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2013

de Been, M, van Schaik, W, Cheng, L, Corander, J & Willems, R J 2013, 'Recent Recombination Events in the Core Genome Are Associated with Adaptive Evolution in Enterococcus faecium' Genome Biology and Evolution, vol 5, no. 8, pp. 1524-U1. DOI: 10.1093/gbe/evt111

http://hdl.handle.net/10138/165535
https://doi.org/10.1093/gbe/evt111

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Recent Recombination Events in the Core Genome Are Associated with Adaptive Evolution in Enterococcus faecium

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Accepted: July 17, 2013

Data deposition: This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accessions AUWV00000000, AUWW00000000 and AUWX00000000. The versions described in this paper are AUWV01000000, AUWW01000000, and AUWX01000000.

Abstract
Reasons for the rising clinical impact of the bacterium Enterococcus faecium include the species’ rapid acquisition of adaptive genetic elements. Here, we focused on the impact of recombination on the evolution of E. faecium. We used the recently developed BratNextGen algorithm to detect recombinant regions in the core genome of 34 E. faecium strains, including three newly sequenced clinical strains. Recombination was found to have a significant impact on the E. faecium genome: of the original 1.2 million positions in the core genome, 0.5 million were predicted to have been affected by recombination in at least one strain. Importantly, strains in one of the two major E. faecium clades (clade B), which contains most of the E. faecium human gut commensals, formed the most important reservoir for donating foreign DNA to the second major E. faecium clade (clade A), which contains most of the clinical isolates. Also, several genomic regions were found to mainly recombine in specific hospital-associated E. faecium strains. One of these regions (the epa-like locus) likely encodes the biosynthesis of cell wall polysaccharides. These findings suggest a crucial role for recombination in the emergence of E. faecium as a successful hospital-associated pathogen.

Key words: BratNextGen, comparative genomics, phylogenomics, whole-genome sequencing, nosocomial pathogen, antibiotic resistance.

Introduction
Enterococci are Gram-positive bacteria that commonly inhabit the gastrointestinal-tract of healthy humans and other animals. However, since the 1980s, the clinical impact of enterococci has continuously increased and they now rank as one of the leading causes of nosocomial infections of the bloodstream, urinary tract, surgical wounds, and other sites (Arias and Murray 2012; Gilmore et al. 2013). In the past, enterococcal hospital-acquired infections (HAIs) were predominantly caused by E. faecalis, but since the 1990s, HAIs have been increasingly associated with E. faecium. Nowadays, E. faecium HAIs are almost as common as HAIs caused by E. faecalis (Hidron et al. 2008).

An important cause for the rise of E. faecium as a nosocomial pathogen has been the species’ capacity to easily acquire novel adaptive traits, including genetic elements encoding antibiotic resistance determinants. For example, nosocomial infections caused by ampicillin-resistant E. faecium were first reported in the United States in the 1980s, after which they spread rapidly around the world (Galloway-Peña et al. 2009). Currently, ampicillin resistance is reported in more than 80% of clinical E. faecium isolates worldwide (Hidron et al. 2008; Willems and van Schaik 2009). Other genetic elements that have been acquired by E. faecium include esp and other genes located on the esp pathogenicity island (ICEefm1), the putative hyaluronidase gene hyl, and several genes encoding surface proteins (Rice et al. 2003; Vankerckhoven et al. 2004; Coque et al. 2005; Klare et al. 2005; Camargo et al. 2006; Hendrickx et al. 2007; van Schaik et al. 2010), a putative phosphotransferase system that contributes to intestinal colonization during antibiotic treatment (Zhang et al. 2013), and specific insertion elements, especially IS16 (Leavis et al. 2007). The cumulative acquisition of these, but probably many more, genetic elements by E. faecium has resulted in the evolutionary development of a genetically distinct E. faecium subpopulation with a progressively increasing fitness in hospitalized patients.

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Recombination in the *E. faecium* Core Genome

(Willems et al. 2011). This subpopulation was initially designated lineage C1 (Homan et al. 2002) and later renamed to clonal complex 17 (CC17) on the basis of multilocus sequence typing (MLST) data analysis (Willems et al. 2005, 2011). Phylogenomic studies, using only a limited number of *E. faecium* strains, strengthened these previous findings as they revealed an ancient phylogenetic split in the *E. faecium* population, essentially dividing animal and human clinical isolates into one clade (clade A) and human commensal isolates into another clade (clade B) (Galloway-Peña et al. 2012; Palmer et al. 2012). However, in a recent study using Bayesian-based population genetic modeling of an MLST data set of more than 1,700 isolates, Willems et al. (2012) found that nosocomial *E. faecium* strains clustered into three, instead of one, subgroups, suggesting different evolutionary trajectories for modern clinical isolates.

