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Interference with histidyl-tRNA synthetase by a CRISPR spacer sequence as a factor in the evolution of *Pelobacter carbinolicus*

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Abstract

Background: *Pelobacter carbinolicus*, a bacterium of the family *Geobacteraceae*, cannot reduce Fe(III) directly or produce electricity like its relatives. How *P. carbinolicus* evolved is an intriguing problem. The genome of *P. carbinolicus* contains clustered regularly interspaced short palindromic repeats (CRISPR) separated by unique spacer sequences, which recent studies have shown to produce RNA molecules that interfere with genes containing identical sequences.

Results: CRISPR spacer #1, which matches a sequence within *hisS*, the histidyl-tRNA synthetase gene of *P. carbinolicus*, was shown to be expressed. Phylogenetic analysis and genetics demonstrated that a gene paralogous to *hisS* in the genomes of *Geobacteraceae* is unlikely to compensate for interference with *hisS*. Spacer #1 inhibited growth of a transgenic strain of *Geobacter sulfurreducens* in which the native *hisS* was replaced with that of *P. carbinolicus*. The prediction that interference with *hisS* would result in an attenuated histidyl-tRNA pool insufficient for translation of proteins with multiple closely spaced histidines, predisposing them to mutation and elimination during evolution, was investigated by comparative genomics of *P. carbinolicus* and related species. Several ancestral genes with high histidine demand have been lost or modified in the *P. carbinolicus* lineage, providing an explanation for its physiological differences from other *Geobacteraceae*.

Conclusions: The disappearance of multiheme *c*-type cytochromes and other genes typical of a metal-respiring ancestor from the *P. carbinolicus* lineage may be the consequence of spacer #1 interfering with *hisS*, a condition that can be reproduced in a heterologous host. This is the first successful co-introduction of an active CRISPR spacer and its target in the same cell, the first application of a chimeric CRISPR construct consisting of a spacer from one species in the context of repeats of another species, and the first report of a potential impact of CRISPR on genome-scale evolution by interference with an essential gene.

Background

Clustered regularly interspaced short palindromic repeats (CRISPR), which consist of direct repeats of a short sequence (21-47 bp) separated by nonrepetitive sequences of similar size, have been identified in the genome sequences of almost all archaea and numerous bacteria, with a variable complement of adjacent CRISPR-associated (*cas*) genes [1-9]. A fraction of the spacer sequences between repeats have been found to match sequences termed "proto-spacers" within genes, from which they may be derived [8,10,11], and the fact

that many of these genes belong to phage or plasmid entities led to the hypothesis that CRISPR and the Cas proteins may function as an RNA interference-based immune system [6]. The link between specific CRISPR spacers and proto-spacers and phage resistance has been established by mutational analysis in *Streptococcus thermophilus* [12,13], and by testing synthetic CRISPR constructs in *Escherichia coli* [14]. Similarly, resistance of *Staphylococcus epidermidis* to a conjugative plasmid has been shown to depend on a CRISPR spacer and the corresponding proto-spacer [15]. Expression of CRISPR loci as long transcripts processed into smaller RNA molecules has been observed in several archaea [5,16-19] and bacteria [14,20]. A complex of Cas proteins has been

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shown to carry out this processing in *E. coli* and to be required for resistance to infection [14]; a different protein (Cas6) processes CRISPR transcripts in *Pyrococcus furiosus* [21]. CRISPR-derived RNAs have been shown to form RNA-protein complexes in *P. furiosus* [19], which leads to degradation of RNAs containing matching proto-spacers [22], whereas DNA was shown to be the target of interference by spacer-containing RNAs in *S. epidermidis* [15]. Although CRISPR are widely regarded as an immunological phenomenon, CRISPR and *cas* genes have also been implicated in spore development of *Myxococcus xanthus* [3,23] and in inhibition of biofilm formation and swarming of *Pseudomonas aeruginosa* by a lysogenic phage [24], and there has been speculation that spacers with matches to housekeeping genes represent a novel mechanism of gene regulation [25].

The *Geobacteraceae*, a Fe(III)-respiring family of *Deltaproteobacteria*, are of interest for their role in bioremediation of U(VI)-contaminated environments and their ability to donate electrons directly to graphite electrodes, producing an electrical current [26,27]. *Pelobacter carbinolicus* is a member of the *Geobacteraceae* that grows by fermentation of acetoin and 2,3-butanediol, as well as by indirect Fe(III) respiration with ethanol as the electron donor and acetate as the end product [28,29]. Unlike its relatives in the genus *Geobacter*, *P. carbinolicus* cannot reduce Fe(III) directly in the absence of sulfur or sulfide [30], or produce electricity [31]. The genome of *P. carbinolicus* was sequenced for the purpose of comparison to those of *Geobacter* species, three of which have been extensively curated: *Geobacter sulfurreducens* [32], *Geobacter metallireducens* [33] and *Geobacter bemidjensis* (Aklujkar *et al.*, submitted). This report explores how evolution of the *P. carbinolicus* genome may have been influenced by a spacer within the CRISPR locus that matches a proto-spacer within histidyl-tRNA synthetase (*hisS*), resulting in the elimination of ancestral genes containing multiple closely spaced histidines. The interfering nature of the spacer was confirmed by introducing it to a transgenic *G. sulfurreducens* strain containing the target gene.

Methods

Analysis of CRISPR spacers

The CRISPR locus was identified when manual curation of the *P. carbinolicus* genome revealed a series of suspiciously repetitive predicted genes. The nonredundant nucleotide sequence database was queried with each of the 111 CRISPR spacers of *P. carbinolicus* using the BLAST algorithm [34], with the minimum possible word size of 7 bp and without filtering out low-complexity regions of the queries. Alignments with five or fewer mismatches out of 32 bases were considered significant.

Phylogenetic analysis of HisS and HisZ proteins

The sequences of all predicted *hisS* gene products of the *Geobacteraceae*, together with HisS and HisZ protein sequences representative of various families of Bacteria and Archaea, were aligned by Toffee [35] and trimmed using Mesquite (Maddison, W. P., and Maddison, D. R. 2006. Mesquite: a modular system for evolutionary analysis. Version 1.12). Phylogenetic trees were constructed using Phylip (Felsenstein, J. 2005. PHYLIP (Phylogeny Inference Package) version 3.6) with 500 bootstrap runs.

Quantitative real-time PCR of reverse-transcribed RNA

P. carbinolicus strain DSM2380 was grown as previously described [36] with ethanol as the electron donor and Fe(III) as the electron acceptor. RNA was isolated from triplicate chemostat cultures as previously described [37,38]. Transgenic *G. sulfurreducens* strains were grown in NBAF medium [39] and RNA was isolated from actively growing triplicate batch cultures at an OD₆₀₀ of 0.20 to 0.31. The absence of DNA contamination was confirmed by PCR as previously described [36] with primer pairs specific for CRISPR spacer #1, for *hisS* and for *hisZ* (Table 1), using *P. carbinolicus* or *G. sulfurreducens* genomic DNA (isolated with the MasterPure DNA Purification Kit from EPICENTRE Biotechnologies, Madison, WI) as a control. Six to twelve clones of each genomic DNA PCR product were sequenced to verify the specificity of the primers. Reverse transcription was performed with the Enhanced Avian First Strand Synthesis Kit (Sigma-Aldrich, St. Louis, MO) as described previously [37], using each primer individually at 2 μM concentration with 400 ng of RNA in 20 μl total volume. Successful reverse transcription and the feasibility of DNA amplification in the presence of RNA were verified by PCR using 5 μl of this reaction. Quantitative real-time PCR (QRT-PCR) was performed with two to four technical replicates (9.5 μl of a tenfold dilution of cDNA, corresponding to 19 ng of RNA) for each of three biological replicates in a Taqman 7500 instrument using 2 × Power SYBR Green PCR master mix (Applied Biosystems, Foster City, CA) and primer pairs at 9 nM concentration in 25 μl total volume, for 50 cycles with an annealing temperature of 60°C and triplicate standards of spacer #1 and *hisS* PCR products from *P. carbinolicus* genomic DNA and a *hisZ* PCR product from *G. sulfurreducens* genomic DNA, encompassing four orders of magnitude.

