Incidence of Type II CRISPR1-Cas Systems in Enterococcus Is Species-Dependent

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Abstract

CRISPR-Cas systems, which obstruct both viral infection and incorporation of mobile genetic elements by horizontal transfer, are a specific immune response common to prokaryotes. Antiviral protection by CRISPR-Cas comes at a cost, as horizontally-acquired genes may increase fitness and provide rapid adaptation to habitat change. To date, investigations into the prevalence of CRISPR have primarily focused on pathogenic and clinical bacteria, while less is known about CRISPR dynamics in commensal and environmental species. We designed PCR primers and coupled these with DNA sequencing of products to detect and characterize the presence of cas1, a universal CRISPR-associated gene and proxy for the Type II CRISPR1-Cas system, in environmental and non-clinical Enterococcus isolates. CRISPR1-cas1 was detected in approximately 33% of the 275 strains examined, and differences in CRISPR1 carriage between species was significant. Incidence of cas1 in E. hirae was 73%, nearly three times that of E. faecalis (23.6%) and 10 times more frequent than in E. durans (7.1%). Also, this is the first report of CRISPR1 presence in E. durans, as well as in the plant-associated species E. casseliflavus and E. sulfureus. Significant differences in CRISPR1-cas1 incidence among Enterococcus species support the hypothesis that there is a tradeoff between protection and adaptability. The differences in the habitats of enterococcal species may exert varying selective pressure that results in a species-dependent distribution of CRISPR-Cas systems.

Introduction

Bacteria and Archaea possessing CRISPR-Cas systems trade off horizontally-acquired adaptation to a changing environment for protection against lethal virus infection. CRISPRs are clustered regularly interspaced short palindromic repeats of DNA; Cas refers to CRISPR-associated proteins. Together, they comprise a uniquely prokaryotic multi-step adaptive immune response that provides defense against bacteriophage infection [1]. In the process, incorporation of transmissible genetic elements is interrupted, including plasmids and DNA with potential advantages for the host cell, such as those conferring antibiotic resistance [2]. Briefly, fragments of non-self DNA called protospacers are acquired by Cas proteins, and incorporated...
as spacers between the DNA repeats of the CRISPR array. These repeat-spacer modules are transcribed and expressed as crRNAs, a small interference-type RNA. If invading nucleic acid has a short sequence with perfect complementarity to the spacer region of the crRNA, a sequence-specific cleavage event is initiated, degrading the foreign nucleic acids [3,4]. CRISPR arrays are widespread among Bacteria and Archaea, in approximately 90% of archaeal and 40% of bacterial genomes examined [5,6]. The diversity of CRISPR systems is extensive. CRISPRs may be broadly divided into those lacking cas genes, thus consisting solely of repeat-spacer arrays (also referred to as orphan CRISPRs), and those comprised of both an array and associated functional genes (CRISPR-Cas). CRISPR-Cas systems are further divided into types and subtypes, defined by presence of subtype-specific Cas proteins [7]. Several Cas proteins are considered universal, with orthologs appearing in every active subtype. One of these is Cas1 [7,8]. Encoded by a single gene (cas1), the ubiquity of Cas1 makes it a suitable marker for the presence of a potentially active CRISPR-Cas system.

We focused on CRISPR1 systems in the genus *Enterococcus*, a clade of commensal bacteria common to animal and human gut microflora. Enterococci emerged as a cause of multidrug-resistant hospital acquired infection in the 1970s, and presently represent one of the most prevalent causes of nosocomial infections in the United States [9]. Two species—*E. faecalis* and *E. faecium*—are primarily responsible for these infections [10]. They are also the predominant enterococcal human gastrointestinal (GI) commensals [11]. Mobile elements, including plasmids, pathogenicity islands, and antibiotic resistance genes, comprise as much as 25% of the genomes of hospital-adapted lineages of both species [12,13,14]. Palmer and Gilmore (15) showed that multiple drug resistance and incidence of CRISPR-Cas are negatively correlated in *E. faecalis* and *E. faecium*. That is, their results suggest that there is a tradeoff between acquisition of drug resistance and CRISPR-mediated protection from foreign DNA. Three Type II CRISPRs have been identified in human GI *E. faecalis*: two with associated cas genes (CRISPR1-Cas and CRISPR3-Cas) and one orphan repeat-spacer array (CRISPR2) [15]. CRISPR2 is present in 95% of *E. faecalis* isolates; as many as half of these strains contain CRISPR1-Cas, and CRISPR3-Cas has been detected in four *E. faecalis* genomes to date [15,16]. This suggests that species under different selective pressures may vary significantly in their incidence of CRISPR.

