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Evolutionary origins of the emergent ST796 clone of vancomycin resistant Enterococcus faecium

Andrew H Buultjens 1, Margaret M C Lam 1, Susan Ballard 2, Ian R Monk 1, Andrew A Mahony 3, Elizabeth A Grabsch 3, M Lindsay Grayson 3, Stanley Pang 4,5, Geoffrey W Coombs 4,5, J Owen Robinson 4,5, Torsten Seemann 6, Paul D R Johnson Corresp. 3,7, Benjamin P Howden Corresp. 2, Timothy P Stinear Corresp. 1

1 Department of Microbiology and Immunology, Peter Doherty Institute for Infection and Immunity, University of Melbourne, Melbourne, Victoria, Australia
2 Microbiology Diagnostic Unit, Department of Microbiology and Immunology, Peter Doherty Institute for Infection and Immunity, University of Melbourne, Melbourne, Victoria, Australia
3 Infectious Diseases Department, Austin Health, Heidelberg, Victoria, Australia
4 School of Veterinary and Life Sciences, Murdoch University, Murdoch, Western Australia, Australia
5 Department of Microbiology, Fiona Stanley Hospital, Murdoch, Western Australia, Australia
6 Victorian Life Sciences Computation Initiative, University of Melbourne, Carlton, Victoria, Australia
7 Department of Medicine, University of Melbourne, Heidelberg, Victoria, Australia

Corresponding Authors: Paul D R Johnson, Benjamin P Howden, Timothy P Stinear
Email address: Paul.Johnson@austin.org.au, bhowden@unimelb.edu.au, tstinear@unimelb.edu.au

From early 2012, a novel clone of vancomycin resistant Enterococcus faecium (assigned the multi locus sequence type ST796) was simultaneously isolated from geographically separate hospitals in south eastern Australia and New Zealand. Here we describe the complete genome sequence of Ef_aus0233, a representative ST796 E. faecium isolate. We used PacBio single molecule real-time sequencing to establish a high quality, fully assembled genome comprising a circular chromosome of 2,888,087 bp and five plasmids. Comparison of Ef_aus0233 to other E. faecium genomes shows Ef_aus0233 is a member of the epidemic hospital-adapted lineage and has evolved from an ST555-like ancestral progenitor by the accumulation or modification of five mosaic plasmids and five putative prophage, acquisition of two cryptic genomic islands, accrued chromosomal single nucleotide polymorphisms and a 80kb region of recombination, also gaining Tn1549 and Tn916, transposons conferring resistance to vancomycin and tetracycline respectively. The genomic dissection of this new clone presented here underscores the propensity of the hospital E. faecium lineage to change, presumably in response to the specialized conditions of hospital and healthcare environments.
Evolutionary origins of the emergent ST796 clone of vancomycin resistant *Enterococcus faecium*

Andrew H. Buultjens 1, Margaret M. C. Lam 1, Susan Ballard 2,3, Ian. R. Monk 1, Andrew A. Mahony 3, Elizabeth A. Grabsch 3, M. Lindsay Grayson 3, Stanley Pang 4,5, Geoffrey W. Coombs 4,5, J. Owen Robinson 4,5, Torsten Seemann 6, Paul D. R. Johnson 1,3,7,*, Benjamin P. Howden 1,2,7,*, and Timothy P. Stinear 1,*

Address: 1 Department of Microbiology and Immunology, Doherty Institute for Infection and Immunity, The University of Melbourne, Victoria, 3010, Australia; 2 Microbiology Diagnostic Unit, Department of Microbiology and Immunology, Doherty Institute for Infection and Immunity, The University of Melbourne, Victoria, 3010, Australia; 3 Infectious Diseases Department, Austin Health, Heidelberg, Victoria, 3084, Australia; 4 School of Veterinary and Life Sciences, Murdoch University, Western Australia, 6150, Australia; 5 Department of Microbiology, PathWest Laboratory Medicine – WA, Royal Perth Hospital, Fiona Stanley Hospital, Murdoch, Western Australia, 6150, Australia; 6 Victorian Life Sciences Computation Initiative, Carlton, Victoria, 3053, Australia; 7 Department of Medicine, University of Melbourne, Heidelberg, Victoria, 3084, Australia.

*Joint senior authors

ABSTRACT

From early 2012, a novel clone of vancomycin resistant *Enterococcus faecium* (assigned the multi locus sequence type ST796) was simultaneously isolated from geographically separate hospitals in south eastern Australia and New Zealand. Here we describe the complete genome sequence of Ef_aus0233, a representative ST796 *E. faecium* isolate. We used PacBio single molecule real-time sequencing to establish a high quality, fully assembled genome comprising a circular chromosome of 2,888,087 bp and five plasmids. Comparison of Ef_aus0233 to other *E. faecium* genomes shows Ef_aus0233 is a member of the epidemic hospital-adapted lineage and has evolved from an ST555-like ancestral progenitor by the accumulation or modification of five mosaic plasmids and five putative prophage, acquisition of two cryptic genomic islands, accrued
chromosomal single nucleotide polymorphisms and a 80kb region of recombination, also
gaining Tn1549 and Tn916, transposons conferring resistance to vancomycin and tetracycline
respectively. The genomic dissection of this new clone presented here underscores the
propensity of the hospital *E. faecium* lineage to change, presumably in response to the specific
conditions of hospital and healthcare environments.