Recombinant DNA techniques

All restriction enzymes were purchased from New England Biolabs; LA Taq polymerase was from Takara Mirus Bio; plasmids were propagated in *E. coli* TOP10 cells from Invitrogen; DNA purification kits for plasmids and agarose gel slices were from QIAGEN, and the

Table 1 Oligonucleotides for QRT-PCR and genetic manipulations.

Primers for QRT-PCR			
Name	Purpose	Location	Sequence (5' to 3')
MA0326	PCR of spacer #1	spacer #2	CCTGGTTGAGGTTAGCGTTGA
MA0327		outside CRISPR	AATTCGGTGGCCAGTTGTTTC
MA0328	PCR of <i>hisS</i> Pcar_1041	sense strand	CAGGAAGCCACCAAGGAT
MA0329		antisense strand	TGGGAGCCGAGTTGATTG
MA0441	PCR of <i>hisZ</i> GSU3307	sense strand	CAAACCTGATTGCCGTTCCCTT
MA0442		antisense strand	AGGCCGATGAGTTCTACGC
Primers for construction of <i>hisS</i> transgenic strain MA159			
Name	Purpose	Sequence (5' to 3')	
MA0330	PCR on 5' side of <i>hisS</i> GSU1659	TGACATCTCGCTGGACCGGG	
MA0331		CTATGCTAGCACTAGTTTGAATCATGAACGTACCTACTC CTTTAATTG	
MA0332	PCR on 3' side of <i>hisS</i> GSU1659	GTACGTTTCATGATTACAACTAGTCTAGCATAGCAATAC CTGCATTG	
MA0333		AGTCCATTCTCTGTGG	
MA0334	PCR of <i>hisS</i> Pcar_1041	AAGGGATCTATCATGAGCATATCAGGCATTAAGGG	
MA0335		GCCGCGCGCGACTAGTTTCTCGTGTCTTTCC	
MA0052	gentamicin marker	TGCATATGGCTCTAGAATAACTTCGTATAGC	
MA0053		TCGATAAGCTTCTAGAATAACTTCGTATAATG	
Oligonucleotides for construction of chimeric CRISPR expression plasmid pMA35			
Name	Purpose	Sequence (5' to 3')	
MA0269	PCR of <i>lacl-taclacUV5</i> promoter	ACATGTCACCTGCCCGCTTTCCAGTC	
MA0270		GCATGCGTGTGAAATTGTTATCCGC	
MA0429	syntheticCRISPR of spacer #1	AATTCGGTTCATCCCCGCGCATGCGGGGAACACATACAT GAGGGCAAACGCTTTTGGCCGCGCGGTTTCATCCCCG CGCATGCGGGGAACACG	
MA0430		GATCCGTGTTCCCGCATGCGCGGGGATGAACCGCCGCC GGCCAAAAGGCGTTTGCCCTCATGTATGTTCCTCCCGCAT GCGCGGGGATGAACCG	
MA36R	sequencing	CGACATCATAACGGTTC	

Note: Within the sequence of the chimeric CRISPR, a single base pair (underlined) has been duplicated in plasmid pMA35-, at the exact centre of spacer #1.

MasterPure kit for genomic DNA extraction was from EPICENTRE. To construct a transgenic strain of *G. sulfurreducens* in which the native *hisS* gene was replaced with *hisS* from *P. carbinolicus*, three primer pairs (Table 1) were used to amplify the 5'-side and 3'-side flanking regions of *hisS* GSU1659 and the coding sequence of *hisS* Pcar_1041. The two flanking regions were digested with *Spe* I and ligated; this product and the Pcar_1041 amplicon were separately TOPO-cloned (Invitrogen) and sequenced. Digestion with *Bsp*H I (overlapping the start codon) and *Spe* I (overlapping the stop codon) was used to insert the Pcar_1041 gene between the flanking regions of GSU1659. As a selectable marker, a gentamicin resistance cartridge was amplified from plasmid pCM351 [40] with *Xba* I site-containing primers (Table 1), maintained as a TOPO clone, and ligated into the *Nhe* I site between the *Spe* I site and the 3' flanking region of GSU1659. (A similar construct, in which only

the marker was inserted without Pcar_1041, was used in unsuccessful attempts to delete GSU1659.) The entire *hisS* replacement construct was excised using *Eco*R I, purified from an agarose gel, and electroporated into the wild type *G. sulfurreducens* strain DL1 as previously described [39]. An isolated gentamicin-resistant colony was streaked for purity before transfer to liquid. The genotype of this strain, called MA159G, was confirmed by PCR of genomic DNA, with primers MA0334 and MA0335 (which amplify *hisS* Pcar_1041 but not *hisS* GSU1659) as well as MA0330 and MA0333 (which give a larger product for MA159G than for DL1, due to the inserted marker). The marker, which had *loxP* sites on either side, was removed from the chromosome of strain MA159G by introducing the Cre recombinase expression plasmid pCM158 [40] by electroporation and selecting for resistance to kanamycin (Sigma). Four colonies of the resultant strain called MA159 were streaked

for purity and their genotypes were confirmed by PCR; amplicons were digested with *Pst* I to distinguish Pcar_1041 from GSU1659. This strain was electroporated with a plasmid called pRG6 (R. Glaven, personal communication), which is incompatible with pCM158 and selectable with spectinomycin (Sigma); it differs from pRG5 [41] only in that it carries the *lacI* repressor gene. The chromosomal genotype of this strain was confirmed by PCR and *Pst* I digestion.

A plasmid vector called pMA36, incompatible with pRG6, was constructed for isopropylthio- β -D-galactopyranoside (IPTG)-inducible expression of a chimeric CRISPR containing spacer #1 from *P. carbinolicus* between two repeats typical of the CRISPR2 locus of *G. sulfurreducens*. The *lacI* repressor gene and *taclacUV5* promoter of plasmid pCD341 [42] were amplified by PCR with *Pci* I and *Sph* I site-containing primers (Table 1), TOPO-cloned and sequenced, and excised for ligation into plasmid pCM66 [43], resulting in plasmid pMA36. The chimeric CRISPR, consisting of annealed oligonucleotides MA0429 and MA0430 (Table 1), was ligated between the *Bam*H I and *Eco*R I sites of pMA36. The sequence of this plasmid, called pMA35-1, was confirmed using the sequencing primer MA36R (Table 1). Serendipitously, two variants were discovered: pMA35-2 in which the chimeric CRISPR had expanded to two copies of spacer #1 with a third copy of the repeat between them, and pMA35-! in which spacer #1 was disrupted by duplication of a single G:C base pair at its exact centre (underlined in Table 1). All three chimeric CRISPR expression plasmids were electroporated into DL1 and MA159(pRG6), in parallel with pMA36 as a control. The genotypes of multiple kanamycin-resistant colonies of each transformation were confirmed by PCR of Pcar_1041 followed by *Pst* I digestion as well as cloning and sequencing, and by sequencing of plasmids present in the genomic DNA extracts (after transformation into *E. coli* to improve DNA quality). Another variant of pMA35-1 called pMA35-0 was serendipitously discovered in which spacer #1 had been deleted by recombination of the repeats on either side.

Growth conditions

Transformants of *G. sulfurreducens* were selected on NBAF medium [39] containing 1.5% Agar Noble (Difco), supplemented with 5 mM cysteine hydrochloride and 0.1% yeast extract in an anaerobic chamber. Growth experiments were carried out with liquid cultures in either NBAF medium supplemented with 1 mM cysteine hydrochloride or FWAFC medium [39] modified to contain 10 mM acetate and supplemented with 1 mM ferrous ammonium sulfate, in an atmosphere of N₂ and CO₂ (80%:20%) in rubber-stoppered 26 ml glass tubes at 30°C.

Bioinformatics

Codon usage was determined using the CodonFrequency algorithm of the Genetics Computer Group Wisconsin Package version 10.3 (Accelrys Inc., San Diego, CA). A script to compute the number of histidines and the distances between them for every protein sequence in a list was written in Perl.