Several studies have investigated CRISPR in clinical and virulent enterococci, but few have addressed the prevalence of these systems in environmental and commensal strains [16,17,18,19]. Additionally, CRISPR content in *E. faecalis* and *E. faecium* has been extensively reported, but a comprehensive survey including other *Enterococcus* species is lacking [15,17,18,20,21]. Since antiviral protection by CRISPR-Cas also prevents incorporation of potentially beneficial genes, retention of a CRISPR locus represents a tradeoff between protection and adaptability. To test the hypothesis that different habitats affect this tradeoff and thus the prevalence of CRISPR, our objective was to determine the frequency of active Type II CRISPR1 systems in *Enterococcus* species. Environmental, non-clinical enterococci were screened for presence of the conserved CRISPR1-cas1 gene, as a marker for the active CRISPR locus most commonly detected in this genus. CRISPR1-cas1 was detected in multiple *Enterococcus* species, including several not previously characterized as containing CRISPR systems. Significant differences in cas1 incidence between species were also observed.

**Methods**

**Enterococcus strains**

*Enterococcus* isolates were cultured from activated sludge, oxygenated wastewater from residential and industrial sources, including storm runoff. Other samples included soil and
sediment, compost, vegetation, marine and freshwater sources, and canine, feline, and avian fecal specimens (S1 Table). No permits were required for the described study, which complied with all relevant regulations. Water, soil, sediment, plant clippings, and fecal samples were taken from public properties where permission was not required, or from private property with permission of the owners. Activated sludge samples were supplied by water treatment plant supervisory personnel.

Activated sludge was diluted to 1:1000, and 10 mL of the dilution was filtered through 0.22-μm pore-size membrane filters, then incubated on mEnterococcus agar (Difco) at 35°C for 24 hours. Isolated colonies were selected from the agar, and streaked for isolation of pure cultures on Enterococcosel agar (BBL). Environmental and fecal samples were enriched by incubation in azide dextrose broth for 24 hours at 35°C, followed by isolation of pure cultures on Enterococcosel agar. Additional Enterococcus strains from beach sand were isolated as previously described [22].

Enterococcus faecalis OG1RF (ATCC 47077), which contains a CRISPR1 locus, was selected as a positive control [20]. The strain was purchased from the American Type Culture Collection (Manassas, VA).

All isolates were Gram-positive, catalase-negative cocc. Species identity of all isolates was determined by 16S rRNA sequence match in the Ribosomal Database Project (http://rdp.cme.msu.edu/index.jsp), and identities were verified by 16S rRNA phylogenetic analysis. Isolate cas1 sequences were confirmed to be Enterococcus cas1 genes by BLASTn (NCBI) sequence match against the nucleotide collection (nr/nt) database.

Identification of CRISPR components in available genome sequences

Enterococcus cas1 genes for primer design were identified by BLASTn of the NCBI nucleotide collection (nr/nt) database, using the E. faecalis OG1RF cas1 sequence (accession number CP002621.1) as the query (Fig A in S1 Text). CRISPR repeat-spacer arrays, and cas genes in proximity to the arrays, were investigated in 13 available Enterococcus genomes in CRISPRdb (E. casseliflavus EC20, accession number CP004856.1; E. faecalis 62, CP002491.1; E. faecalis D32, CP003726.1; E. faecalis OG1RF, CP002621.1; E. faecalis str. Symbioflor 1, HP558530.1; E. faecalis V583, AE016830.1; E. faecium Aus0004, CP003351.1; E. faecium Aus0085, CP006620.1; E. faecium DO, CP003583.1; E. faecium NRRL B-2354, CP004063.1; E. hirae ATCC 9790, CP003504.1; E. mundtii QU 25, AP013036.1; Enterococcus sp. 7L76, FP929058.1 [5]. Additional draft genomes (E. durans ATCC 6056, accession number GCA_000406985.1; E. faecium FB129-CNAB-4, GCA_000315405.1; E. durans I PLA 655, GCA_000350465.1) were downloaded from GenBank and analyzed for CRISPR content using CRISPRfinder (Table A in S1 Text) [6].

PCR and sequencing

Nucleic acid extractions were performed using the MoBio UltraClean® Microbial DNA Isolation Kit. The variable region of the 16S rRNA gene was amplified using universal bacterial DNA primers, forward, 5'–CTACGGGAGGCAGCAG–3' ; reverse, 5'–GCTACGGGAGGCAGCAG–3' [23].