Pan-genome analyses of *E. faecium* indicated that *E. faecium* has an open genome which means that the total available gene pool within this species is essentially unlimited (van Schaik et al. 2010). This also suggests that the *E. faecium* genome is highly plastic and few barriers exist for the acquisition of foreign genetic elements. Evidence for high levels of recombination was found in recent comparative genomes analyses which noted the existence of hybrid *E. faecium* strains, of which the genomes were an amalgam of clade A and clade B backgrounds (Galloway-Peña et al. 2012; Palmer et al. 2012). Similarly, MLST data analysis showed that certain *E. faecium* subpopulations displayed high levels of admixture, indicating that up to 14% of the MLST gene sequences originated from other *E. faecium* subpopulations (Willems et al. 2012). Furthermore, the majority of gene tree topologies of individual MLST genes were incongruent (Willems et al. 2005), which together with the above suggest that the impact of recombination on the population structure and evolution of *E. faecium* may be considerable. However, recombination in *E. faecium* has so far only been studied using either a limited set of strains or only few genes (i.e., seven core genes in the case of MLST data). Recent sequencing efforts have significantly increased the number of *E. faecium* whole-genome sequences (WGS) available in the public domain. These WGS provide an excellent opportunity to study recombination on a high-resolution whole-genome level. Here, we use the Bayesian Recombination Tracker (BratNextGen) algorithm (Marttinen et al. 2008, 2012; Castillo-Ramirez et al. 2012; McNally et al. 2013) to study the impact of recombination in *E. faecium* using a set of 34 WGS, including those of three clinical strains that were sequenced for the purpose of this study. We show that most of the recombination detected in clade A originated from a clade B type background and that purging recombinant signals from a genome-wide alignment prior to phylogenetic tree-building widened the ancient split between clade A and B. Moreover, we found that contemporary clinical isolates share specific recombination signals pointing to recombination-driven adaptations to selection pressures associated with hospitalization (e.g., antibiotic treatment).

**Materials and Methods**

**Sequence Information and Genome Sequencing**

WGS were retrieved from GenBank (ftp://ftp.ncbi.nih.gov/genomes/Bacteria, last accessed August 3, 2013 and ftp://ftp.ncbi.nih.gov/genomes/Bacteria_DRAFT, last accessed August 3, 2013). *Enterococcus faecium* WGS were downloaded on May 2012 and included 31 available genomes: 1 complete and 30 draft genomes. Available WGS of 66 other enterococci were downloaded in July 2012. Besides using publicly available WGS, three *E. faecium* isolates were newly sequenced for the purpose of this study. These included strains E155, E525, and E1165, which all belong to MLST ST17 (Homan et al. 2002; Willems et al. 2005) and have been isolated from hospitalized patients in the United States (Chicago), Australia (Melbourne), and Italy (Genoa), respectively (table 1). Genomic DNA was isolated from cell pellets of these strains using the Wizard Genomic DNA Purification kit (Promega, Leiden, The Netherlands) according to the manufacturer’s instructions. The strains were sequenced using Illumina GALL sequencing technology generating 50-bp paired-end reads from a library with an average insert size of 250 bp (Baseclear B.V., Leiden, The Netherlands). After confirming the quality of the raw illumina reads with FastQC v0.10.1 (Simon Andrews, Babraham Bioinformatics, http://www.bioinformatics.babraham.ac.uk/projects/fastqc/, last accessed August 3, 2013), reads were assembled de novo using Velvet v1.2.01 (Zerbino and Birney 2008). For each Velvet assembly, a range of different k-mer (hash) lengths (15–49 bp) was empirically tested to gain an optimal N50. In addition, Velvet was run with the options “-exp_cov” (expected k-mer coverage) and “-cov_cutoff” (coverage cutoff). The expected k-mer coverage (Ck) was calculated using \[C_k = \frac{C \times (L - k + 1)/L}{S}\], where \(L\) and \(k\) are the read and k-mer lengths (in bp), respectively. \(C\) is the estimated sequence coverage given by \([C = L \times N/S]\), where \(N\) is the number of reads and \(S\) is the estimated size of the genome. For the three genome assemblies, an estimated *E. faecium* genome size (\(S\)) of 2.75 Mb was used. The coverage cutoff was set at \(0.25 \times C_k\) as suggested by the Velvet developers (Zerbino DR, personal communication).

**Functional Annotation and Core Genome Prediction**

The Prokaryotic Annotation pipeline of ISGA (default settings) (Hemmerich et al. 2010) was used to annotate the three newly sequenced *E. faecium* genomes, as well as one of the available draft genomes (LCT-EF90) for which no annotation was available from GenBank. Annotated proteins were assigned to Clusters of Orthologous Groups (COG) (Tatusov Materials and Methods).
et al. 2001) as described previously (Snel et al. 2002). Protein domains were predicted using SMART v7.0 (Letunic et al. 2012) and Pfam v26.0 (Finn et al. 2010). To determine groups of orthologous proteins, the 97,972 annotated proteins present in the 34 \textit{E. faecium} strains were used as input for an all-versus-all sequence similarity search using BLASTP v2.2.24 (Altschul et al. 1997) (default settings, except for: \texttt{-F "m S"; -e 1.0E-05; -z 97,972 [i.e., the total number of proteins]}. From the BLAST output, groups of orthologous proteins were predicted using OrthoMCL v2.0.2 (Li et al. 2003). Orthologous groups with exactly one representative protein from each of the 34 input strains were considered to be part of the \textit{E. faecium} core genome.