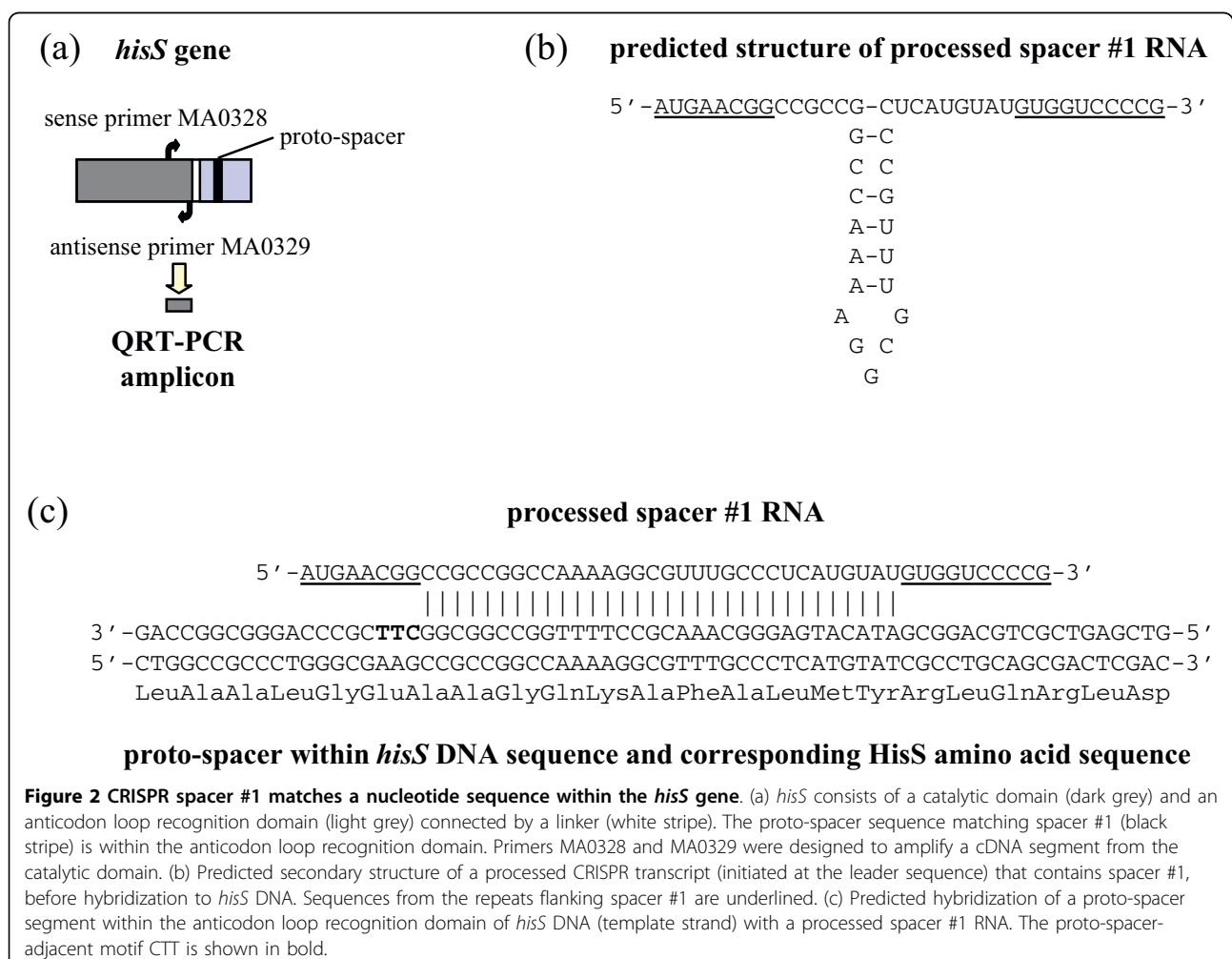
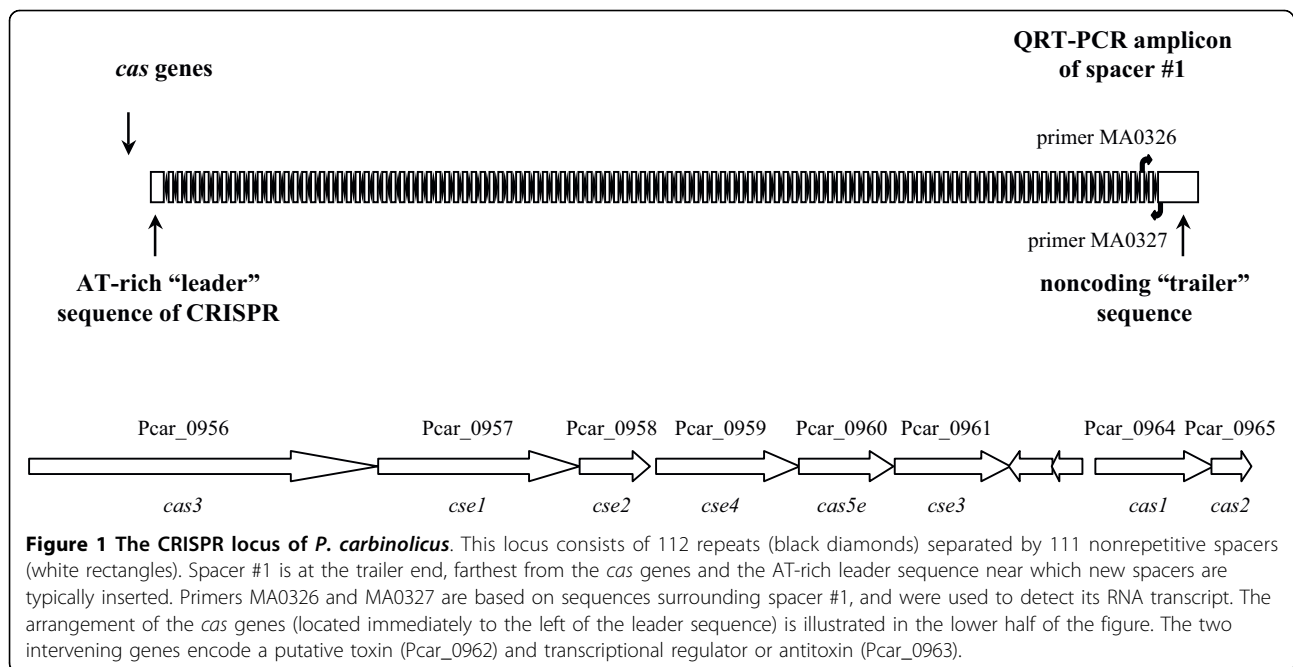
Results

The CRISPR locus of *P. carbinolicus* includes a spacer matching its own histidyl-tRNA synthetase

During manual curation of the *P. carbinolicus* genome annotation, the CRISPR locus was identified as 112 repeats of the sequence 5'-GAGTT**CCCCGCA**-GAT**GC**GGGGATGAACCG-3' (bases in bold predicted to form a hairpin), separated by spacer sequences of 32 bp (Figure 1). This repeat sequence belongs to phylogenetic cluster 2 of the CRISPR classification system [44] and the adjacent *cas* genes (Figure 1) are of the subtype "Ecoli" [3]. The nonredundant nucleotide sequence database was queried in an attempt to identify genes from which the 111 CRISPR spacers of *P. carbinolicus* might be derived. The only hits with five or fewer mismatched bases were hits with zero mismatches within the *P. carbinolicus* genome itself: spacer #1, located at the "trailer" end of the locus, farthest from the AT-rich "leader sequence" and *cas* genes encoding CRISPR-associated proteins (Figure 1), matched a sequence within the histidyl-tRNA synthetase (*hisS*) gene Pcar_1041 (Figure 2); spacer #43 matched the adjacent spacer #44; and spacer #28 matched the nonadjacent spacer #50. Spacer #1 is likely to be the oldest spacer because new spacers are added next to the leader sequence upon exposure of streptococci to bacteriophage [12,13,45-47], and closely related strains of bacteria and archaea contain identical spacers only near the trailer ends of their CRISPR [2,5,11,18,45,48-51]. This observation led to the hypothesis that *P. carbinolicus* has experienced interference with the *hisS* gene, encoding an essential housekeeping enzyme, over a significant period of its evolutionary history.