To screen isolates for CRISPR1-cas1, primers amplifying a 212-bp internal region of the cas1 gene (forward, 5'–ATGGGCTGGCGAAC–3' ; reverse, 5'–ATTACCGCGGCTGCTGG–3' ) were used. Multiple alignment of Enterococcus CRISPR1-cas1 nucleotide sequences available at that time (E. faecalis OG1RF, accession number CP002621.1; E. faecalis D32, CP003726.1; E. hirae ATCC 9790, CP003504.1) was performed by MUSCLE [24,25] to locate conserved regions of the cas1 homologs. Primers were designed manually, and their compatibility was...
confirmed using Primer3 (http://bioinfo.ut.ee/primer3/) [26,27]. Primers were deemed compatible, as Tm differed by 0.75°C, and no complementarity (self, pair, and primer hairpin) was detected. Target specificity of the primer set was further confirmed by Primer BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/) against all Enterococcus (taxid: 1350), using all variations of the reverse primer, which contains a degenerate base. The primer set amplified in silico in *E. faecalis* OG1RF, *E. faecalis* D32, and *E. hirae* ATCC 9790. Amplification was optimized for the following program: 2 minutes at 94°C, 30 cycles of [1 minute at 94°C, 1 minute at 48.9°C, 1 minute at 72°C], 10 minutes at 72°C.

PCR products were submitted to Massachusetts General Hospital DNA Sequencing Core Facility or Eton Biosciences, Boston, MA for sequencing. Sequences were curated manually, and 16S rRNA gene sequences were deposited in GenBank (S1 Table).

### Analysis and phylogeny

To test whether CRISPR1-cas1 distribution significantly differed by species or source, data were analyzed by Chi square and Fisher’s exact tests (Tables 1–3).

Phylogeny was constructed using SeaView 4 (http://doua.prabi.fr/software/seaview). Multiple sequence alignment was performed within Seaview 4 using MUSCLE [25], and gap-only sites were removed. A maximum likelihood tree (PhyML) was generated, using the GTR model and aLRT branch support, with all other parameters set to default (nucleotide equilibrium frequencies: empirical; Ts/Tv ratio: fixed, 4.0; invariable sites: none; across site rate variation: optimized; tree searching operations: NNI; starting tree: BioNJ, optimized tree topology). FigTree 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree) was used for tree visualization. Two of the sequences used to design the cas1 primers were used as reference sequences in the cas1 phylogenetic tree; *E. faecalis* D32 was omitted, as it is identical to that of *E. faecalis* OG1RF.

| Source                  | cas1-positive | No. of isolates | % cas1- positives |
|-------------------------|---------------|-----------------|-------------------|
| Activated sludge        | 40            | 131             | 30.5              |
| Environmental samples   | 38            | 113             | 33.6              |
| Animal fecal            | 12            | 31              | 38.7              |

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| Source                  | cas1-positive | No. of isolates | % cas1- positives |
|-------------------------|---------------|-----------------|-------------------|
| Activated sludge        | 9             | 38              | 23.7              |
| Environmental samples   | 17            | 69              | 24.6              |
| Animal fecal            | 0             | 3               | -                 |

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| Source                  | cas1-positive | No. of isolates | % cas1 positives |
|-------------------------|---------------|-----------------|-----------------|
| Activated sludge        | 29            | 39              | 74.4            |
| Environmental samples   | 16            | 25              | 64.0            |
| Animal fecal            | 12            | 14              | 85.7            |

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Results

The predominant Enterococcus species isolated were E. faecalis (40.0% of 275 total isolates), E. hirae (28.4%), E. durans (20.4%), and E. faecium (5.1%). Additional enterococcal species were isolated less frequently, and include E. casseliflavus, E. sulfureus, E. mundtii, E. malodoratus, E. termitis, and E. sanguinicola (Table 4).

