Phage-like *Streptococcus pyogenes* chromosomal islands (SpyCI) and mutator phenotypes: control by growth state and rescue by a SpyCI-encoded promoter

Julie Scott1, Scott V. Nguyen2†, Catherine J. King1, Christina Hendrickson1 and W. Michael McShan1,2 *

1 Department of Pharmaceutical Sciences, The University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA
2 Department of Microbiology and Immunology, The University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA

We recently showed that a prophage-like *Streptococcus pyogenes* chromosomal island (SpyCI) controls DNA mismatch repair and other repair functions in M1 genome strain SF370 by dynamic excision and reintegration into the 5′ end of mutL in response to growth, causing the cell to alternate between a wild type and mutator phenotype. Nine of the 16 completed *S. pyogenes* genomes contain related SpyCI integrated into the identical attachment site in mutL, and in this study we examined a number of these strains to determine whether they also had a mutator phenotype as in SF370. With the exception of M5 genome strain Manfredo, all demonstrated a mutator phenotype as compared to SpyCI-free strain NZ131. The integrase gene (int) in the SpyCIM5 contains a deletion that rendered it inactive, and this deletion predicts that Manfredo would have a pronounced mutator phenotype. Remarkably, this was found not to be the case, but rather a cryptic promoter within the *int* ORF was identified that ensured constitutive expression of *mutL*, and the downstream genes encoded on the same mRNA, providing a striking example of rescue of gene function following decay of a mobile genetic element. The frequent occurrence of SpyCI in the group A streptococci may facilitate bacterial survival by conferring an inducible mutator phenotype that promotes adaptation in the face of environmental challenges or host immunity.

Keywords: prophages, chromosomal islands, SpyCI, DNA mismatch repair, mutator phenotype, group A streptococcus, *Streptococcus pyogenes*

INTRODUCTION

Bacteria and bacteriophages exist in a dynamic relationship. In *Streptococcus pyogenes* (group A streptococcus), prophages are prominent components of bacterial chromosomes (Banks et al., 2002; Canchaya et al., 2002). The integration of phage DNA into a bacterial chromosome imparts an increased metabolic burden on the host and serves as a constant threat to survival should conditions change and the prophage enter the lytic cycle. However, the prevalence of lysogeny in *S. pyogenes* implies that prophage acquisition benefits the host under some conditions. Phage-encoded fitness factors such as toxin genes act as means of bringing positive selection pressure upon the host bacterium to maintain the phage DNA in its chromosome, and while such fitness factors may not be required for the phage life cycle, they may act to promote host bacterial growth or survival (Brussow et al., 2004).

We recently reported a novel prophage-like chromosomal island (CI) in M1 genome strain SF370 that integrates and excises from the chromosome in response to cell growth stage (Scott et al., 2008). This element, originally annotated as prophage SF370.4, is now identified as *S. pyogenes* Chromosomal Island M1 (SpyCIM1) to conform to the nomenclature proposed by Novick et al. (2010). SpyCIM1 integrates between the DNA mismatch repair (MMR) genes *mutS* and *mutL*, leading to the silencing of *mutL* and several downstream genes, resulting in the inactivation of MMR. However, during exponential growth the SpyCI excises and replicates as an episome, allowing *mutL* to be transcribed, and restoring MMR to correct errors following DNA replication (Figure 1). As cell density increases, SpyCIM1 re-integrates into the chromosome, interrupting transcription of the polycistronic message and creating a mutator phenotype (Scott et al., 2008). Three genes directly downstream of *mutL* are predicted to be included on this polycistronic message: the multidrug resistance transporter *lmrP*, the Holliday junction DNA helicase *ruvA*, and the base excision repair glycosylase *tag*. The SpyCI are members of a larger group of bacteriophage-related CI found in the low %G+C Gram-positive organisms. The best-characterized members of this group are the prophage-like *Staphylococcus aureus* pathogenicity islands (SaPI), which are the vectors for the toxic shock syndrome toxin and other virulence factors (Novick et al., 2010). The SaPI disseminate to new host staphylococcal cells by hijacking and remodeling the capsids of helper bacteriophages (Tormo et al., 2008; Ubeda et al., 2009), and a similar strategy appears to be used by the SpyCI to infect new host streptococci (Nguyen and McShan, unpublished results). Thus, these CI may be considered a unique subset of prokaryotic viruses.