Quantitative detection of CRISPR spacer #1 transcripts

In an attempt to determine whether spacer #1 is transcribed into RNA that could have interfered with the *hisS* gene at one time, and which strand of trailer end RNA is predominant in *P. carbinolicus*, two oligonucleotide primers were designed flanking spacer #1 (Figure 1, Table 1): MA0326 within spacer #2 and MA0327 just outside the outermost repeat of the CRISPR. Reverse transcription of *P. carbinolicus* RNA into cDNA was attempted with each single primer, followed by quantitative real-time PCR amplification with both primers. The amount of spacer #1-containing RNA including the



sequence found on the sense strand of *hisS*, detected by primer MA0327, was not significantly different from the amount of spacer #1-containing RNA corresponding to the antisense of *hisS*, detected by primer MA0326 (Figure 3). Control PCR reactions without reverse transcription yielded no product, indicating that DNA contamination was negligible and only RNA of both strands was detected.

The sense strand spacer #1-containing RNA detected in this experiment may represent the 3' end of a long transcript initiated near the leader sequence, whereas the antisense strand spacer #1-containing RNA may be produced independently from a promoter at the opposite end of the CRISPR. It is also possible that one strand is produced from the other by an unidentified RNA-directed RNA polymerase. If the sense strand spacer #1-containing RNA undergoes processing similarly to the *E. coli* CRISPR transcript [14], which belongs to the same phylogenetic cluster of repeat sequences as the *P. carbinolicus* CRISPR [44], cleavage within the stem-loops of the repeats flanking spacer #1, followed by 3' end trimming, would release a short RNA with predicted secondary structure (Figure 2b). This or the corresponding antisense strand spacer #1-containing RNA

may hybridize to the proto-spacer DNA sequence within the *hisS* gene (Figure 2c).

Phylogenetic and experimental evidence that interference with *hisS* cannot be compensated

It is surprising that spacer #1 is retained by *P. carbinolicus* if it interferes with the essential function of histidine activation for protein synthesis. Comparative genome analysis revealed that *P. carbinolicus* and its close relatives, the *Geobacteraceae*, possess two full-length *hisS*-like genes, whereas other bacteria have only one. Interference with Pcar_1041 by spacer #1 might have had negligible effect if Pcar_0202 also produced histidyl-tRNA synthetase activity. However, both phylogenetic and mutational studies suggest that Pcar_1041 is essential, being the only real *hisS*, as detailed below.

In some bacteria there is a *hisS*-related gene called *hisZ*, which produces a protein that lacks the C-terminal anticodon loop recognition domain of a true histidyl-tRNA synthetase, functioning instead as a regulatory subunit of ATP phosphoribosyltransferase, the first enzyme of histidine biosynthesis [52]. In bacteria that possess *hisZ*, the *hisG* gene encoding the catalytic domain of ATP phosphoribosyltransferase is shorter than in bacteria that do not possess *hisZ* [53]. A short *hisG* gene is present in *P. carbinolicus* and all other *Geobacteraceae*, but unlike previously described *hisZ* genes, both *hisS*-like genes contain obvious anticodon loop recognition domains. Phylogenetic analysis showed that the Pcar_1041 gene product and orthologous protein sequences of *Geobacteraceae* cluster among the HisS proteins of other bacteria, whereas the Pcar_0202 gene product and its orthologs belong among the HisZ proteins (Figure 4). Furthermore, the ortholog of Pcar_0202 in *G. sulfurreducens* (GSU3307) could be deleted (Aklujkar and Lovley, manuscript in preparation), whereas the ortholog of Pcar_1041 (GSU1659) could only be replaced with Pcar_1041 (this study). Three electroporation attempts failed to delete GSU1659 outright. This result indicates that Pcar_0202 and its orthologs lack significant histidyl-tRNA synthetase activity, and suggests that interference with Pcar_1041 by spacer #1 would exert severe pressure on *P. carbinolicus*.

The phylogenetic tree also demonstrates that the *hisS* gene Pcar_1041 was not acquired laterally; it is clearly an ancestral gene containing a proto-spacer that is not present in its closest relatives.

Spacer #1 inhibits growth of a transgenic *G. sulfurreducens* strain containing *hisS* of *P. carbinolicus*

It is not yet possible to make mutations in *P. carbinolicus*. Therefore, interference of spacer #1 with *hisS* was tested in the more genetically tractable species

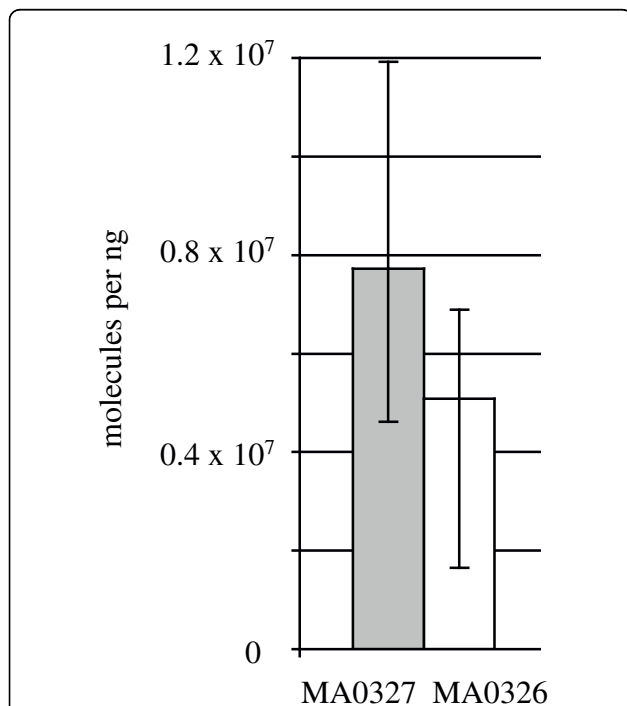
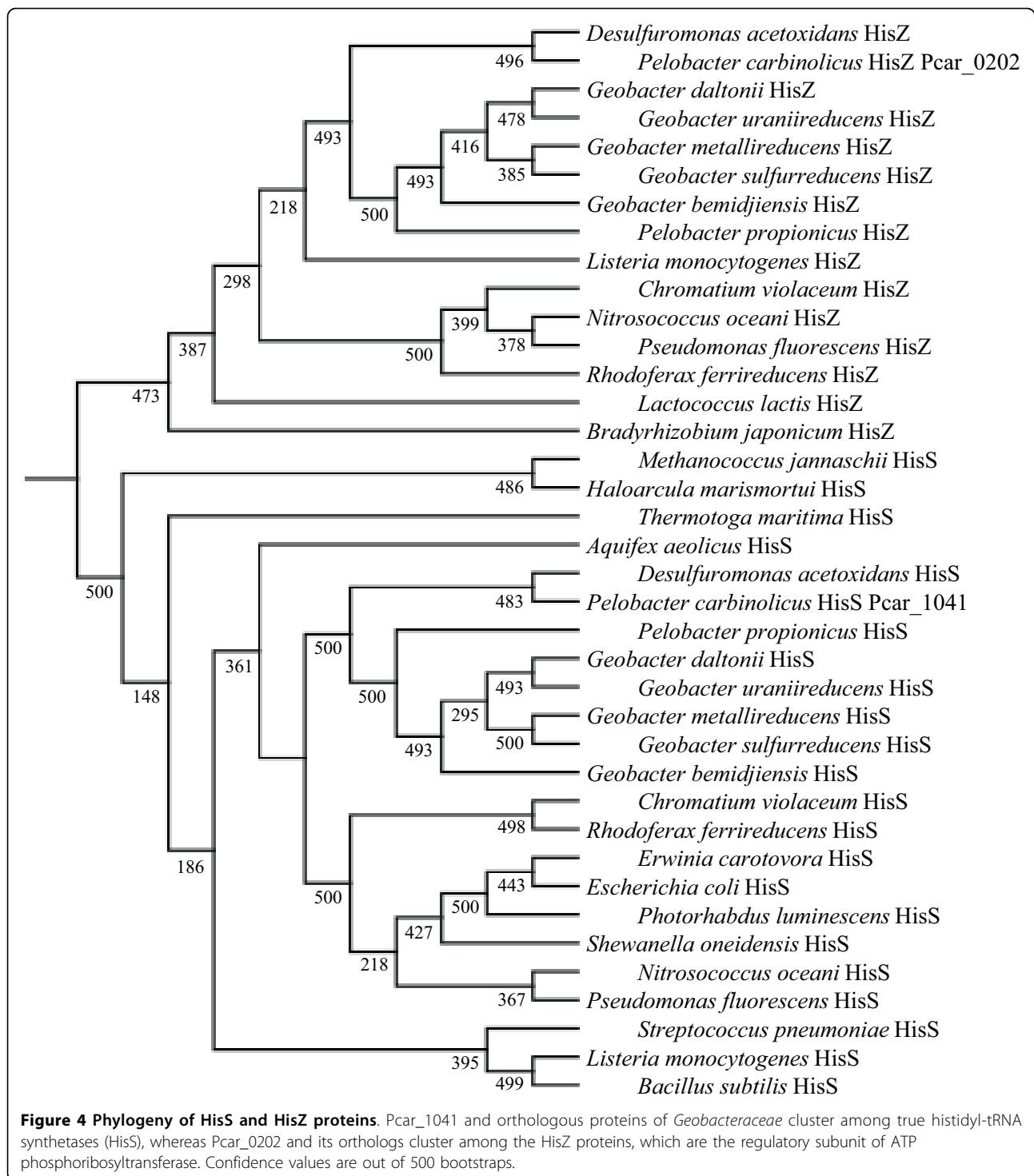