**INTRODUCTION**

*Enterococcus faecium* is a human and animal gastrointestinal tract (GIT) commensal but a
lineage within the species has rapidly evolved to become a significant opportunistic pathogen
(Coombs et al. 2014c; Deshpande et al. 2007; Lebreton et al. 2013b; Pinholt et al. 2014).

Early genotyping methods such as amplified fragment length polymorphism (Willems et al.
2000), restriction endonuclease analysis (Quednau et al. 1999), multi-locus sequence typing
(MLST) (Leavis et al. 2006; Top et al. 2008) and more recent analyses using whole genome
datasets (Lebreton et al. 2013), have shown the *E. faecium* population separates into two major
clades largely according to source origin, designated as clades A and B. Clade B strains are
community-associated and mostly of non-clinical origin while clade A strains are hospital-
associated and mostly of clinical origin (Galloway-Peña et al. 2012; Leavis et al. 2007; Palmer et
al. 2012). Clade A has been found to further divide into clade A1, which contains epidemic
hospital strains, and clade A2, which encompasses animal strains and strains linked to sporadic
human infections (Lebreton et al. 2013).

Clade A1 or clonal complex 17 (CC17, a MLST designation) has adapted to the hospital
environment and is adept at GIT colonization with the potential to cause invasive disease (Top
et al. 2008; Willems et al. 2005). Members of clade A1 are characterised by larger genomes and
harbour a greater abundance of virulence factors and genes conferring antibiotic resistance
compared to non-A1 *E. faecium* lineages, a reflection of adaptation to healthcare environments
In Australia, as in other countries, we have observed the sequential emergence of new \textit{E. faecium} clones within the clade A1 hospital lineage which spread rapidly and displace previously endemic clones. For example, from 1994 to 2005, Australian hospital acquired \textit{E. faecium} VRE was uncommon and mostly caused by ST17 strains. The situation changed suddenly from 2005 when there was a nationwide wave of by \textit{E. faecium} ST203 blood stream infections (BSI), a significant and rising proportion of which are \textit{vanB} VRE (Coombs et al. 2014; Johnson et al. 2010; Lam et al. 2012). Previous work comparing ST17 and ST203 genomes revealed that ST203 possesses 40 unique genes with inferred functions of riboflavin metabolism, ion transport and phosphorylation, and harboured a larger vancomycin resistance-conferring Tn1549 transposon (Lam et al. 2013).

At the Austin Hospital in Melbourne, improved cleaning protocols following our local ST203 outbreak were associated with a reduction in VRE BSI between 2009 and 2011 (Grabsch et al. 2012). However, despite retaining these protocols we once again observed an abrupt increase in \textit{vanB} VRE \textit{E. faecium} BSI from 2012 onwards that was caused by a completely new ST. We originally recognised the change in strain using PFGE and a high-resolution melt method (Tong et al. 2011) but have now switched to whole genome sequencing for epidemiological typing. We lodged the alleles of the new ST with the MLST Database and received the new designation ST796 in September 2012 (Mahony et al. 2014). ST796 was unknown before 2011 but by 2013 \textit{vanB} ST796 \textit{E. faecium} had caused a large outbreak of colonisation in a Melbourne Neonatal Intensive Care (Lister et al. 2015) and in the same year was responsible for 40% of \textit{E. faecium} VRE BSI in 5 geographically separate Melbourne hospitals, largely replacing its ST203 predecessor strains. In 2015, ST796 \textit{vanB} \textit{E. faecium} was responsible for 62 of 117 (53%) of all patient episodes of all \textit{E. faecium} bacteraemia in Melbourne Hospitals, compared with 10 of 117 (8.5%) for ST203.

In the current study, we used single molecule real-time sequencing to establish a high quality, fully assembled genome sequence of ST796 \textit{E. faecium} isolate Ef_aus0233, a representative of
this emerging clone and then employed population based comparative genomics to better understand the genetic changes that have accompanied the emergence.

METHODS

Bacterial strains. A list of the isolates examined in the study is provided (Table S1). *E. faecium* were cultured as previously described (Johnson et al. 2010).