## Detection of Recombination

The BratNextGen algorithm (Marttinen et al. 2008, 2012) was used to detect recombination events in \textit{E. faecium}. A
concatenated multiple sequence alignment of the *E. faecium* core genome was used as input for this algorithm. This concatenated alignment was built as follows: for each orthologous group, the corresponding nucleotide sequences were extracted and aligned using Muscle v3.7 (Edgar 2004), after which gaps were stripped from the alignment using trimAl v1.2 (Capella-Gutierrez et al. 2009). Stripped alignments were concatenated in the order of orthologous group numbers as assigned by OrthoMCL. Because OrthoMCL assigns orthologous group numbers on a “first-come, first-served” basis, orthologous genes ended up in the concatenated alignment in the order in which they were read by OrthoMCL, thus following the order in which they are located on the genomic contigs and scaffolds. As the first genomic contigs that entered OrthoMCL were those of strain 1,141,733, the alignment followed the exact gene order in the contigs of this strain. Analysis of the alignment showed that the gene order used also exactly matched that of 11 other strains and nearly exactly matched that of the remaining 22 strains (a median of only 3 synteny breaks was observed for these 22 strains with a maximum of only 10 and 11 synteny breaks for strains Aus0004 and 1,231,408, respectively). Thus, the concatenated alignment of orthologous genes accurately reflects the synteny of the average *E. faecium* genome. The estimation of recombination was performed with the default BratNextGen settings as in (Marttinen et al. 2012) starting out with a partition of the genome alignment into 5 kb blocks and using 20 iterations of the estimation algorithm, which was assessed to be sufficient since changes in the hidden Markov model parameters were already negligible over the last 30% of the iterations. The alpha parameter was estimated using the default options in BratNextGen and the resulting value was 4.96. Significance of a recombination region was determined as in (Marttinen et al. 2012) using a permutation test with 100 permutations executed in parallel on a cluster computer (threshold of 5% was used to conclude significance for each region).

Phylogeny and the Identification of Recombinant Sequence Origins

Phylogenetic trees were built using RAxML v7.2.8 (Stamatakis 2006) under the GTRCAT model. To assess the effect of recombination on the evolution of *E. faecium*, two phylogenetic trees were built: one using the aligned SNPs from the concatenated core genome alignment, and one using the same information, but filtered for significant recombination signals as determined by BratNextGen. Confidence was inferred by running 1,000 bootstrap replicates under the GTRCAT model. The phylogenetic origin of the identified recombinant sequences was assessed as follows: first, for each recombination event, the corresponding sequence alignment was extracted from the larger *E. faecium* core genome alignment and a separate phylogenetic subtree was built. Recombinant sequences were classified as belonging to the same recombination event when they had received exactly the same recombination start and end positions by BratNextGen. To prevent obscuring the individual subtrees with signals from other recombination events, all sequences that partially or completely belonged to another recombination event were removed from the alignment prior to building the tree (note that recombination events do not overlap within one sequence). Second, the resulting trees were midpoint rooted using the Phangorn package v1.6-5 (http://cran.r-project.org/web/packages/phangorn/index.html, last accessed August 3, 2013) in R (R Development Core Team, http://www.R-project.org, last accessed August 3, 2013), after which a decision scheme was followed using the topology of the first major split closest to the root (supplementary fig. S1, Supplementary Material online). In the whole-genome recombination-filtered tree, *E. faecium* was separated into the two clades A and B (see also Results and Discussion). This knowledge was taken into account in the decision scheme, such that recombinant sequences were predicted to have originated from either an (ancestral) clade A or clade B strain. Alternatively, when the recombinant sequences clustered separately from both clades (i.e., as an outgroup), the sequence was predicted to have originated from a species other than *E. faecium*. Because BratNextGen uses a conservative approach for predicting recombinant sequences to avoid false positives (Marttinen et al. 2012), it is likely that not all recombination events were detected. Therefore, undetected overlapping recombination events could still obscure individual subtrees and blur the separation into clades A and B. To take this into account, a maximum of two supposedly nonrecombinant sequences were deleted from the tree if needed to get a “pure” clade A/B separation, thereby keeping a minimum of three clade A and three clade B strains in the tree. If the tree could not be purified in this way, the phylogenetic source of the recombinant sequence was considered to be either *E. faecium* or inconclusive (supplementary fig. S1, Supplementary Material online).

Analysis of Recombinant Sequences Originating from Outside *E. faecium*

In cases where recombinant sequences were predicted to originate from outside *E. faecium*, an attempt was made to identify the potential donor species. For this, recombinant sequences were split up into their corresponding gene sequences (partial gene sequences <30 bp were not considered). Recombinant gene sequences were then screened against GenBank’s nucleotide sequence database (nt) (ftp://ftp.ncbi.nlm.nih.gov/blast/db/, last accessed August 3, 2013; date of download August 2012) and against the assemblies of 66 non-*E. faecium* enterococci (discussed earlier) using BLASTn v2.2.24 (default settings, except for: -F F). Target sequences were recovered using an E-value cutoff of 1.0E-10 and a query sequence coverage of \( \geq 75\% \). Recovered
sequences were aligned with the corresponding *E. faecium* gene sequences, after which midpoint rooted individual gene trees were built using RAxML. Trees were purified as described above if necessary and were inspected manually to assign potential donor species.