Figure 3 Spacer #1 is transcribed into RNA in *P. carbinolicus*, with both strands similarly abundant. Reverse transcription was performed with either primer MA0327 (grey bar) or primer MA0326 (white bar), and the amount of cDNA was quantified by QRT-PCR. The mean of three biological replicates is shown; error bars represent the minimum and maximum.



G. sulfurreducens, in which the repeat sequence of the CRISPR2 locus (5'-GTGTTCCCCGCATGCGCGGG-GATGAACCG-3') is very similar to that of the *P. carbinolicus* CRISPR. A plasmid called pMA35-1 was designed for IPTG-inducible expression of a chimeric CRISPR construct consisting of spacer #1 of *P.*

carbinolicus between two copies of the *G. sulfurreducens* repeat, and a transgenic strain of *G. sulfurreducens* called MA159 was generated in which the native *hisS* gene GSU1659 was replaced by Pcar_1041, the *hisS* gene of *P. carbinolicus*. The tRNA-His sequences of the two species are very different (Figure 5), suggesting that

mere presence as DNA, is largely incompatible with the *hisS* gene of *P. carbinolicus* containing the matching proto-spacer, which is present in the MA159 host, but not with the native *hisS* gene of *G. sulfurreducens* in the DL1 host.

Growth experiments provided further proof that spacer #1 interferes with *hisS* of *P. carbinolicus*. The *G. sulfurreducens* transformants were first checked by PCR, restriction digestion, and sequencing to verify that the *hisS* transgene and spacer #1 were intact. In one transformant, spontaneous recombination of the repeats on either side of spacer #1, eliminating it from the chimeric CRISPR expression plasmid, resulted in a strain called MA159(pMA35-0) that possessed both *hisS* of *P. carbinolicus* and the repeat sequence on the plasmid, but no spacer #1. Compared to this control that grew similarly to MA159, the presence of spacer #1 in the other MA159 transformants (i.e., with *hisS* of *P. carbinolicus*) resulted in long lag periods and somewhat reduced growth rates in NBAF medium with fumarate as the electron acceptor (Figure 6a), and very poor growth in FWAF medium with Fe(III) citrate as the electron acceptor (Figure 6b). This effect was the same with either one copy of spacer #1 in MA159(pMA35-1) or two copies in MA159(pMA35-2), or with a single base pair insertion in spacer #1 in MA159(pMA35-!) - for the sake of clarity, only MA159(pMA35-2) is shown. The only exception was that in one experiment, triplicate cultures of MA159(pMA35-!) grew especially poorly after three transfers in NBAF (not shown). Wild type growth patterns were observed when any of the three plasmids was present in the DL1 host (i.e., with *hisS* of *G. sulfurreducens*) - for clarity, only DL1(pMA35-2) is shown (Figure 6). Although expression of spacer #1 is expected to be low in the absence of IPTG, growth inhibition of the MA159 strains was observed, and addition of IPTG had no effect, indicating that expression of chimeric CRISPR RNA was not the limiting factor for inhibition of growth.

Spacer #1 reduces the amount of *hisS* RNA in transgenic *G. sulfurreducens* no more than it affects *hisZ* RNA

Total RNA was isolated from NBAF-grown cultures of strains containing *hisS* of *P. carbinolicus*. The amount of *hisS* RNA was higher in the control MA159(pMA35-0) strain than in the growth-inhibited MA159(pMA35-2) strain with spacer #1 (Figure 7), and lowest in the MA159(pMA35-!) strain that had the most severe growth defect in a parallel growth experiment using the same inoculum. However, when the amount of *hisZ* RNA was compared across the same three strains as a control, a similar pattern was observed (Figure 7), suggesting that reduced expression of other housekeeping

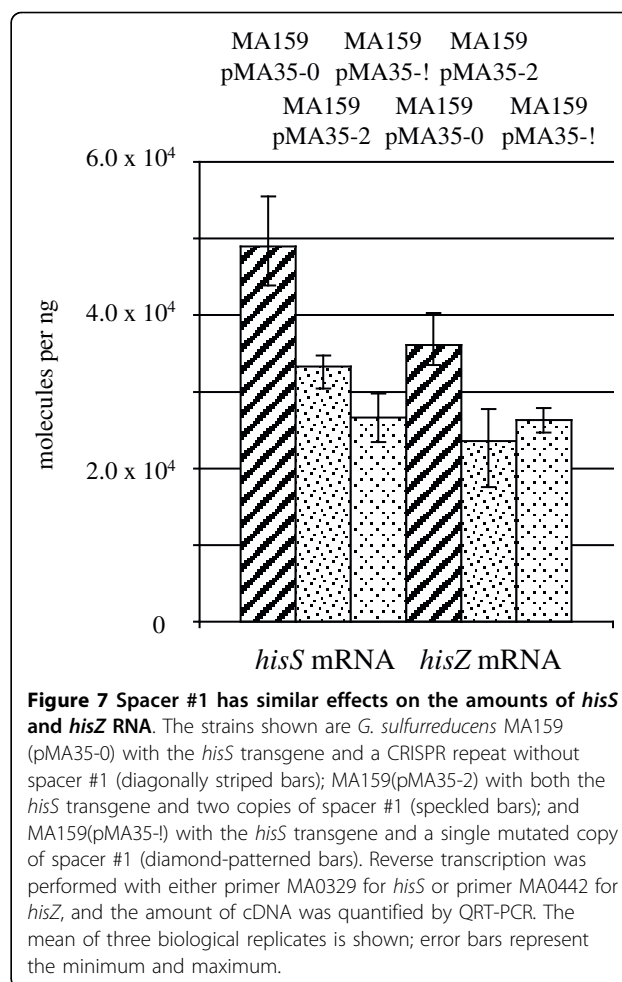


Figure 7 Spacer #1 has similar effects on the amounts of *hisS* and *hisZ* RNA. The strains shown are *G. sulfurreducens* MA159 (pMA35-0) with the *hisS* transgene and a CRISPR repeat without spacer #1 (diagonally striped bars); MA159(pMA35-2) with both the *hisS* transgene and two copies of spacer #1 (speckled bars); and MA159(pMA35-!) with the *hisS* transgene and a single mutated copy of spacer #1 (diamond-patterned bars). Reverse transcription was performed with either primer MA0329 for *hisS* or primer MA0442 for *hisZ*, and the amount of cDNA was quantified by QRT-PCR. The mean of three biological replicates is shown; error bars represent the minimum and maximum.

genes besides *hisS* occurs when growth is slowed by the incompatibility between spacer #1 and *hisS*.