The CRISPR1-cas1 gene was detected in 32.7% of all Enterococcus isolates (Table 4). Within the three most predominant species isolated, frequency of cas1 detection varied significantly. The incidence of CRISPR1-cas1 genes between E. faecalis, E. durans, and E. hirae is significantly different (Table 4; p < 0.0001). The frequency of remaining species was not considered in this analysis due to small sample size. CRISPR1-cas1 was detected in 23.6% of E. faecalis isolates, while 73.1% of E. hirae and 7.1% of E. durans strains contain the gene. Cas1 was also detected in isolates of E. faecium, E. casseliflavus and E. sulfureus. The few strains of E. malodoratus, E. sanguinicola, E. mundtii, and E. termitis that were isolated did not contain cas1 (Table 4). The origin of the bacterial strain and presence of a CRISPR1-cas1 gene were not significantly correlated. This observation was consistent for all Enterococcus species analyzed, as well as intraspecific analyses of the two most commonly isolated species, E. faecalis and E. hirae (Tables 1–3). A phylogenetic tree of partial cas1 sequences formed two strongly distinct clusters around the E. faecalis OG1RF and the E. hirae ATCC 9790 cas1 reference sequences (Fig 1). All but 4 of the 26 E. faecalis cas1 genes clustered with the E. faecalis OG1RF-like cas1 gene. The remaining four strains of E. faecalis (MWRA37, MWRA22, 176T, and 158T) contained an E. hirae-like cas1 homolog. All identified E. hirae strains possess a cas1 homolog similar to that of E. hirae ATCC 9790. Cas1 sequences for E. casseliflavus, E. faecium, and E. sulfureus share identity with the E. hirae gene. E. durans strains contained cas1 genes homologous to both the E. faecalis OG1RF and E. hirae ATCC9790 cas1 types.

Cas1 sequences are conserved in the region amplified in this study, and the E. hirae and E. faecalis homologs are distinctly different from each other, perhaps reflecting species-level evolution. Within this region, the sequences differ by 16 transitions, 20 transversions, and a 3 bp indel, and not a continuum of differences between the two clusters (Fig 2). E. faecalis strains usually contain an E. faecalis cas1 homolog, and E. hirae-like cas1 genes typically appear in strains identified as E. hirae. Additionally, three of four E. durans cas1-positive strains contain E. hirae homologs, but one contains an E. faecalis-like gene. Horizontal transfer of CRISPR components in enterococci has yet to be demonstrated.

Table 4. Detection of CRISPR1-cas1 in Enterococcus, by species. Differences in CRISPR1-cas1 detection between E. faecalis, E. hirae, and E. durans isolates are significant, P value < 0.0001. Species for which a low number of strains were isolated are indicated in italics.

| Species          | Cas1-positives | Total isolates | Percent cas1 positive |
|------------------|----------------|----------------|-----------------------|
| E. faecalis      | 26             | 110            | 23.6                  |
| E. hirae         | 57             | 78             | 73.1                  |
| E. durans        | 4              | 56             | 7.1                   |
| E. faecium       | 1              | 14             | 7.1                   |
| E. casseliflavus | 1              | 7              | 14.3                  |
| E. sulfureus     | 1              | 2              | 50.0                  |
| E. mundtii       | 0              | 2              | 0.0                   |
| E. sanguinicola  | 0              | 1              | 0.0                   |
| E. malodoratus   | 0              | 4              | 0.0                   |
| E. termitis      | 0              | 1              | 0.0                   |
| **Total**        | **90**         | **275**        | **32.7**              |

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Incidence of cas1 in Enterococcus

This study is the first systematic analysis of Type II CRISPR1-Cas incidence in non-clinical enterococci. The incidence of CRISPR1-associated cas1 in *E. hirae* (73.1%) is significantly higher than in other species. Fig 1 shows the phylogenetic tree of CRISPR1-cas1 partial sequences, highlighting the E. faecalis-like cas1 cluster in red branches and the E. hirae cas1 cluster in blue branches.

**Fig 1.** Phylogenetic tree of CRISPR1-cas1 partial sequences. Red branches represent the *E. faecalis*-like cas1 cluster; blue branches represent the *E. hirae* cas1 cluster.

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**Discussion**

**Incidence of cas1 in Enterococcus**

This study is the first systematic analysis of Type II CRISPR1-Cas incidence in non-clinical enterococci. The incidence of CRISPR1-associated cas1 in *E. hirae* (73.1%) is significantly higher than in other species. Fig 2 compares partial CRISPR1-cas1 sequences of representative isolates (*E. hirae* MWRA15 and *E. faecalis* AS003) and reference strains (*E. hirae* ATCC 9790 and *E. faecalis* OG1RF) using MUSCLE. Bases conserved between all analyzed sequences are indicated with asterisks; spaces denote transitions and transversions, and dashes represent indel regions.

**Fig 2.** Comparison of partial CRISPR1-cas1 sequences. Representative isolates (*E. hirae* MWRA15 and *E. faecalis* AS003) and reference strains (*E. hirae* ATCC 9790 and *E. faecalis* OG1RF) were aligned using MUSCLE. Bases conserved between all analyzed sequences are indicated with asterisks; spaces denote transitions and transversions, and dashes represent indel regions.