Nine of the 16 published *S. pyogenes* genomes have a SpyCI integrated between *mutS* and *mutL*. Although integration in this...
location appears to be a common occurrence in *S. pyogenes*, limited sequence identity has been observed between these modified prophages with the exception of highly conserved modules for site-specific integration, DNA replication, and maintenance (Scott et al., 2008). Here we report that *S. pyogenes* strains with a SpyCI integrated between the MMR genes *mutS* and *mutL* exhibit an increased spontaneous mutation rate and an increased sensitivity to UV irradiation due to the inhibition of RuvA as compared to strains lacking these elements. The unexpected exception was the M5 genome strain Manfredo that is always wild type for mutation rate and UV sensitivity, despite having a CI permanently integrated between *mutS* and *mutL* due to a 128 bp deletion in the integrase ORF. We found that this paradox of harboring a SpyCI without a mutator phenotype is a result of the transcription of *mutL* and the downstream genes from a cryptic promoter within the inactivated SpyCI integrase pseudogene, providing a novel example of host gene expression rescue following CI sequence decay.

**MATERIALS AND METHODS**

**BACTERIAL STRAINS AND CULTURE CONDITIONS**

Bacterial strains used in this work are listed in Table 1. *S. pyogenes* NZ131 is an M49 strain isolated from a patient in New Zealand with acute post-streptococcal glomerulonephritis; its genome has been completely sequenced and does not have a SpyCI integrated between the MMR genes, *mutS* and *mutL* (McShan et al., 2008). Thus, it is wild type for MMR and has been used previously as a control (Scott et al., 2008). The following strains, whose genomes have been completely sequenced, were available for these studies, and all contain a SpyCI integrated into the identical attachment site in *mutL*: SF370 (M1), MGAS10270 (M2), MGAS10750 (M4), Manfredo (M5), MGAS10394 (M6), and MGAS6180 (M28; Table 1). *S. pyogenes* strains 270fer, 3470-04, and 5474-00 are serotype M49 clinical isolates whose genomes have not been sequenced but have a prophage-like SpyCI integrated at *mutL* (McShan et al., 2008). These SpyCI containing clinical strains were included to provide serotype-matched controls for NZ131. *S. pyogenes* strains were grown in Todd–Hewitt broth (BD, Franklin Lakes, NJ, USA) supplemented with 2% yeast extract (THY medium) at 37°C. For some strains, brain heart infusion (BHI) supplemented with 5% heated horse serum was used for growth media. *Escherichia coli* strains were grown in L broth supplemented with the appropriate antibiotics as needed (Sambrook and Russell, 2001). Growth was monitored by absorbance at 600 nm ($A_{600}$) using a Genesys20 spectrophotometer (Thermo Scientific, Waltham, MA, USA), and cells were harvested at appropriate growth stages as needed for subsequent experimental analysis. For induction of SpyCI, cultures were grown until the $A_{600} = 0.2$ and then split into two 40 ml portions. Samples were induced by the addition of mitomycin C from *Streptomyces caespitosus* (Calbiotech, Spring Valley, CA, USA) to a final concentration of 0.2 µg/ml. Incubation at 37°C was continued, and samples were collected for each strain at approximately $A_{600} = 0.2$, 0.4, 0.6, and 0.8 and stored in RNA later (Ambion, Austin, TX, USA) at 4°C. The remainder of each culture was left at 37°C overnight, and final samples were collected for each strain the following day.