The *P. carbinolicus* genome has fewer genes with numerous or closely spaced histidine codons than closely related genomes

The evidence that *P. carbinolicus* expresses CRISPR spacer #1, and that spacer #1 inhibits growth of a *G. sulfurreducens* strain that is dependent on *hisS* of *P. carbinolicus*, led to the question of whether any effect of this interference during recent evolution could be discerned in the genome of *P. carbinolicus*. If the expected shortage of histidyl-tRNA were occasionally severe enough for ribosomes to stall during translation of genes with numerous histidine codons, one would expect these genes to be predisposed for elimination from the genome, because abortive expression wastes energy and because any selective advantage of the genes would be diminished. Missense mutations of closely spaced histidine codons would also be favoured as long as they did not interfere with an essential function.

Therefore, the number of histidine codons in every gene and the harmonic mean distance between histidine codons in every gene were computed for *P. carbinolicus* and its closest relative with a nearly completely sequenced genome, *Desulfuromonas acetoxidans* [GenBank:NZ_AAEW00000000], as well as for the completely sequenced and manually curated genome annotations of the more distantly related *G. sulfurreducens* [32], *G. metallireducens* [33] and *G. bemidjiensis* (Aklujkar *et al.*, submitted). A plot of the fraction of protein sequences in each genome that have a given minimum number of histidines shows that the *P. carbinolicus* genome is deficient in genes with 35 or more histidine codons, and possesses none with 45 or more (Figure 8a). To identify ancestral genes that might have been counterselected in *P. carbinolicus* due to close spacing of histidine codons, an index of histidine demand was computed as the number of histidine codons in a gene divided by the harmonic mean distance between them. Fewer genes with histidine demand above 5.0 are present in the *P. carbinolicus* genome, and none has an index above 10.0 (Figure 8b). Despite these trends, the overall frequency of histidine codons in the *P. carbinolicus* genome is 22.50 per thousand, very similar to *D. acetoxidans* (23.94 per thousand), *G. sulfurreducens* (20.55 per thousand), *G. metallireducens* (20.42 per thousand) and *G. bemidjiensis* (19.76 per thousand). This observation is consistent with the expected effect

of an acute histidyl-tRNA shortage in the vicinity of gene transcripts with multiple or closely spaced histidine codons undergoing translation, whereas a defect in histidine biosynthesis prior to histidyl-tRNA synthetase would be expected to affect histidine codon usage in general.

The *P. carbinolicus* genome has lost ancestral genes with numerous or closely spaced histidines

Genes of *D. acetoxidans*, *G. sulfurreducens*, *G. metallireducens* and *G. bemidjiensis* that contain 35 or more histidine codons, or have a histidine demand index above 5.0, were examined in order to identify ancestral genes that have reduced their histidine demand or have been lost specifically in the *P. carbinolicus* genome (Additional file 1: Table S1). Many genes found in *Geobacter* species are not necessarily ancestral to *P. carbinolicus*; they lack homologs in either the unfinished *D. acetoxidans* genome or the partial genome sequences of a mixture of *D. acetoxidans* and *D. palmitatis* (D. R. L. and coworkers, unpublished). Other genes that are present in *D. acetoxidans*, but not *Geobacter* species, could have been acquired after divergence from *P. carbinolicus*. Five gene families actually show increased histidine demand in *P. carbinolicus* compared to other *Geobacteraceae*, and in many other cases, a *P. carbinolicus* gene has lower histidine demand than its orthologs, but is still above the cutoff value of 5.0, or contains a similar

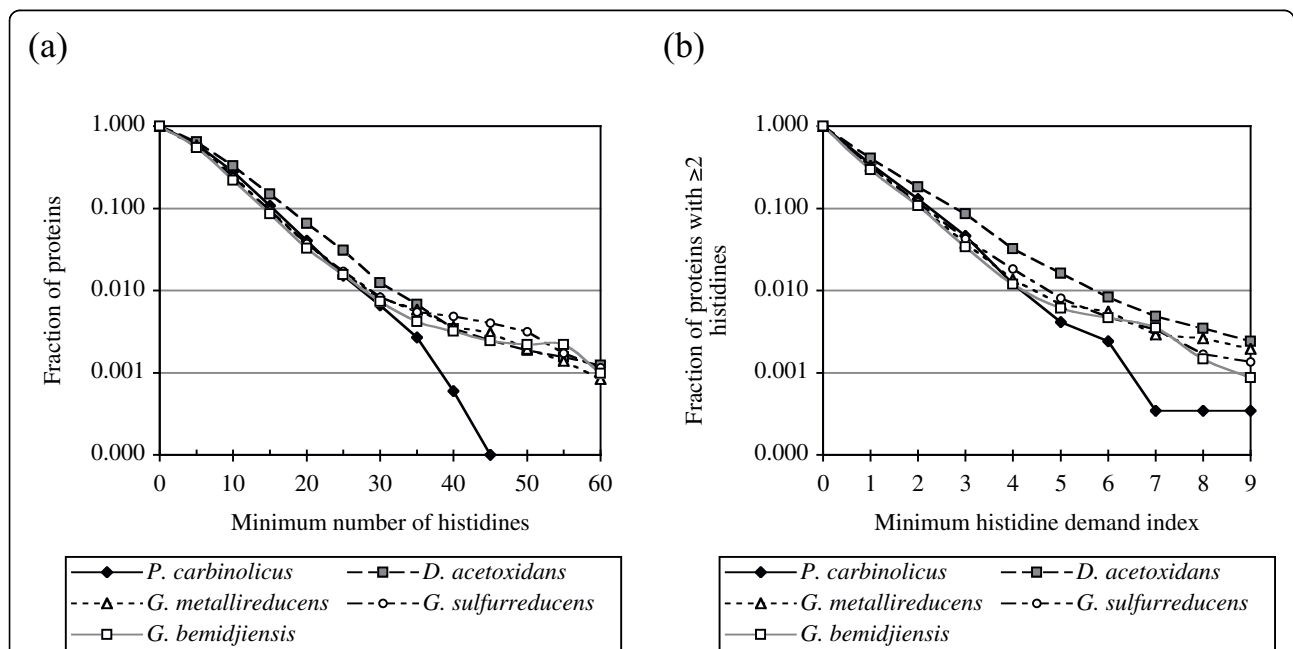


Figure 8 The *P. carbinolicus* genome encodes fewer proteins with multiple closely spaced histidines. (a) The fraction of proteins with a given minimum number of histidines, plotted for the genomes of *P. carbinolicus* (black diamonds), *D. acetoxidans* (grey squares), *G. sulfurreducens* (white circles), *G. metallireducens* (white triangles) and *G. bemidjiensis* (white squares). (b) The fraction of proteins with two or more histidines and a given minimum histidine demand index, plotted for the same five genomes.

number of histidine codons. However, after all these considerations there remain sixteen clearly ancestral gene families with typically high histidine demand that are missing or have reduced histidine demand specifically in *P. carbinolicus* (Additional file 1: Table S1). The functional annotations of these gene families are briefly described below.

The *nuoL-1* gene family, with a high-histidine-demand representative in every *Geobacter* species and *D. acetoxidans*, encodes the L subunit of an NADH dehydrogenase I complex that *P. carbinolicus* has entirely lost. The NuoL protein is thought to have a proton-pumping function [54], for which the imidazole groups of its clustered histidines are well suited.

The *znuA* (or *zntC*) gene encodes the periplasmic protein of an ATP-binding cassette transporter, within which a cluster of histidines is thought to bind zinc with high affinity [55]. The *znuA* gene of *D. acetoxidans* has eight histidines in its putative metal-binding cluster, whereas its closest relative, the *P. carbinolicus znuA* gene (Pcar_3026), has only four.