Database Submission

The whole genome sequence data of *E. faecium* strains E155, E525, and E1165 have been deposited at DDBJ/EMBL/GenBank under the accessions AUWX00000000, AUWW00000000, and AUWW00000000, respectively.

**Results and Discussion**

*Enterococcus faecium* Phylogeny Based on the Core Genome

To study the effects of recombination on the genome evolution of *E. faecium*, we used the genome sequence information of 34 *E. faecium* strains, including 31 publicly available strains and 3 strains that were newly sequenced for the purpose of this study. The newly sequenced strains were E155, E525, and E1165 and were all modern clinical isolates, originating from hospitalized patients in the United States, Australia, and Italy, respectively, and belonging to ST17, which is an important node in the hospital-associated subpopulation of *E. faecium* (Homan et al. 2002; Willems et al. 2005, 2011, 2012) (table 1). De novo assembly of the sequence reads resulted in an average genome size of 2.94 Mb (±0.14 Mb) and a mean scaffold N50 of 43.4 kb (±2.9 kb). The number of scaffolds of size ≥200 bp ranged from 185 to 242 (supplementary table S1, Supplementary Material online). An annotation of the new genomes using the ISGA pipeline resulted in an average set of 2,971 (±170) protein-encoding genes per genome.

A total of 97,927 protein sequences were extracted from the 34 *E. faecium* genomes and used as input for assessing the core genome. This resulted in a set of 5,075 orthologous groups, of which 1,512 were considered to belong to the core genome as they contained exactly one protein representative from each of the 34 input strains. After stripping the individual core gene alignments for gaps, all alignments were concatenated to generate a whole-genome core alignment of 1,223,137 bp. From this alignment, we extracted 85,488 SNPs, which were used for the construction of a phylogenetic tree. The topology of the resulting tree (fig. 1A and supplementary fig. S2, Supplementary Material online, for details on bootstrap support) was in agreement with what has been observed before by others who used an *E. faecium* core genome extracted from 8 to 29 isolates to build a phylogenetic tree (Galloway-Peña et al. 2012; Lam et al. 2012; Palmer et al. 2012; Willems et al. 2012). The tree displayed a deep phylogenetic split between two major *E. faecium* clades. These clades were previously observed by Palmer et al. (2012), who designated them clades A and B, and by Galloway-Peña et al. (2012), who designated them clades HA (hospital-associated) and CA (commensal/community-associated), respectively. Clade A contained most of the clinical and hospital-associated isolates as well as animal isolates, whereas clade B consisted mainly of human commensal isolates. The clinical isolate 1,231,408 clustered consistently (100% bootstrap support) with clade A, albeit relatively distant from all other clade A strains. This is also in line with previously built phylogenetic trees and with the observation that 1,231,408 is a hybrid strain that appears to have received a large part of its genome from a clade B background (Palmer et al. 2012) (discussed later). As expected on the basis of their ST (table 1), the newly sequenced clinical isolates clustered closely together with other modern clinical isolates of the same ST, as well as with strain C68 (ST16), which is a single locus variant of ST17 (Homan et al. 2002).

*Enterococcus faecium* Phylogeny Based on a Recombination-Filtered Core Genome

To estimate the effects of recombination on the evolution of *E. faecium*, we used the algorithm BratNextGen (Marttinen et al. 2008, 2012). Running this algorithm on the *E. faecium* core genome alignment resulted in the prediction of 1.2 Mb of recombinant sequence, divided over 383 recombination events (supplementary fig. S3, Supplementary Material online). Of the original 1,223,137 positions in the core genome alignment, 534,961 (44%) were predicted to be affected by recombination in at least one strain. On the SNP level, this rate was comparable with 38,665 of the original 85,488 positions (45%) identified as being recombinant. The remaining 46,823 nonrecombinant SNPs were used to build a recombination-free phylogenetic tree. As shown in figure 1B (supplementary fig. S2, Supplementary Material online, for details on bootstrap support), the topology of this tree was largely similar to that of the unfiltered tree presented in figure 1A in the sense that the ancient split between clade A and B was still observed after purging for recombinant. Interestingly, BratNextGen did not detect any recombination signals in clade B. This does not necessarily mean that clade B strains are completely unaffected by recombination. BratNextGen uses a conservative approach for predicting recombinant sequences to avoid false positives (Marttinen et al. 2012). Consequently, true recombination signals may go undetected (i.e., false negatives) when there is low statistical power. The reasons for the absence of detectable recombination in clade B may be that 1) clade B strains were underrepresented in the available WGS data set used here, and 2) the diversity within clade B was relatively large as compared to the diversity within clade A (fig. 1A). Because BratNextGen has higher statistical power to detect significant recombination events when they are shared across strains, these factors (i.e., limited sample of strains and high diversity between them) may have hampered the detection of recombinant
sequence segments within clade B. Indeed, we noticed that, when using our decision scheme for predicting recombinant sequence origins (supplementary fig. S1, Supplementary Material online), tree purifications were made 74 times corresponding to the removal of 103 sequences. This observation may point to a recombinant nature for the corresponding sequences although other reasons for sequence divergence (i.e., mutation) cannot be ruled out. Of the 103 removed sequences, 60 belonged to clade B strains, mostly to strains LCT-EF90 (33 cases) and E980 (23 cases), thus potentially indicating the presence of recombinant sequences in clade B. Nevertheless, the fact that BratNextGen detected no recombination signals at all in clade B at least strongly suggests that, in general, clade A is more prone to recombination than clade B. Consequently, the recombination-filtering procedure effectively meant a purification of clade A only. As became evident by visually comparing the trees in figure 1A and B, the purification resulted in a strongly increased cohesiveness of the phylogenetic tree for clade A. To quantitatively support this observation, we extracted all strain-to-strain pairwise branch-length distances from both trees and compared the average distance within clade A and B and between both clades, pre- and post-recombination-filtering. Potential oversampling of lineages was taken into account by including only one strain (the one furthest from the root) for branches where all pairwise strain distances were less than 5.0E–04 in the purified tree (e.g., strain TX0133a01 was picked as a representative for all 5 TX0133 strains). This analysis showed that upon recombination-filtering, the average distance within clade A significantly decreased (from 0.1 to 0.03; P < 0.01; Wilcoxon signed rank test), whereas the average distance between clade A and B significantly increased (from 1.4 to 2.0; P < 0.01). The average distance within clade B (0.2) remained the same. In summary, recombination-filtering widened the ancient clade A/B split by bringing the clade A strains closer together.

