faecium* isolates to investigate the reasons for failure of infection control strategies. |
| Evolution of VRE           | Lam et al. [12] Published 3.0 MB complete genome of an ST17 isolate from Australia |
|                            | Qin et al. [13] Published a fully assembled, finished *E. faecium* genome |
|                            | Galloway-Peña et al. [14] Study the diversity of *E. faecium* and identify any hospital or community-associated clades. The sequences were used to define the core genome and to infer the role of recombination after isolation. |
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| Hospital/outbreak studies  | Lam et al. [12] Published 3.0 MB complete genome of an ST17 isolate from Australia |
|                            | Qin et al. [13] Published a fully assembled, finished *E. faecium* genome |
|                            | Galloway-Peña et al. [14] Study the diversity of *E. faecium* and identify any hospital or community-associated clades. The sequences were used to define the core genome and to infer the role of recombination after isolation. |
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VRE emerged owing to importation of a successful VREfm clone that rapidly transmitted to most hospitals in the region.
Table 2. Characteristics of van operons in Enterococcus spp.

| Van operon | Level of resistance | Location and mobility (transposon) | References |
|------------|---------------------|-----------------------------------|------------|
| vanA       | High                | Chromosome, transferable (Tn1546), plasmid | [28, 29] |
| vanB       | High (variable)     | Chromosome, transferable (Tn1547, Tn1549, Tn5382) | [28, 30–33] |
| vanC       | Low                 | Chromosome                         | [34–37] |
| vanD       | Low to high (variable) | Low to high (variable) | [38–40] |
| vanE       | Low to moderate     | Chromosome                         | [28, 35, 41] |
| vanG       | Low                 | Chromosome, transferable            | [42] |
| vanL       | Low                 | Unknown, transferable              | [43] |
| vanM       | High                | Plasmid                            | [44] |
| vanN       | Low                 | Susceptible                        | [45] |

Note: table adapted from [8] with additional material as per the references column.

number of significant limitations, including poor discriminatory power [21]. Additionally, some VREfm isolates have been identified that lack the pstS gene, one of the seven MLST loci [19], and recombination has been shown to affect the VREfm genome in the regions of key MLST loci [17, 20]. The highly dynamic nature of the E. faecium genome, with the constant flux of accessory genes and horizontal gene transfer events [52, 53], further exacerbates the shortcomings of the MLST method. WGS has proven to be the ultimate molecular typing method for VREfm [21, 54, 55]. Comparative analyses have been based on pairwise single-nucleotide polymorphism (SNP) comparisons, and more recently a core genome MLST (cgMLST) approach using 1423 target genes that has a similar resolution to that of an SNP-based approach [15].

VREfm clonal types associated with human disease

The majority of sequenced strains are human clinical isolates, with an overrepresentation of isolates from Europe and North America, but these only form a small part of the total E. faecium population [52]. A comparison of 100 core genes in all available E. faecium genomes initially found that most of the genes split into 2 clades, differing by 3.5–4.2% nucleotide divergence [14]. The two distinct, highly genetically divergent clades can be grouped by origin: clade A (hospital-associated; consisting mainly of common clinical isolates and some isolates of community origin) and clade B [community associated; consisting almost entirely of isolates from the community (both human and animal)] [14, 16–18]. Initial comparative genome studies suggested that clade A could be subdivided into clade A1 (hospital-associated) and clade A2 (animal-associated) [16, 18], a distinction not fully supported by some later studies [17]. The majority of hospital-associated VREfm are part of clonal complex 17 (CC17) and common STs within CC17 include ST17, ST18, ST80 and ST203 [56]. Genomic studies on global populations of E. faecium, including VREfm, have revealed that in some cases close genetic relationships exist between vanB VREfm isolates and vancomycin-susceptible E. faecium (VSE) from the same institutions, suggesting possible frequent generation of new vanB VREfm clones within resident VSE isolates [20], while other studies indicate frequent intra-regional and inter-regional spread of VREfm clones [17, 21, 53].

Role of genomic plasticity in adaptation to the healthcare environment

WGS has revealed that horizontal gene transfer – involving plasmids, prophages, genomic islands, homologous recombination and the recently identified enterococcus cassette chromosome [57] – is an important driver of the diversity and adaptive evolution of the dynamic E. faecium genome [52, 58, 59]. Complete sequencing of VREfm genomes has revealed a large number of these accessory elements, as well as many insertion sequences in clade A VREfm genomes [12]. The ability of E. faecium to acquire foreign DNA has been central in the evolution of drug resistance and hospital adaptation in this species. At the core genome level, chromosome regions have been identified that predominantly recombine in specific hospital-associated E. faecium strains [59]; these regions include those involved in the biosynthesis of cell wall polysaccharides and carbohydrate uptake and metabolism. Genes involved in these processes have been shown to be important in successful colonization and overgrowth of hospital-adapted E. faecium during antibiotic therapy [60], with the gain and loss of carbohydrate metabolism genes reflecting the environmental/colonizing niche of the isolates. Clade B isolates tend to contain genes involved in complex carbohydrate utilization, while clade A, hospital-adapted VREfm, contain genes involved in the utilization of amino sugars found at the epithelial surface [61].
A pan-genome analysis by van Schaik et al. [58] indicated that the total available gene pool within the species is expanding (open pan-genome), with E. faecium able to acquire genes from many different bacteria. However, the majority of imported DNA is from other E. faecium populations [53, 59], with clade B (community-associated) being the most important reservoir for foreign DNA for clade A (hospital-associated) [59]. Multidrug-resistant, hospital-adapted E. faecium have also been shown to lack clustered regularly interspaced short palindromic repeats (CRISPR) self-defence systems that protect from genomic modification by plasmids and phage, thereby making them more susceptible to changes through horizontal gene transfer [62]. A recent study by Pinholt et al. [24] comparing over 800 VREfm strains suggested that the success of a dominant VREfm clone was due to the acquisition of a heterogeneous accessory genome, with no single successful combination of accessory genes. This flexibility and large variation in the accessory genes is an important driver of the evolution of E. faecium and its adaptation to the healthcare environment [53, 58, 59], as it is readily able to acquire genes that increase fitness under adverse conditions.