D. acetoxidans possesses two clearly ancestral genes related to arsenite *S*-adenosylmethyltransferase (Dace_2134, Dace_3081), with homologs in *Geobacter lovleyi* and other *Deltaproteobacteria* that contain numerous histidines. The closest homolog of Dace_3081 is in *P. carbinolicus* (Pcar_2089), with fewer histidines and a much reduced histidine demand index. *P. carbinolicus* lacks an ortholog of Dace_2134.

Two ancestral genes encoding polyketide synthase-type enzymes, with representatives in *D. acetoxidans*, *G. lovleyi*, *Geobacter uraniireducens*, *Geobacter bemidjensis*, *Geobacter* sp. M21, *Geobacter* sp. M18, *Geobacter daltonii*, and *Pelobacter propionicus* that contain numerous histidines, have clearly been lost by *P. carbinolicus*. Both polyketide synthases are composed of unusual domain combinations. Dace_0979 and its orthologs contain just one elongation domain, one acyl carrier protein acylation domain, two to five acyl carrier protein domains, one reductase domain, and no dehydratase or thioesterase domains, suggesting that they build up a long-chain fatty acid with multiple hydroxyl groups, or possibly a storage polymer of a precursor resembling 3-hydroxybutanoate. Dace_1838 and its orthologs contain two elongation domains, one acyl carrier protein acylation domain, three to four dehydratase domains, and no acyl carrier protein or reductase or thioesterase domains, suggesting that their product may be polyhydroxylated, polyunsaturated, or cyclic. The loss of these two enzymes likely means that *P. carbinolicus* does not make secondary metabolites that are present in most other *Geobacteraceae*.

Three other high-histidine-demand gene families that *P. carbinolicus* has lost encode a DUF323 domain-

containing methyltransferase (Dace_1886) with homologs in *G. uraniireducens*, *G. bemidjensis*, *Geobacter* sp. M21, and *Geobacter* sp. M18, a glycoside hydrolase (GSU2359) with homologs in *D. palmitatis*, *G. bemidjensis*, *Geobacter* sp. M21, *Geobacter* sp. M18, *G. daltonii*, and *P. propionicus*, and a predicted *c*-type heme-binding, GAF domain-containing phosphohydrolase (GSU2622) with homologs in *D. palmitatis*, *G. metallireducens*, *G. uraniireducens*, *G. bemidjensis*, *Geobacter* sp. M21, *Geobacter* sp. M18, and *G. daltonii*. The specific reactions catalyzed by these enzymes are not known.

The remaining eight ancestral gene families are multi-heme *c*-type cytochromes represented in *D. acetoxidans* or *D. palmitatis* and one or both of *G. sulfurreducens* and *G. metallireducens*. At least one gene of each family was most probably inherited by *P. carbinolicus* from a common ancestor of the *Geobacteraceae*, and then lost. Several other cytochrome families, found in the *Geobacter* genomes but not the incomplete *D. acetoxidans* and *D. palmitatis* genomes, may have been inherited and lost by *P. carbinolicus*, but the evidence is inconclusive. Nevertheless, it is notable that there are no cytochromes among the eighteen proteins of *P. carbinolicus* that have either more than 35 histidines or a histidine demand index above 5.0 (Additional file 1: Table S1).

Discussion

Although CRISPR spacers that match phage/plasmid genes have been shown to confer immunity against infection [12,15], and it is known that their mode of action is distinct from previously described mechanisms of phage resistance [13], the significant number of spacers that match host genes have not been investigated [25]. The activity of CRISPR spacers against genes encoding essential housekeeping enzymes has not been demonstrated before, nor have its consequences for genome-scale evolution been examined. The present study establishes that when spacer #1 of the *P. carbinolicus* CRISPR and its putative target, the *hisS* gene of the same species, are both present in a transgenic *G. sulfurreducens* strain, interference occurs that severely affects growth. Very few cells containing the *hisS* transgene were able to take up the chimeric CRISPR expression plasmids. Transformants in which spacer #1 was present grew more poorly. Attempts to induce expression of spacer #1 with IPTG had no effect, but growth was almost totally inhibited by switching from fumarate to Fe(III) as the electron acceptor, under which condition expression of Cas proteins and the chromosomal CRISPR2 is upregulated [56]. Therefore, the limiting factor for interference with *hisS* may be the amount of one or more Cas proteins, or the leader sequence, rather than the amount of spacer #1 transcript. Alternatively, growth by respiration of Fe(III) may require protein

factors that cannot be expressed properly when histidyl-tRNA synthetase activity is low. Induction of CRISPR expression with IPTG in *S. epidermidis* has also shown no effect on interference with a target plasmid [15], but that CRISPR was inactive without a leader sequence in *cis*, whereas severe interference between spacer #1 and *hisS* requires neither the native leader sequence of the *G. sulfurreducens* CRISPR nor that of the *P. carbinolicus* CRISPR in *cis*.

The small decrease in the amount of *hisS* RNA in the presence of spacer #1, comparable to the decrease in the amount of transcript for a control housekeeping gene (*hisZ*) that accompanies the growth defect, is unlike the extent of decimation that one would expect if spacer #1-containing RNA catalyzed degradation of *hisS* RNA. It is even less likely that any transgenic strains could be viable if spacer #1-containing RNA targeted *hisS* DNA for degradation. An alternative possibility is that spacer #1 RNA hybridizes with the *hisS* gene without marking it for degradation (but perhaps recruiting one or more proteins), and must be displaced by RNA polymerase in order to complete transcription. In this scenario, even leaky expression of spacer #1 would saturate the available targets, and overexpression of spacer #1-containing RNA would not prevent RNA polymerase from displacing the one molecule obstructing transcription of the *hisS* gene. Consistent with these predictions, induction of spacer #1 expression with IPTG was not required for inhibition of growth, and did not cause stronger inhibition. It will be interesting to examine whether any putative nucleases have mutated in the few viable transformants that carry both spacer #1 and its target. None of the Cmr proteins implicated in RNA-targeted CRISPR function in *P. furiosus* [22] has a homolog in *G. sulfurreducens* or *P. carbinolicus*, nor does the CRISPR-associated double-stranded DNA/RNA-specific endonuclease of *Sulfolobus solfataricus* [57].

Sequencing confirmed that neither *hisS* nor spacer #1 was mutated in the viable transformants. Therefore, in contrast to earlier studies that showed absolute incompatibility between host spacers and phage proto-spacers, evaded only by mutation of the proto-spacers or proto-spacer-adjacent motifs [12,13], these results indicate that it is possible to establish a spacer that persistently interferes with an essential housekeeping gene. Both spacer and target can be maintained intact over numerous generations of inhibited growth. Insertion of a single base pair in the middle of the spacer did not reduce its efficacy, indicating that a small bulge in the region of complementarity between spacer and proto-spacer does not necessarily mitigate interference.

The experiments described herein with the chimeric CRISPR construct also show that a spacer from one species can be active in the context of the somewhat

different repeats and *cas* genes of another species. The *P. carbinolicus* and *G. sulfurreducens* genomes encode homologs of all five components of the Cascade complex (*cse1* CasA Pcar_0957, GSU1385; *cse2* CasB Pcar_0958, GSU1386; *cse4* CasC Pcar_0959, GSU1387; *cas5e* CasD Pcar_0960, GSU1388; and *cse3* CasE Pcar_0961, GSU1389), which processes CRISPR transcripts into target-active RNAs in *E. coli* [14], along with the Cas3 helicase (Pcar_0956, GSU1384) that is required for their activity against targets [14], the Cas2 endoribonuclease (Pcar_0965, GSU1393) that may degrade them [58], and the Cas1 endodeoxyribonuclease (Pcar_0964, GSU1392) that may aid in the acquisition of new spacers [59], and promote DNA/RNA annealing [60]. However, the sequences of most of these Cas proteins are so divergent between the two species (e.g. 28% identity for Cas3) that it is noteworthy that the *G. sulfurreducens* system seemingly required no context other than its cognate repeat to recognize spacer #1 as a guide and *hisS* of *P. carbinolicus* as a target. There is evidence that proto-spacer-adjacent motifs are determinants of target recognition [13,20]. The proto-spacer within *hisS* is followed by a CTT motif (Figure 2), typical of sequences captured by CRISPR of phylogenetic cluster 2 [61], to which the *P. carbinolicus* CRISPR and *G. sulfurreducens* CRISPR2 loci both belong [44]. The facile reconstitution of interference between species with similar repeats and proto-spacer-adjacent motifs despite Cas protein divergence is encouraging for future development of CRISPR-based gene silencing technology with synthetic spacers.