Comparison of the two trees showed that most of the strains that clustered tightly together in the unfiltered tree also clustered tightly together upon recombination-filtering. The most notable exceptions were 1) the separation of strains E1039 and 1,231,501 from each other and from the larger cluster containing for instance strain E1679, 2) the shift of the 5 × TX0133 branch to a position within the ST17 cluster of modern clinical isolates, and 3) the shift of the branch
1,231,502/U0317 to a position just outside the ST17 cluster (fig. 1 and supplementary fig. S2 [Supplementary Material online] for details). These last two modifications resulted in a robust branch consisting only of modern clinical isolates of ST17 (and 16) and with ST78 and ST203 isolates together forming a distinct lineage. Another major change that occurred upon recombination-filtering was the displacement of strain 1,231,408 from a “hybrid” position in-between clade A and B toward a position branching deeper inside clade A and tightly clustering with strain 1,230,933, which clustered with strain DO in the unfiltered tree. The strong shift of strain 1,231,408 toward clade A and away from clade B fits with the previously observed hybrid nature of this strain (Palmer et al. 2012). However, a similar post-recombination-filtering displacement of strain 1,231,408 was not observed in a previous study (Qin et al. 2012), in which the authors used PHI (Bruen et al. 2006) to detect and remove recombination signals prior to phylogenetic tree building. In their resulting tree, strain 1,231,408 still occupied a “hybrid” position similar to the one observed in the unfiltered tree presented here (fig. 1A).

Quantities of Recombination in Clade A

Recombination signals were detected in each of the 27 clade A strains included in this study. However, the amount of detected recombinant DNA as a percentage of the 1.2 Mb core genome varied widely over the different clade A strains (fig. 1C). When correcting for potential over-sampling of genetic lineages, the average amount of recombination detected in clade A was 4.4% of the core genome. The strain with the highest amount of recombination was strain 1,231,408 with as much as 26.9% of its core genome predicted to be recombinant. Other strains with a relatively high load of recombination were E1039 (10.1%) and 1,231,501 (8.2%). At the other end of the spectrum were strains TX0133a01, E1162, and TX0082 with only 0.3%, 0.7%, and 0.7% of their core genome predicted to be of foreign origin, respectively (fig. 1C). Interestingly, the modern clinical isolates of ST16 and ST17 had a significantly lower amount of recombination (average of 1.1%) than the other strains in clade A (6.0%) (P < 0.01; Mann–Whitney U test). This finding appears to be in agreement with a previous observation made by Willems et al. (2012), who applied a genetic admixture analysis on MLST sequence data from a large collection of E. faecium strains and found that hospital isolates displayed lower levels of recombination than isolates from nonhospitalized persons and pigs. This may point to genetic and/or ecological isolation of E. faecium hospital isolates and fits with the previously proposed hypothesis that novel, highly invasive, E. faecium strains may arise through horizontal gene transfer, but once they have adopted their new lifestyle, they become genetically isolated from other bacteria and become less prone to recombination (Willems et al. 2012).