WGS has also shown that the E. faecium genome contains a variety of loci encoding key virulence factors that make it particularly suited to persistence in a hospital environment and cause invasive infections, as summarized in a review by Gao et al. [63]. A recent study identified a unique botulinum neurotoxin gene cluster (host species target to be determined) in a commensal strain of E. faecium and demonstrated the ability of E. faecium to horizontally acquire and possibly disseminate this gene cluster [64].

Role of VREfm genomics in hospital infection control

WGS is providing the capacity for highly discriminatory identification of pathogen outbreaks and transmission, including in the hospital setting. The application of WGS to putative hospital outbreaks of VREfm has been reported, with clear evidence of utility (Table 1), and in some cases has resulted in changes in infection control practices [22, 54, 65]. WGS of VREfm has particular value in identifying complex transmission networks, especially where multiple wards are involved, where patients move wards between VREfm acquisition and the development of a clinical infection, and where environmental contamination is contributing [22]. Although many unknowns remain with regard to percentage genome-wide nucleotide identity and SNP thresholds for identifying transmission events, using WGS data complemented by space–time epidemiological data can allow these more complex transmission links to be defined, informing appropriate interventions. In Australia, WGS has additionally been used to detect and describe the emergence of new clones of VREfm that have become regionally important, including pstS null strains (now ST1421-ST1425), which lack one of the MLST typing genes [19, 66].

Mechanisms and emergence of resistance to last-line antibiotics

Treatment options for VREfm infections, especially vanA VREfm, are limited and last-line antibiotics such as linezolid, daptomycin, tigecycline and possibly quinupristin/dalfopristin are often needed. The majority of linezolid-resistant VREfm cases have been found in North America and Europe [67] and although the majority of linezolid-resistant cases appear to emerge after linezolid treatment, linezolid-resistant enterococci have been shown to emerge in the absence of linezolid treatment [68, 69]. Of particular concern are the optrA, poxA and cfr resistance genes, which can be rapidly disseminated in E. faecium on mobile genetic elements [70–72], particularly under the selective pressures found in clinical settings. Daptomycin non-susceptibility has been shown to emerge in VREfm both during daptomycin therapy [73] as well as in the absence of previous exposure to the antibiotic [74]. The recent review by Ahmed and Baptiste [8] outlines the genotypes associated with resistance to last-resort antibiotics such as these. Teicoplanin remains an option for therapy for some vanB VREfm infections, although the emergence of resistance during therapy has been reported, especially in high-bacterial-load infections, and WGS-based studies have shown that this may be due to mutations in the vanS region of the VanB operon [75]. Genomics has been an important tool in understanding the mechanisms of resistance to last-line agents, especially given the often polygenic nature of resistance mechanisms.

In addition to growing drug resistance, a recent study has also suggested that strains of VREfm are becoming more tolerant to alcohol-based disinfectants, with isolates collected after 2010 being significantly more tolerant to killing by alcohol compared to older isolates in Australia through mutations occurring in cell wall-associated transport proteins [76]. Other studies have shown that repeated exposure to chlorhexidine, an antiseptic used in infection control practices, can lead to reduced chlorhexidine susceptibility [77, 78]. The wider consequences of these findings on hospital-based infection prevention strategies remain to be seen, but these studies further highlight the highly adaptive nature of VREfm in the face of selective pressures. These data provide a warning about potential future adaptations that will pose further challenges to the control of this important nosocomial pathogen.

Conclusions and view to the future

VREfm represent a significant public health threat, causing opportunistic invasive infections that are often broadly resistant to available antibiotic treatments. E. faecium are highly adaptive and evolve rapidly; the level of variation within a population is rarely accurately represented by traditional typing approaches, although the use of WGS is now providing important new insights into the genetic mechanisms underlying the evolution and adaptation of this nosocomial pathogen. Studies implementing WGS have been able to define both local and large-scale spread of clones, as
well as elucidating genomic aspects related to host specificity, resistance and hospital adaptation. These studies have also highlighted the importance of vancomycin-susceptible *E. faecium* (VSE) as a nosocomial pathogen that underpins the further evolution and spread of VREfm. These studies have helped identify risk factors, adaptations and mechanisms for transmission and treatment failure that now need to be leveraged to inform and direct efforts to limit the further evolution and spread of VREfm. While in some regions the burden of VREfm in the healthcare setting may appear overwhelming, the precision of WGS has the potential to reveal the complexity of this problem and provide the evidence for real action.

**Funding information**
B. P. H. receives a Practitioner Fellowship from the National Health and Medical Research Council, Australia (NHMRC, GNT1105905).

**Author contributions**
C. G. and C. H. were primarily responsible for preparing the review. All authors contributed to writing the review and editing the final version.

**Conflicts of interest**
The authors declare that there are no conflicts of interest.

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Author/s:
Gorrie, C; Higgs, C; Carter, G; Stinear, TP; Howden, B

Title:
Genomics of vancomycin-resistant Enterococcus faecium

Date:
2019-07-01

Citation:
Gorrie, C., Higgs, C., Carter, G., Stinear, T. P. & Howden, B. (2019). Genomics of vancomycin-resistant Enterococcus faecium. MICROBIAL GENOMICS, 5 (7), https://doi.org/10.1099/mgen.0.000283.

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