Whole Genome Sequencing. Short fragment DNA libraries were generated using the Illumina NexteraXT DNA preparation kit and fragment sequencing was undertaken with the Illumina NextSeq 500 platform using 2x150 bp chemistry. Highly intact and high quality genomic DNA was extracted from Ef_aus0233 and subjected to Pacific Biosciences SMRT sequencing according to the manufacturer’s instructions and sequenced with two SMRT cells on the RS II platform (Pacific Biosciences) using P5-C3 chemistry. Genome assembly was performed using the SMRT Analysis System v2.3.0.140936 (Pacific Biosciences). Raw sequence data were *de novo* assembled using the HGAP v3 protocol with a genome size of 3 Mb. Polished contigs were error corrected using Quiver v1. The resulting assembly was then checked using BridgeMapper v1 in the SMRT Analysis System, and the consensus sequence corrected with short-read Illumina data, using the program Snippy (https://github.com/tseemann/snippy). The final chromosome assembly was validated by reference to a high-resolution NcoI optical map using MapSolver (version 3.10; OpGen, Maryland USA). Common bacterial DNA base modifications and methyltransferase motifs were assessed using the protocol, RS_Modification_and_Motif_Analysis in the SMRT Analysis System v2.3.0.140936 (Pacific Biosciences).

Plasmid Copy Number. The approximate number of plasmid copies per cell for the Ef_aus0233 genome was inferred using differences in Illumina sequence read depth. The read depth of plasmid sequences was compared to the average chromosomal coverage to estimate copy number multiplicity.
Comparison of completed genomes. Artemis Comparison Tool (Carver et al. 2005) was used to align the chromosomes of four fully assembled *E. faecium* genomes. BRIG (Alikhan et al. 2011) was used to visualize DNA:DNA comparisons using BLASTn for comparisons of the EfAus0233 chromosome against other fully assembled *E. faecium* chromosomes.

De novo assembly and genome annotation. Illumina sequence reads were *de novo* assembled into contigs using Spades v3.6.1 (Nurk et al. 2013). The closed EfAus0233 genome and Spades contigs were annotated using Prokka (v1.12b) (Seemann 2014). Multilocus sequence types (STs) were determined using an *in silico* tool (https://github.com/tseemann/mlst). CRISPR databases were used to search for CRISPR sequences (http://crispi.genouest.org and http://crispr.uppsud.fr/Server/) (accessed 19th of May 2016). Sequence files were uploaded to the web based ISsaga (Varani et al. 2011)(accessed 11th of February 2016) to detect both the abundance and diversity of insertion elements. Phage discovery was undertaken using the web based resource PHAST (accessed 15th of February 2016) (Zhou et al. 2011).

Variant detection and Bayesian population clustering. Snippy was used to map short read data against the full-assembled EfAus0233 genome to call core genome single nucleotide polymorphism (SNP) differences. Hierarchical Bayesian clustering was performed upon a core SNP alignment to assign genomes into discrete populations using hierBAPS with BAPS6 (a prior of 10 depth levels and a maximum of 20 clusters were specified) (Cheng et al. 2013). Nested clustering analyses were undertaken upon subsets of the original SNP alignment to a total depth of three levels or until no further clustering could be achieved.

Recombination and phylogenomic analysis. Recombination within the core genome was inferred using ClonalFrameML v1.7 (Didelot & Wilson 2015) using the whole genome alignment generated by Snippy. The ML tree generated with FastTree v2.1.8 was used as a guide tree for ClonalFrameML. Positions in the reference genome that were not present in at least one genome (non-core) were omitted from the analysis using the “ignore_incomplete_sites true”
option and providing ClonalFrameML with a list of all non-core positions. Maximum likelihood
trees with bootstrap support were constructed using a recombination free SNP alignment with
the program FastTree (Price et al. 2010). Bootstrap support was derived from comparisons
between the original tree against 1000 trees that were built upon pseudo-alignments (sampled
from the original alignment with replacement).

Pan genome analysis. Orthologous proteins were identified through reciprocal blast using
Proteinortho5 v5.11 (Lechner et al. 2011). A blast cutoff of 95% identity and alignment
coverage of 30% were used. The resulting matrix of ortholog presence and absence was
visualized using Fripan (https://github.com/drpowell/FriPan) (downloaded on the 28th of April
2016). The matrix of pan genome content has been deposited in Figshare

Sequence alignment and visualization. The alignment of homologous sequences was
undertaken using Mauve (Darling et al. 2004). Sequences and alignments were visualized using
Geneious Pro (version 8.1.8, Biomatters Ltd. [www.geneious.com]).

RESULTS AND DISCUSSION

Genome overview. Assembly of the 158,885 sequence reads from PacBio SMRT sequencing of
Ef_aus0233 (N50 read length 8,952 bp) resulted in reconstruction of a 3,272,427 bp genome,
comprising a circular chromosome and five circular plasmids (Table 1). Remaining 1bp
homopolymer insertion errors were corrected using Illumina reads. The structural integrity of the
chromosome assembly was confirmed correct by reference to a NcoI optical map (Fig. 1A).
DNA base modification analysis indicated an absence of adenine methylation.