A CRISPR transcript containing spacer #1 and flanking sequences is expressed in *P. carbinolicus*, and potentially processed by the Cas proteins into a *hisS*-interfering RNA. There are at least four possible explanations why the spacer and proto-spacer, which are incompatible in the transgenic *G. sulfurreducens* strain, still co-exist in *P. carbinolicus*. Firstly, being the spacer farthest from the leader sequence in *P. carbinolicus*, spacer #1 may produce comparatively few target-active RNA molecules, as processed RNAs containing spacers distal to the leader sequence are underrepresented in a clone library from *P. furiosus* [19]. Secondly, the repeats on either side of spacer #1 may not be good targets for processing because they deviate from the consensus repeat sequence of the *P. carbinolicus* CRISPR more than it has diverged from that of *D. acetoxidans* since the time of their last common ancestor (Figure 9). Thirdly, pairing of the two strands of RNA derived from spacer #1 might prevent targeting of the proto-spacer. The fourth possibility is that *P. carbinolicus* in its natural environment experiences only growth conditions under which the incompatibility between spacer #1 and *hisS* is permissive, as it is for the transgenic strain

<i>P. carbinolicus</i> CRISPR consensus	GAGTTCCCCGCAGATGCGGGGATGAACCG
repeat on leader side of spacer #1	GTGGTCCCCGCAGGTGCGGGGATGAACGG
repeat on trailer side of spacer #1	GTGGTCCCCGCAGGTGCGGGGATGAACGA
<i>D. acetoxidans</i> CRISPR consensus	ATGTTCCCCGCAGATGCGGGGATGAACCG
<i>G. sulfurreducens</i> CRISPR consensus	GTGTTCCCCGCATGCGCGGGGATGAACCG

Figure 9 Alignment of the repeats on either side of spacer #1 with the CRISPR consensus sequences of *P. carbinolicus*, *D. acetoxidans* and *G. sulfurreducens*.

growing by respiration of fumarate, and has evolved to avoid growth conditions under which the incompatibility is absolute, comparable to respiration of Fe(III) by the transgenic strain.

Both strands of spacer #1 were detected at similar levels in *P. carbinolicus* RNA from actively growing cultures, in contrast with previous observations that both strands are present as RNA in *Sulfolobus acidocaldarius* only during stationary phase [5], that unequal amounts of RNA were detected with probes for the two strands in *P. furiosus* [19] and *Thermus thermophilus* [62], and that only transcripts from one strand were detected in *E. coli* [14], *S. epidermidis* [15], and *Xanthomonas oryzae* [20]. The presence of RNA representing both strands is consistent with CRISPR spacers on one strand of a cluster matching both the sense and antisense of proto-spacers [6], and with artificial spacers in both orientations having activity against their targets, although the efficacy may differ by several orders of magnitude [14].

Together, the evidence of spacer #1 expression and the proof that spacer #1 can inhibit growth in a *hisS*-dependent manner indicate that interference with *hisS* by spacer #1 almost certainly occurred during the recent evolutionary history of *P. carbinolicus*. It is likely that this interference resulted in a growth-limiting shortage of histidyl-tRNA. As expected under this selective pressure, genes with numerous and/or closely spaced histidine codons have evidently been counterselected in the genome of *P. carbinolicus*. Although loss or mutation of ancestral genes is easiest to surmise from comparative genomics, acquisition of low-histidine-demand genes and failure to acquire new high-histidine-demand genes also contribute to this difference.

Among the ancestral gene losses from the *P. carbinolicus* genome attributable to interference with *hisS* by spacer #1, the most obvious are the multiheme *c*-type cytochromes typical of *Geobacter* species. The *P. carbinolicus* genome encodes only 14 *c*-type cytochromes [36], whereas the *D. acetoxidans* genome encodes at least 80 such genes (M. A., unpublished). These cytochromes are not well conserved even between closely

related *Geobacter* species [33,63], and it has been hypothesized that they have a generic function as capacitors [64]. Therefore, in addition to the loss of eight ancestral multiheme *c*-type cytochrome families by *P. carbinolicus*, the non-evolution of new families may also be an effect of interference with *hisS*.

Of the other eight families of ancestral genes that exhibit loss or reduced histidine demand in *P. carbinolicus*, two have especially interesting metabolic and physiological implications. The *nuoL-1* gene encodes a subunit of NADH dehydrogenase I with a conserved histidine cluster, implicated in proton pumping. Abortive expression of *nuoL-1* due to interference with *hisS* by spacer #1 in an ancestor of *P. carbinolicus*, leading to a loss of function for the entire NADH dehydrogenase I complex, would have favoured the elimination of all fourteen structural genes of the complex from the genome, which is what has occurred. The reason why this deletion was not lethal is probably that another ancestral NADH dehydrogenase I complex, for which the *nuoL-2* gene does not contain a cluster of histidine codons, has been retained by *P. carbinolicus*. Conceivably, loss of a major respiratory enzyme complex by *P. carbinolicus* caused it to rely more on laterally acquired fermentative pathways.

Interference with *hisS* by spacer #1 also offers an explanation for the diminutive histidine cluster in the *znuA* gene product of *P. carbinolicus*, which functions to bind zinc in the periplasm for active transport into the cell. Zinc is a cofactor of key metabolic enzymes such as carbonic anhydrase, acetate kinase and phosphotransacetylase [65-67]. If the mutations in the histidine cluster of *znuA* reduce the efficiency of zinc uptake, it could have an impact on metabolism related to the unexplained inability of *P. carbinolicus* to oxidize acetate [28] despite the presence of a complete set of tricarboxylic acid cycle genes in the genome.

Conclusions

This paper reports genetic and comparative genomic evidence that housekeeping genes can be targets of chronic CRISPR interference. Spacer #1 is shown to be

transcribed into RNA with comparable amounts of both strands in *P. carbinolicus*, and to inhibit the growth of a transgenic strain of *G. sulfurreducens* that relies on *hisS* of *P. carbinolicus*, without drastically reducing the level of *hisS* RNA. The genome of *P. carbinolicus* exhibits the expected effect of a histidyl-tRNA shortage. Overall, the ancestral genes lost or mutated by the *P. carbinolicus* genome as part of its tendency towards fewer histidines per gene and lower histidine demand illustrate the de-emphasis of the metal-respiring metabolism that is typical of other *Geobacteraceae*. More generally, while previous studies have approached CRISPR as a microbial immune system, another important role of the system may be to exert pressure on endogenous and essential genes, resulting in dramatic changes in the genome content and physiology of the host species.

Additional material

Additional file 1 : Table S1. Analysis of gene families of *P. carbinolicus*, *D. acetoxidans* and three *Geobacter* species for which one or more members have histidine content or histidine demand above the cutoffs (35 histidines or demand index 5.0).

Abbreviations

cDNA: complementary deoxyribonucleic acid; CRISPR: clustered regularly interspaced short palindromic repeats; DNA: deoxyribonucleic acid; IPTG: isopropylthio- β -D-galactopyranoside; PCR: polymerase chain reaction; QRT-PCR: quantitative real-time polymerase chain reaction; RNA: ribonucleic acid; tRNA: transfer ribonucleic acid.

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Authors' contributions

MA conceived of the study, performed experiments and wrote the manuscript. DL offered guidance. Both authors read and approved the final manuscript.

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