The Recombinant DNA in Clade A Mainly Had a Clade B Origin

Interestingly, the vast majority of the E. faecium recombinant sequence was predicted to have originated from a clade B type E. faecium background (fig. 1C). More specifically, we found that of the 1.2 Mb of recombinant sequence, 71% had a predicted clade B type source, followed by 19.6% with a predicted source from outside E. faecium and 7.6% with a predicted clade A type origin. For only 0.1% of the recombinant sequence, it was unclear whether it originated from a clade A or clade B type background. Also, for only 1.7% of the recombinant sequence, the phylogenetic source was inconclusive according to the decision scheme that we employed (supplementary fig. S1, Supplementary Material online). These results indicated that clade B E. faecium strains have formed, and possibly still form, an important source for donating core genomic DNA to clade A E. faecium strains. Especially strains 1,231,408, E1039, and 1,231,501 were found to have received a large amount of core genomic DNA from a clade B type source, totalling 327, 103 and 84 kb, respectively. Also with respect to the total amount of foreign DNA detected in their core genomes, the amount of DNA with a potential clade B type source in these strains was high at 99%, 84%, and 84%, respectively. In figure 2, we plotted the detected recombinant sequences and their potential sources onto the core genome sequence of the completed strain Aus0004, as determined in this study. The results presented in figure 2 expand on previous results of Palmer et al. (2012), who studied genome mosaicism in five clade A and three clade B E. faecium strains. For each gene in these eight strains they determined whether it had either a clade A- or clade B-type signature. Mosaicism was most evident for two clade A strains they analyzed, namely strains 1,231,408 and 1,231,501. Here, when using the currently much larger data set of 34 E. faecium genomes, we found that strains 1,231,408 and 1,231,501 still ranked among the top strains with respect to the amount of recombinant sequence coming from a clade B type background. In accordance with the observations made by Palmer et al. (2012), we found that these recombinant sequences clustered into a few long genomic stretches taking up around 27% and 7.8% of the 1,231,408 and 1,231,501 core genomes, respectively (fig. 2).

Recombinant DNA with a Predicted Source Outside E. faecium

As mentioned earlier, almost 20% of the recombinant core genomic DNA in E. faecium was predicted to come from a source outside E. faecium. To find potential donor species for these recombinant segments, we extracted the corresponding gene sequences from each recombinant segment and screened them against a database of enterococcal genome sequences (Materials and Methods) and GenBank’s nt database. Similar sequences were extracted and used, together
with the corresponding recombinant and nonrecombinant E. faecium gene sequences, to build phylogenetic trees (Materials and Methods). The resulting trees were manually inspected to find close homologs of (i.e., potential donor species for) the recombinant E. faecium sequences. In cases where one recombination event covered multiple genes, all the resulting gene trees had to be in agreement with each other concerning potential donor species. If not, for instance because one of the recombinant genes did not yield BLAST hits outside E. faecium, the donor species was deemed unknown. Unfortunately, we did not find potential donor species for the majority (84 out of 111) of the corresponding recombination events. This was due to the fact that for these 84 events, no similar sequences were detected for at least one of the recombinant genes, other than E. faecium sequences themselves. The most likely explanation for obtaining so few BLAST hits may be that current microbial sequence databases simply do not (yet) contain sequences from the actual donor species or from species closely related to these donors. Still, for the remaining 27 recombination events (24%), we did find more specific evidence concerning the nature of the potential donor species. For instance, for 25 out of the 27 recombination events, the recombinant gene sequences clustered in between E. faecium and other enterococcal species or clustered together with multiple different enterococcal species other than E. faecium, suggesting that these recombinant sequences have originated from the genus Enterococcus (results not shown). Finally, for two recombination events, we found even more specific evidence about the nature of the potential donor species. One of these events was found in strain C68 and involved genes efxg_00252–efxg_00259. Each of these genes clustered together with a gene from E. hirae ATCC 9790 (ehr_06405–ehr_06440, respectively), suggesting that E. hirae or a closely related species has been the responsible recombination donor for these genes. Similarly, we found that gene efme1071_1769 in strain E1071 had an E. faecalis signature. These observations indicate that there is also detectable recombination between E. faecium and other species in the genus Enterococcus thereby significantly enlarging the potential reservoir for genetic diversity in E. faecium.

**Functional Analysis of Recombinant Genes**

To describe the functional impact of recombination on the E. faecium genome, we analyzed whether certain functional gene categories were more affected by recombination than others. To this aim, we assigned COGs to each orthologous protein group in the E. faecium genome. We then compared the composition of the COG functional categories found in
the recombinant regions versus those found in the recombinant-free regions. After correcting for potential over-sampling, we found that 1,832 proteins were affected by recombination compared with 40,502 recombinant-free proteins, corresponding to 2,086 and 45,654 assigned COG functional categories, respectively. As shown in supplementary figure S44, Supplementary Material online, a linear regression analysis between the COG category compositions of these two data sets showed a strong linear correlation ($R^2 = 0.87$), suggesting that in general most functional groups are equally prone to recombination in the *E. faecium* core genome. Still, when we used Grubb’s test for outliers using a 99% confidence level, we found that COG categories C (energy production and conversion) and S (function unknown) were identified as outliers, potentially being relatively more (category C) and less (category S) prone to recombination in the *E. faecium* core genome (supplementary fig. S44, Supplementary Material online). Further analysis of the categories C and S showed that their skewed representation in recombinant DNA was not due to specific COGs that were over- or underrepresented, but was caused by a proportional change of all COGs in the entire category (results not shown).