Antimicrobial resistance gene content. One of the major drivers behind the success of E.
faecium in the clinical environment is their ability to acquire genes conferring antibiotic
resistance (Handwerger et al. 1993; Iwen et al. 1997; Murray 2000). In silico antibiotic
resistance screening of the Ef_aus0233 genome confirmed the presence of seven loci, conferring resistance to many major classes of antibiotics including trimethoprim and vancomycin (Table 2). Vancomycin resistance in Ef_aus0233 is conferred by the Tn1549 transposon, harbouring the vanB operon that has integrated into the chromosome. Here, Tn1549 was the larger (57 kB) of the two reported versions and inserted into signal peptidase 1 gene, an insertion site that was previously reported in a comparative analysis of ST203 genomes (Howden et al. 2013; Lam et al. 2013a). Of particular interest, the Tn1549 transposon in Ef_aus0233 and that of the fully assembled ST203 representative genome (Ef_aus0085) (Lam et al. 2013), shared 100% pairwise nucleotide identity across the full length of the element, implying a common Tn1549 origin for these two clones. The majority of ST796 isolates exhibit vancomycin resistance. However, three ST796 vancomycin susceptible enterococci (VSE) have been isolated to date, one of which (Ef_aus1016) is included in our study and is discussed later.

Virulence gene content. In addition to antimicrobial resistance genes, virulence related genes are particularly enriched among hospital adapted E. faecium strains and are thought to enhance fitness in the hospital environment (Rice et al. 2003). In silico comparative analysis (see methods) of the Ef_aus0233 genome revealed the presence of several genes associated with virulence including collagen-binding adhesin (Rice et al. 2003) (chromosome coordinates 2,235,651-2,233,486), enterococcal surface protein (Shankar et al. 1999; Van Wamel et al. 2007) (chromosome coordinates 2,786,822-2,780,895), hemolysin (Cox et al. 2005) (chromosome coordinates 1,025,987-1,027,363), all of which are also present in the fully assembled genomes of ST17 and ST203 isolates (Lam et al. 2012; Lam et al. 2013). The Ef_aus0233 enterococcal surface protein shared identity with the ortholog in Ef_aus0004, a fully assembled ST17 genome (Lam et al. 2012), however it had a 358 bp insertion compared to Ef_aus0085, a fully assembled ST203 genome (Lam et al. 2013). In comparison, esp from Ef_aus0004=4938nt (6 Rib repeats), Ef_aus0085=5199nt (7 Rib repeats), Ef_aus0233=5928nt (9 rib repeats). The hemolysin of Ef_aus0233 shared complete identity with orthologs in Ef_aus0004 and Ef_aus0085, suggesting that this CDS may be under strong selection. Several other genes encoding factors that associate with E. faecium virulence were absent in
Ef_aus0233 and included gelatinase (Hancock & Perego 2004), aggregation substance (Olmsted et al. 1991), hemagglutinin (Elsner et al. 2000) and hyaluronidase (Rice et al. 2003).

**Insertion sequence content.** The Ef_aus0233 chromosome was found to contain 80 distinct elements (9 families) while Ef_aus0233_p1 had 41 (6 families), Ef_aus0233_p3 had 8 (5 families), Ef_aus0233_p4 had 8 (2 families) and no IS elements were detected on Ef_aus0233_p2 or Ef_aus0233_p5. Several of these IS families have been found not only in enterococci but additionally in species of other genera, including *Carnobacterium* and *Lysinibacillus*, reflecting the ease to which *E. faecium* can acquire exogenous DNA (Guzman Prieto et al. 2016).

**CRISPR content.** Akin to an adaptive immune system, the clustered regularly interspaced short palindromic repeats (CRISPR) systems of prokaryotes function as a sequence-specific security to defend genomes against viral predation and exposure to invading nucleic acid (Horvath & Barrangou 2010). Unlike members of the community-associated *E. faecium* lineage, genomes belonging to the CC17 *E. faecium* have been found to lack CRISPR systems (van Schaik et al. 2010). Given the advantages associated with the acquisition of extraneous DNA carrying antimicrobial resistance genes, CRISPRs are thought to be under negative selection among multi-drug resistant enterococci (Palmer & Gilmore 2010). Despite this, two distinct CRISPR loci were detected on Ef_aus0233_p1 (chromosome coordinates 168197-168396) and Ef_aus0233_p2 (chromosome coordinates 2630-2860), both containing three spacers and imperfect direct repeats. A single CRISPR associated gene (cas2) was detected. However, no cas1 ortholog was detected. Due to the practical necessity of cas1 for the operation of CRISPR systems (Yosef et al. 2012), it is unlikely that these detected CRISPR systems are functional.

**Prophage content.** The Ef_aus0233 genome was found to contain five putative prophages. Prophages Ef_aus0233_chr_phage-1 (chromosome coordinates 260,208 - 308,229: 52 CDS), Ef_aus0233_chr_phage-2 (chromosome coordinates 916,578 - 956,914: 61 CDS), Ef_aus0233_chr_phage-3 (chromosome coordinates 2,366,359 - 2,425,249: 78 CDS) and
Ef_aus0233_chr_phage-4 (chromosome coordinates 2,601,769 - 2,627,494: 19 CDS) were located on the chromosome (Fig. 1B) while Ef_aus0233_p1_phage-1 (plasmid-1 coordinates 62,059 - 94,736: 30 CDS) was identified on Ef_aus0233_p1. Alignment of these prophage elements signified that several common blocks of co-linearity existed (Fig. S1), however an overall lack of prophage genome conservation implies that these phage represent five distinct elements. Prophage gene content among a diverse collection of *E. faecium* genomes is discussed below.