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Cas1 phylogeny indicates horizontal transfer of CRISPR loci

The tight clustering of the partial cas1 sequence phylogeny was striking. Therefore, the four strains of E. faecalis that clustered with the E. hirae cas1 sequences indicates horizontal transfer of CRISPR elements between Enterococcus species (Fig 1). CRISPR1-cas1 genes identified in E. sulfureus, E. casseliflavus, and E. faecium cluster with the cas1 homologs in E. hirae strains (Fig 1). This is further indication of horizontal transfer, or CRISPR-Cas systems may be conserved with high levels of sequence similarity between these species. A more comprehensive description of the CRISPR-Cas systems in E. durans, E. faecium, E. casseliflavus, and E. sulfureus is needed to answer this question, as well as to shed light on differences in cas genes and array content that may explain interspecific CRISPR diversity.

CRISPR1 in E. durans, E. casseliflavus, and E. sulfureus

This is the first report of the presence of CRISPR1-Cas systems in E. durans, E. casseliflavus, and E. sulfureus. E. durans is a minor component of human and animal gut flora, and is also found in food of animal origin, especially dairy products [11,28]. Lack of virulence genes, including those that confer antibiotic resistance, indicate a probiotic role for E. durans [30]. CRISPR1 incidence in E. durans is low, but phage pressure in typical habitats of this species are not well characterized. E. casseliflavus and E. sulfureus are primarily plant-associated species [31,32,33]. Recent studies of E. casseliflavus have implicated the bacterium in human infection; however, these cases remain infrequent [34,35,36,37]. Reports implicating E. sulfureus in human disease could not be found in scientific literature. The rarity with which these species are pathogenic suggests an inverse correlation between virulence and CRISPR content, as was demonstrated in Escherichia coli [15,38,39]. Accurate frequencies of CRISPR1 loci in these species will require more comprehensive testing. In this study, only a few isolates of these species were cultured and screened for the cas1 gene.

CRISPR1-cas1 was not detected in isolates of E. mundtii and E. malodoratus. CRISPR loci have not been reported in two E. mundtii genomes previously analyzed, and incidence in E. malodoratus has also not been reported [40,41]. However, these sample sizes are too small to conclude that these species do not possess CRISPR1 loci. Additionally, the Type II-specific cas1
primers used in this study are unlikely to amplify all cas1 genes within Enterococcus, as species may contain CRISPR-Cas systems of different types [42]. With the three additional species reported here to contain CRISPR1-cas1, six species of Enterococcus are reported to possess CRISPR. But, as many as 40 other Enterococcus species have yet to be investigated [43]. Although more thorough characterization is warranted, the presence of cas genes in the species reported here indicates that CRISPR1-Cas systems may be widespread among the Enterococcus genus. The primers designed here successfully amplified a conserved region of the cas1 gene in multiple enterococcal species, making it an efficient marker for screening for CRISPR1 loci. Furthermore, widespread incidence of active CRISPRs and omnipresence of the clade in many environments make Enterococcus an ideal model for investigation of CRISPR dynamics.

Conclusions

Immunity against lytic phages is a recognizable evolutionary benefit for a bacterium, demonstrated both experimentally and in mathematical models of CRISPR-Cas/phage interaction [1,44]. Often considered as beneficial, indiscriminate insertion of foreign genetic elements, such as genomic islands, prophages and plasmids, on the other hand, can result in disruption of essential gene function and incorrect regulation of acquired genes [45]. CRISPR-mediated prevention of these detrimental insertions may also confer an evolutionary advantage [2]. However, horizontally-acquired genes may increase fitness by conferring habitat adaptations. Such adaptations in Enterococcus include antibiotic resistance, enhanced biofilm formation, resistance to metal toxicity, and expanded metabolic capacity [46,47]. Maintaining a functional CRISPR-Cas system also incurs an energetic cost for the organism [45]. Thus, for the bacterium possessing CRISPR loci, there is a tradeoff between adaptability and protection. Significant differences in Type II CRISPR1-cas1 incidence seen here indicate that selective pressures exerted by this tradeoff may influence CRISPR-Cas distribution in a species-dependent manner. The nature of this selection remains an area of future research.

Supporting Information

S1 Fig. Phylogenetic tree of Enterococcus isolates. (TIF)
S1 Table. 16S rRNA gene sequences and accession numbers deposited in GenBank. (DOCX)
S1 Text. Supplemental Methods. (DOCX)

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Author Contributions

Conceived and designed the experiments: CL MS. Performed the experiments: CL NR MAB. Analyzed the data: CL MS. Contributed reagents/materials/analysis tools: MS. Wrote the paper: CL MS.
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