By a similar approach, we also evaluated whether specific functional categories originated from specific recombination donors. Supplementary figure S4B–D, Supplementary Material online, shows the COG distributions for the recombinant genes originating from one of the three major donors of recombinant DNA (clade A, clade B, and outside *E. faecium*) compared with the COG distributions for the recombinant genes originating from all donors, except the one of interest. This analysis revealed that recombinant genes belonging to COG category G (carbohydrate transport and metabolism) mainly originated from clade A. Two individual COGs from category G that were overrepresented in recombinant DNA from clade A, were COG2723 and COG1486, which both encode putative beta-glucosidases. Together these two COGs made up 62% of all recombinant COGs of category G originating from clade A, whereas they only made up 9% of all recombinant COGs of this category originating from other sources. Other functions that originated mostly from clade A included a putative sugar uptake permease (COG4975), a dihydroxyacetone kinase (COG2376), and an ABC type polysaccharide transporter (COG1134) (results not shown). These results suggest that genes involved in sugar uptake and metabolism are extensively transferred between clade A *E. faecium* strains. This is of interest because genes involved in these processes may be important for successful colonization of the gut during antibiotic treatment, as has been demonstrated recently (Zhang et al. 2013).

Functional Analysis of Recombination Hotspots

To locate the position of conserved major recombinant regions, we searched for recombination events that were detected in at least two strains (correcting for potential over-sampling) with a total summed length of at least 5 kb. This revealed a set of 18 recombination events (listed in supplementary table S2, Supplementary Material online), which merged into eight regions of overlapping and closely flanking recombination events (fig. 2). Interestingly, six of these eight regions (regions 2–7), including the four largest ones with a length ranging from 26 to 59 kb, were located in the third quadrant of the Aus0004 core genome. These data indicate a nonrandom distribution of recombination events over the *E. faecium* core genome and suggest the existence of a hot-spot for recombination in *E. faecium*, which, when translating to the Aus0004 genome, roughly lies between open reading frames efau004_01354 and efau004_02178. Besides identifying recombination events that were conserved across multiple isolates, we searched for sites in the *E. faecium* core genome with clear traces of recombination-driven gene acquisition. For this, we screened for genomic regions of at least five consecutive accessory genes that were flanked on both sides by the same recombination event. The accessory genes and the flanking recombinant core genes had to be located on one single contig or scaffold. A total of 17 recombination events met these criteria, several of which were clearly associated with mobile genetic elements, such as phages, transposons, recombinases, and integrases (supplementary table S2, Supplementary Material online). Of the 17 events, five were found in strain 1,231,408, corresponding to the first quadrant of the Aus0004 core genome. Still, 9 of the remaining 12 events were found within the recombination hotspot described earlier, which further indicates that this region on the *E. faecium* genome is relatively prone to recombination.

Recombination of an Ampicillin Resistance Determinant

As mentioned earlier, we found that the clinical *E. faecium* strains of ST16/17 displayed relatively low levels of recombination. Considering this, it is of interest to note that 6 of the 18 conserved recombination events listed earlier were associated with ST16/17 strains, indicating that the recombination-driven diversity of these genomic regions may reflect adaptation to selection pressures associated with hospitalization. Four of the corresponding recombination events (events 8–11, supplementary table S2, Supplementary Material online) were located within region 5 (fig. 2). Among these were events 8 and 9, which were predicted by BratNextGen to be independent recombination events, but which covered exactly the same set of core genes. Together, these two events were found in all ST16/17 strains analyzed, except TX0133. Other events that partly overlapped with events 8 and 9 were found in strains E4452 and E1071 (results not shown). One of the genes within the affected region was *ddcP* (efau004_01934 in Aus0004), which has previously been identified as a membrane-associated α-alanyl-α-alanine carboxypeptidase that confers resistance to ampicillin and
Recombination in the *E. faecium* Core Genome

lysozyme in *E. faecium* E1162 (Zhang et al. 2012). Recombinant *ddcP* was predicted to stem from an ancestral clade B type *E. faecium*. On the protein level, three positions gave recombinant DdcP a clear clade B type signature, including an Ile-to-Val at position 101, an Asp-to-Ala at position 298 and an Asp-to-Glu at position 392 (supplementary fig. S5, Supplementary Material online). Although these amino acid variations do not by themselves fully explain DdcP-mediated ampicillin resistance in *E. faecium* (e.g., strain E1071 and all clade B strains are ampicillin sensitive), they do suggest a scenario in which clinical strains, like E1162, have recombinated their *ddcP* gene with a clade B type *ddcP* gene, which in combination with other factors like mutations in *ppb5*, has resulted in clinical strains becoming more prone to developing ampicillin resistance.