**Comparisons with other completed *E. faecium* genomes.** In addition to diversity within the core and accessory genome, structural rearrangements represent an additional layer of genomic variation that may contribute to *E. faecium* phenotypic differences (Lam et al. 2012; Lam et al. 2013; Matthews & Maloy 2010). To assess how the genomic organization of the Ef_aus0233 chromosome compared to that of other *E. faecium* genomes, a whole chromosome alignment of Ef_aus0233, Ef_aus0004, Ef_aus0085 and DO was undertaken. The BLASTn based alignment revealed substantial conservation of genome content (Fig. 1B) and chromosomal architecture (Fig. 2A). Like Ef_aus0085 and DO, Ef_aus0233 does not exhibit the replichore inversion observed in Ef_aus0004 (Lam et al. 2012; Lam et al. 2013).

**E. faecium population genomic comparisons.** In order to contextualize the 21 ST796 genomes within the global diversity of *E. faecium* as a species, we compared these genome data with a diverse collection of 89 published, fully assembled and draft *E. faecium* genomes (Table S1). To investigate the structure and evolutionary relatedness of the strains, we employed an unsupervised Bayesian clustering technique (BAPS) to distinguish distinct genomic populations and estimated a rooted phylogenomic tree using maximum likelihood.

Here, we found that our BAPS groups unambiguously classified genomes into the two previously reported A and B clades (Lebreton et al. 2013) (Fig. 3A). BAPS-1 corresponded with clade A and BAPS-2 corresponded with clade B, while BAPS-1, BAPS-1.1-4 and BAPS-1.5 overlapped with clades A1 and A2 (Fig. 3B). All ST796 isolates clustered within clade A1.
When inspecting the phylogeny of the ST796 genomes, it was noted that ST796 and ST555 share a most recent common ancestor (MRCA) (Fig. 3C). Almost without exception, ST555 and ST796 genomes formed distinct monophyletic clades, although the single vancomycin sensitive (VSE) ST796 included in this study clustered among the ST555 clade. Despite this phylogenomic incongruence, the BAPS groupings of ST555 and ST796 genomes (BAPS 1.3.3 and 1.3.4, respectively) were in harmony with the MLST designations. The ST555 clone is another recently-emerged hospital adapted ST, however its discovery in the hospital environment predates that of ST796 (Coombs et al. 2014). Another major difference between these two STs is that ST796 appears to have been localized to south east Australia and New Zealand (Carter et al. 2016; Coombs et al. 2014), while ST555 has been reported nationally in the Northern Territory, South Australia and Western Australia (Coombs et al. 2014), in China (Liu et al. 2011) and among wild birds in the United States (Oravcova et al. 2014). The phylogenetic position of the ST555 clade being basal to the VRE ST796 clade, suggests that VRE ST796 emerged from an ST555-like ancestral progenitor. The national and international pervasiveness of ST555 and relatively limited geographical dispersal of ST796 in southeast Australia is consistent with a scenario in which the evolutionary emergence of ST555 predates that of ST796.

Recombination analyses indicated that both the ST555-796 and ST796 MRCAs have evolved in part by recombination. Inspection of the inferred recombining segments for these two ancestors revealed a single hotspot of 170kb that contained two overlapping clusters of increased SNP density (ST555-796_MRCA: chromosome coordinates 1,783,249-1,953,029, ST796_MRCA: chromosome coordinates 1,857,926-1,937,284) (Fig. 2B). The spatial clustering of these inferred ancestral recombination events suggests that this region may contain non-clonally derived alleles (particularly those in the ST796 MRCA) that may have been under positive selection and perhaps contributed to the emergence of ST796. Inspection of this region revealed a preponderance of cell-wall associated transport CDS, including CDS encoding putative copper and cadmium-translocating P-type ATPases, amino acid permeases, OxaA-like
membrane protein, as well as housekeeping CDS such as Glycyl-tRNA synthetase subunits, RecO DNA repair proteins and a GTP-binding protein.

In order to assess the evolutionary divergence between the ST555 and ST796 clades, a ST555-796 specific core genome was established and pairwise SNP distances were calculated. As we were primarily interested in SNPs derived through clonal evolution, we removed SNPs within the above mentioned 170kb region of ancestral recombination. Inspection of SNP distribution in VSE Ef_aus1016 revealed several dense clusters, indicating substantial recombination, the majority of which was not detected by ClonalFrameML. As the recombination was so extensive and limited to Ef_aus1016, this genome was removed from the inter-clade comparisons.

Consistent with two distinct groups, within clade comparisons revealed smaller mean SNP differences (within-ST555: 11 SNPs, within-ST796: 49 SNPs) than that between clades (ST555-796: 151 SNPs) (Fig. 3D).

Ancestral single nucleotide polymorphisms. Forty-one core-genome SNPs differentiated the ST555-ST796 MRCA from its predecessors, while only two core-genome SNPs were predicted in the ST796 MRCA compared to ST555 (Table S2). Analysis of these SNPs showed a range of CDS impacted. Among the 41 SNPs, 22 were predicted to change amino-acid sequence and potentially alter protein function, including non-synonymous mutations in four CDS encoding putative regulatory proteins. While the function of these regulatory proteins and the consequence of the predicted mutations is unknown, such changes can have profound impacts on phenotype (Howden et al. 2011; Howden et al. 2008).

Accessory gene content comparisons. The clustering of all predicted CDSs into orthologous groups allowed for inter-ST comparisons at the gene-content level. In total there were 10,740 orthologous clusters among the 110 genomes, of which 1,437 were core and 9,303 were variably present (accessory) - representing the E. faecium pan genome (Table S3). Using this approach, orthologous clusters that were diagnostic of the ST555-ST796 and ST796 populations were identified.
Lineage specific genomic islands. The acquisition of genomic islands has been reported in previous studies that compared the genomes of hospitalized and non-hospital derived isolates, suggesting that such novel elements may offer possessing strains a competitive advantage (Heikens et al. 2008). In this study, subsets of the accessory genome that were associated with ST555-ST796 and ST796 genomes were found to cluster on the Ef_aus0233 chromosome (Fig. 2C). The contiguous location of these CDS and conserved inheritance patterns, indicated that these elements collectively formed larger genomic islands and were likely to have been acquired through horizontal gene transfer events. Given the ancestral relationships among the genomes and the conservation of ortholog presence amongst these lineages, it is reasonable to infer that these events occurred at various stages along the evolutionary paths of the ST555-ST796 and ST796 ancestries.

Two genomic islands were conserved among ST555-ST796 genomes while being almost entirely absent from genomes of other STs (ST555-ST796_GI-1, chromosome coordinates 39,093-53,122: 13 CDS, and ST555-ST796_GI-2 chromosome coordinates 2,316,643-2,373,309: 63 CDS), suggesting that these elements are likely to have been acquired by the ST555-ST796 MRCA. Assessment of the CDS annotation for ST555-ST796_GI-1 (56kb) suggests it is a mosaic integrative element. Two 3kb regions spanning a site-specific tyrosine recombinase and excisionase at the 5’ end of this element and replication proteins at the 3’ extreme were identical to a previously described Enterococcus faecalis pathogenicity island (Shankar et al. 2002). A 13kb region harbouring a putative beta-galactosidase and other sugar modifying CDS was identical to a region in ST203 Ef_aus0085 (Lam et al. 2013). The function of CDS in the remaining 33kb was more difficult to infer with few database matches to indicate function. However, a role for this region in cell wall modification (potentially DNA transfer) was suggested by the presence of CDS encoding cell-wall binding proteins, peptidases, ATP/GTP-binding proteins and peptidoglycan-binding proteins. ST555-ST796_GI-2 (14kb) is another integrative element with CDS encoding a site-specific tyrosine recombinase, replication proteins, and putative sugar kinases, hydrolases and permeases. This potential carbohydrate
utilization/transport locus shared complete nucleotide sequence identity with a region of the
The phage CDS content and their predicted products is provided (Table S4)

Two other elements were found to be exclusively present among ST796 genomes in this
comparison. One of these was the Tn916-like transposon (chromosome coordinates 635,179 -
658,601: 19 CDS), carrying tetracycline resistance (Franke & Clewell 1981) and the second was
the Tn1549 transposon (chromosome coordinates 803,567- 861,054: 61 CDS), carrying
vancomycin resistance (Garnier et al. 2000). Given the phylogeny, it appears likely that the
ST796 MRCA acquired these two transposons and then spread (Fig. 2C). The exception to this
pattern was the single VSE ST796 genome (Ef_aus1016) that lacked these elements that were
universally conserved among VRE ST796 genomes. Ef_aus1016 exhibits the same genomic
island presence/absence profile that is observed among the ST555 genomes. It appears that
Ef_aus1016 may actually be a ST555 genome that has recombined with ST796. This explains
why BAPS (when run on a non-recombination filtered alignment) clusters Ef_aus1016 as an
ST796 and why the phylogeny (which was built upon a recombination filtered alignment)
clusters this genome into the ST555 clade (Fig. 3C). This hypothesis also provides an explanation
as to why Ef_aus1016 exhibits an ST555-like genomic island presence and absence profile (Fig.
2C). An alternative explanation would be that Ef_aus1016 might be an extant descendent of a
ST796 evolutionary intermediate that had not yet acquired the ST796 specific GIs, such as
Tn1549. The horizontal acquisition of Tn1549 has been demonstrated (Launay et al. 2006) and
evidence for a VSE version of an emergent *E. faecium* clone to precede the VRE version has
been previously documented with the emergence of ST17, ST203 and ST252 (Johnson et al.
2010).