**Recombination of the epa-Like Locus Is Reminiscent of Pneumococcal “Capsule-Switching”**

The two remaining recombination events that were associated with ST16/17 *E. faecium* strains were both found in region 2 (fig. 2) and were predicted to have originated from an unidentified source outside *E. faecium*. Both events covered and flanked almost the same set of core and accessory genes, respectively, namely *efau004_01354* to *efau004_1385* in strain Aus0004 and its orthologs in strains E1165 and E525 (event 2, supplementary table S2, Supplementary Material online) and a somewhat larger region extending toward the orthologs of *efau004_01387* in strains 1,231,410, E155, 1,230,933, DO, 1,231,502, and U0317 (event 3). In Aus0004, the region between *efau004_01354* to *efau004_1387* mostly consists of accessory genes, including two ISL3-family transposases and a retroviral integrase, further pointing toward a history of lateral gene transfer for this complete genomic region. The region involved has previously been described as a downstream extension of the *E. faecium* epa-like locus (Palmer et al. 2012; Qin et al. 2012), a gene cluster that is putatively involved in the biosynthesis of antigenic cell wall polysaccharide, based on homology with the *E. faecalis* epa locus (Xu et al. 1998; Teng et al. 2009). The extended region of the epa-like locus (corresponding to *efau004_01365–efau004_01382* or *efau004_01384* in Aus0004) is known to be highly variable across *E. faecium* and has been divided into four locus variants (Qin et al. 2012). Analysis of the current set of 34 *E. faecium* strains showed that the same division into four locus variants still holds. Figure 3 shows representatives of these four locus variants as well as the regions affected by recombination. Notably, all the epa-like loci affected by recombination events 2 and 3 belonged to locus variant four. Only one epa-like locus of this type, the one present in the clade B strain E980, did not contain recombination signals according to BratNextGen. This does not necessarily mean that this locus has not recombined in this strain, but may suggest that the potential recombination event was not recent enough, so that any strong recombination signals may have been neutralized over time. Besides recombination events 2 and 3, additional independent events were detected within the same genomic region of strains E1165, E4453, E1679, E4452, E1636, and E1039, but these events covered only small segments within epaMNO, never extending beyond three consecutive core genes (results not shown). Nevertheless, the observation that multiple independent recombination events were detected in the downstream epa-like extension, suggests that this region of the *E. faecium* genome is prone to recombination. As shown in figure 3, the variable epa-like region that was flanked and thus potentially affected by events 2 and 3 contained a predicted class C beta-lactamase gene (*efau004_01384* in Aus0004) and sialic acid biosynthesis genes *neuABC* (*efau004_01366* and *efau004_01370–efau004_01372*) specific for locus variant four. Sialic acid decoration of the bacterial cell surface may be a form of molecular mimicry that affects detection by the host’s immune system (Vimr and Lichtensteiger 2002). Interestingly, events 2 and 3 both extended beyond the epa-like locus into a core genomic region containing yet another class C beta-lactamase gene (*efau004_01360*). A recombinant cell surface biosynthesis cluster flanked by two beta-lactamase genes has also been described in *Streptococcus pneumoniae*. In this species, serotype variants have arisen through recombinational replacements within and around this so-called cps locus (Trzcinski et al. 2004; Brueggemann et al. 2007; Croucher et al. 2011; Golubchik et al. 2012; Martinen et al. 2012). Our observations suggest that recombination may have a role in capsule variation in *E. faecium*.

**Conclusions**

In this study, we have analyzed the impact of recombination on the whole-genome level in *E. faecium*. We show that filtering for recombination prior to phylogenetic tree building had a purifying effect on the ancient *E. faecium* clade A/B split. Closer inspection of the origin of the recombinant sequences indicated that this purifying effect was mainly caused by the fact that clade A strains have received foreign DNA mainly from clade B type strains. However, no recombination signals were detected in clade B strains themselves, suggesting that the transfer of DNA within the *E. faecium* population is mainly a unidirectional event: from clade B to clade A, and not vice versa. However, it is important to note that the number of clade B strains in our data set was relatively low, which may have hampered the detection of recombination signals within this clade. The inclusion of more clade B strains in future recombination detection analyses will identify whether the apparent lack of recombination within clade B is genuine or is merely a result of a lack of resolution. Still, the results presented here strongly indicate that clade A is more prone to recombination than clade B.

Given the observation that recombination seems to act as a coalescent force for clades A and B, it is of interest to note...
that, within clade A, contemporary clinical isolates of ST16/17 contained relatively low levels of recombination. This fits with previous hypotheses (Willems et al. 2012; McNally et al. 2013) that novel, highly successful, clinical strains may arise through gene acquisition, but become genetically isolated and less prone to recombination once they have adopted their new lifestyle. Still, several recombination events were associated with contemporary clinical isolates. These events involved the ampicillin resistance determinant \( \text{ddcP} \) and the putative cell wall polysaccharide biosynthesis \( \text{epa} \)-like locus. These findings point to important recombination-driven mechanisms that facilitate adaptation of \( E. \text{faecium} \) to selection pressures associated with hospitalization.

### Supplementary Material

Supplementary figures S1–S5 and tables S1 and S2 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org).

### Acknowledgments

This work was supported by the European Union Seventh Framework Programme (FP7-HEALTH-2011-single-stage) “Evolution and Transfer of Antibiotic Resistance” (EvoTAR) (grant number 282004 to M.B., W.S., and R.J.W.;) the European Research Council (ERC) (grant number 239784 to J.C.;) the Academy of Finland (grant number 251170 to J.C.;) and the Sigrid Juselius Foundation (grant number 4702281 to J.C.). The Centre for Molecular and Biomolecular Informatics (CMBI, Radboud University Nijmegen, Nijmegen, the Netherlands) is acknowledged for providing computing space. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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