Prophage gene content comparisons. Alignment of the orthologs found within the prophages
that were identified in the Ef_aus0233 genome revealed the extent to which these elements
are conserved among the greater *E. faecium* population (Fig. 4). The vast majority of orthologs
within these prophages were found to exist in non-ST796 genomes, suggesting that at the gene-
In a content level, such prophages are not unique to ST796. Prophages Ef_aus0233_chr_phage-2, Ef_aus0233_chr_phage-4 and Ef_aus0233_p1_phage-1 showed the greatest degree of ortholog conservation with non-ST796 genomes. Prophages Ef_aus0233_chr_phage-1 and Ef_aus0233_chr_phage-3 did contain orthologs present in non-ST796 genomes, however the presence of these orthologs outside of ST796 genomes was limited. Overall the prophages in Ef_aus0233 form a substantial contribution to the accessory genome but do not contain CDS that are unique to ST796 (Table S4).

Plasmid gene content comparisons. Plasmids form an important component of the E. faecium accessory genome that can spread horizontally through a population and carry genetic elements that may confer enhanced fitness (Fiedler et al. 2016). Approximately 12% of the Ef_aus0233 genome (384,340 bp) is comprised of plasmid DNA. In order to assess the conservation of plasmid gene content among ST796 genomes and across the greater E. faecium population, the presence and absence of plasmid genes within the ortholog clusters were inspected (Fig. 5). Patterns of individual ortholog presence and absence demonstrated that all plasmid orthologs were found in non-ST796 genomes, however in varying degrees. Plasmids Ef_aus0233_p1, Ef_aus0233_p4 and Ef_aus0233_p5 contain orthologs found outside ST796, however they are rarely seen elsewhere in their entirety. A list of all plasmid CDS content and their predicted products is provided (Table S5).

Given these gene content patterns and the aforementioned phylogenomic relationships between ST555 and ST796 genomes, it appears likely that the ST555-ST796 MRCA acquired these plasmids, as they are not observed in their entirety in surrounding clades. Interestingly, Ef_aus0233_p2 was not only scarce among non-ST796 genomes but lacked conservation among the ST796 genomes. Overall, no single plasmid ortholog was specific to ST796, however these plasmids in their entirety are diagnostic of the ST555-ST796 lineage. Furthermore, the intra-ST796 differences in plasmid gene content, particularly in Ef_aus0233_p2, indicate there are appreciable amounts of diversity within the ST796 accessory genome, variation that might be useful during outbreak investigations involving this clone (Lister et al. 2015).
CONCLUSION

The hospital environment presents a challenging ecological niche for the adaptation of bacterial pathogens. Historically, *E. faecalis* was the leading causative agent of enterococcal nosocomial infections, however *E. faecium* infections have escalated in the last decade (Galloway-Peña et al. 2009; Guzman Prieto et al. 2016; Leavis et al. 2006; Willems et al. 2011; Willems & Van Schaik 2009). Following this apparent interspecies replacement, population-based studies have observed substantial intraspecies dynamics with clonal replacement of *E. faecium* STs in hospitals (Bender et al. 2016; Johnson et al. 2010). Here we have described the genomic basis for the emergence of a new highly hospital adapted *E. faecium* ST early in its evolutionary history. The preparation of a fully assembled ST796 genome facilitated a comprehensive genomic analysis of this lineage and enabled detailed comparisons among other clinically relevant draft and fully assembled *E. faecium* genomes.

We demonstrate that the emergence of ST796 was preceded by several genomic events including the acquisition of two genomic islands, plasmid and phage activity, modest SNV accumulation and recombination. These analyses highlight genetic elements within the *E. faecium* core and accessory genome that may have been important drivers for the evolution of the ST555-ST796 and ST796 lineages. Given the likely significance of genomic island acquisition for the emergence of CC17 (Heikens et al. 2008), the GIs presented in this study presumably reflect adaptive responses to the clinical environment, either through acquired antibiotic resistance or perhaps enhanced capacity to utilize carbohydrates and thus augment gastrointestinal colonization. Our finding that ST796 evolved from an ST555-like ancestral progenitor is another example of newly emergent VRE arising from a VSE MRCA, although ST796 itself is almost exclusively VRE when identified in human BSIs unlike ST555 which causes both VSE and VRE BSI in Australian hospitals in about equal proportions (Coombs et al. 2014).
This analysis focused upon providing an overview of the first fully assembled ST796 genome and genomic differences that were assessed at the inter-ST population level. In order to explore specific diversity within the ST796 lineage, an intra-ST population study focusing upon diversity among a large collection ST796 genomes is currently underway. Our observation of substantial variation within the ST796 accessory genome, in particular plasmid presence and absence, suggest a means for effective intra-ST796 genotyping that could potentially be more useful than core genome analysis in the tracking of outbreaks.

In this study we have described the genomic events that have shaped the evolution of *E. faecium* ST796. While the extent to which each genomic event has contributed to the ST796 emergence is not yet understood, our findings lay a foundation for testing specific hypotheses that have arisen from this work.

**REFERENCES**


10.1101/gr.2289704
14/7/1394 [pii]


Table 1: Characteristics of *E. faecium* ST796 Ef_aus0233 complete genome

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Length (bp)</th>
<th>% G+C</th>
<th>Copy number</th>
<th>No of CDS</th>
<th>No of non-paralogous CDS</th>
<th>No of tRNA</th>
<th>No of rRNA</th>
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<tbody>
<tr>
<td>Ef_aus0233_p1</td>
<td>197,153</td>
<td>35.4</td>
<td>1</td>
<td>210</td>
<td>197</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Ef_aus0233_p2</td>
<td>79,293</td>
<td>33.8</td>
<td>1</td>
<td>96</td>
<td>96</td>
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<td>-</td>
</tr>
<tr>
<td>Ef_aus0233_p3</td>
<td>77,977</td>
<td>35.2</td>
<td>1</td>
<td>84</td>
<td>84</td>
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<td>-</td>
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<tr>
<td>Ef_aus0233_p4</td>
<td>22,080</td>
<td>35.6</td>
<td>2</td>
<td>28</td>
<td>23</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Ef_aus0233_p5</td>
<td>7,837</td>
<td>33.5</td>
<td>8</td>
<td>8</td>
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<td>-</td>
</tr>
<tr>
<td>Resistance</td>
<td>Product</td>
<td>Gene</td>
<td>Location</td>
<td>Reference</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>(nucleotide positions)</td>
<td></td>
<td></td>
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<tr>
<td>Trimethoprim</td>
<td>Dihydrofolate reductase</td>
<td>dfrG</td>
<td>331,475 - 331,972 (chromosome)</td>
<td>Sekiguchi et al. 2005</td>
<td></td>
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<td>Tetracycline</td>
<td>Tetracycline resistance protein</td>
<td>tetM (Tn916)</td>
<td>652,734 - 654,653 (chromosome)</td>
<td>Burdett et al. 1982</td>
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<td>Macrolides</td>
<td>ABC transporter protein</td>
<td>msrC</td>
<td>2,711,468 - 2,712,946 (chromosome)</td>
<td>Portillo et al. 2000</td>
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<td></td>
<td>rRNA adenine N-6-methyltransferase</td>
<td>ermB</td>
<td>13,080 - 13,842 (plasmid 4)</td>
<td>Trieu-Cuot et al. 1990</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Aminoglycosides</td>
<td>Bifunctional aminoglycoside modifying enzyme</td>
<td>aac(6')-aph2''</td>
<td>60,698 - 62,008 (plasmid 1)</td>
<td>Patterson &amp; Zervos 1990</td>
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</tbody>
</table>

Table 2: Antibiotic resistance genes and mutations present in Ef_aus0233 and other ST796
**Fig. 1**: Ef_aus0233 chromosomal optical map and BRIG plot. (A) Optical map of the Ef_aus0233 chromosome. (B) Referenced based alignment of blast hits of Ef_aus0085, Ef_aus0004 and DO genomes against the aus0233 chromosome. Prophage elements and the Tn1549 VanB containing transposon are annotated as arcs in the outermost ring.
Fig. 2: Comparisons of chromosomal architecture, genomic islands and recombining segments associated with the ST555-796 and ST796 clades. (A) Alignment of fully assembled chromosomes of Ef_aus0233, Ef_aus0085, Ef_aus0004 and DO. (B) Recombining segments in the ST555-796 and ST796 MRCAs. (C) Core genome phylogeny aligned with gene content blocks for identified genomic islands. Colours indicate the MLST designations.
**Fig. 3:** Nested core genome SNP phylogeny of the greater *E. faecium* population. (A) *E. faecium* population tree containing the major division between the community and hospital associated clades (B and A). (B) Nested tree focusing on the sub-clade containing the ST555 and ST796 genomes. (C) ST555-796 pairwise core SNP differences. (D) Pairwise SNP comparisons of within and between core genome nucleotide diversity of the ST555 and ST796 clades. Y-axis depicts the number of SNP differences, error bars indicate one standard deviation above and below the mean and points represent the minimum and maximum values.
Fig. 4

**Fig 4:** Prophage gene content comparisons: the presence and absence of orthologs within each of the five phages that were identified in the Ef_aus0233 genome. The phylogeny depicts the evolutionary relationships among the genomes. Colours indicate the MLST designations.
Fig. 5: Plasmid gene content comparisons: the presence and absence of orthologs within each of the five plasmids that were identified in the Ef_aus0233 genome. The phylogeny depicts the evolutionary relationships among the genomes. Colours indicate the MLST designations.
Supplementary Fig. S1

Supplementary Fig. 1: Mauve alignment of prophages detected in the Ef_aus0233 genome.