**Table 1 | Streptococcus pyogenes** strains used in this work.

| Strain   | Relevant genotype | Associated disease                         | Accession number (ref.) |
|----------|-------------------|-------------------------------------------|-------------------------|
| NZ131    | *emm*49, MMR wild type | Acute post-streptococcal glomerulonephritis | CP000829 McShan et al. (2008) |
| SF370    | *emm*1, SpyCIM1   | Wound infection                           | AE004092 Ferretti et al. (2001), Scott et al. (2008) |
| MGAS10270| *emm*2, SpyCIM2   | Pharyngitis                               | CP000260 Beres et al. (2006) |
| MGAS10750| *emm*4, SpyCIM4   | Pharyngitis                               | CP000262 Beres et al. (2006) |
| Manfredo | *emm*5, SpyCIM5   | Rheumatic heart disease                   | AM295007 Holden et al. (2007) |
| MGAS10394| *emm*6, SpyCIM6   | Pharyngitis                               | CP000003 Banks et al. (2004) |
| MGAS6180 | *emm*28, SpyCIM28 | Invasive disease                          | CP000056 Green et al. (2005) |
| 3470-04  | *emm*49, SpyCIM49.1| Invasive disease                          | McShan et al. (2008)  |
| 5474-00  | *emm*49, SpyCIM49.2| Invasive disease                          | McShan et al. (2008)  |
| 270fer   | *emm*49, SpyCIM49.3| Invasive disease                          | McShan et al. (2008)  |
| OKM48    | Manfredo (pWM483) | This work                                 |                         |
| OKM49    | Manfredo (pWM483) | This work                                 |                         |
| OKM50    | NZ131 (pWM483)    | This work                                 |                         |
| OKM51    | NZ131 (pWM483)    | This work                                 |                         |
| OKM52    | NZ131 (pDL276)    | This work                                 |                         |
| OKM54    | NZ131 (pWM488)    | This work                                 |                         |
| OKM55    | NZ131 (pWM489)    | This work                                 |                         |
| OKM56    | NZ131 (pWM490)    | This work                                 |                         |
| OKM57    | Manfredo (pWM488) | This work                                 |                         |
| OKM58    | Manfredo (pWM488) | This work                                 |                         |
| OKM59    | Manfredo (pWM489) | This work                                 |                         |
| OKM60    | Manfredo (pWM490) | This work                                 |                         |
| OKM61    | Manfredo (pWM489) | This work                                 |                         |
| OKM62    | Manfredo (pWM489) | This work                                 |                         |
NUCLEIC ACID ISOLATION AND cDNA SYNTHESIS
Isolation of streptococcal DNA was done as previously reported (Pitcher et al., 1989; McShan et al., 1998). DNA concentrations were determined by spectrophotometry at 260 nm using a NanoDrop ND-1000 spectrophotometer (NanoDrop Products, Wilmington, DE, USA). RNA was prepared using the RiboPure system for bacteria (Ambion) following the manufacturer’s recommended protocol. RNA samples were tested for DNA contamination by PCR using primers specific for the variable region of the 16srna gene (Beall et al., 1996). RNA samples were converted to cDNA using Superscript II (Invitrogen, Carlsbad, CA, USA) and random hexamer priming by following the manufacturer’s protocol.

PCR AND QUANTITATIVE REAL-TIME PCR (qRT-PCR)
PCR was used to detect transcription of mutS, mutL, and the intergenic region between int and mutL. Ten nanograms of cDNA was used for each reaction. PCR was also performed on RNA and DNase-treated RNA as controls. PCR was carried out in a Techne TC-312 Thermocycler (TechneInc, Burlington, NJ, USA). For each reaction the following program was used: an initial denaturation at 95°C for 5 min; 35 cycles each consisting of denaturation at 95°C for 30 s, annealing at 55°C for 30 s, and elongation at 72°C for 60 s; and a final elongation period at 72°C for 5 min.

qRT-PCR was used to evaluate excision of the SpyCI from the host chromosome and to evaluate transcription of MMR genes, mutS and mutL. Primers (Table 2) were designed using the Primer3 program1 and synthesized by Integrated DNA Technologies (Coralville, IA, USA). For each real-time PCR plate evaluated, primers to a 126 base pair (bp) region of the 16S ribosomal subunit housekeeping gene were also run on each sample for normalization of the data. All data was analyzed with respect to the uninduced, housekeeping gene were also run on each sample for normalization of the data. All data was analyzed with respect to the uninduced, housekeeping gene for each strain being equal to one. For each reaction, 10 ng of cDNA was used for qRT-PCR analysis of mutS, mutL, and 16S ribosomal subunit expression. Water blanks were run as negative controls, and all samples and controls were run in triplicate. qRT-PCR was carried out on a BioRad iCycler equipped with the real-time optical fluorescent detection system using SYBR Green PCR Master Mix (BioRad Laboratories, Hercules, CA, USA). The following program was employed for all qRT-PCR: an initial denaturation (95°C, 3 min); followed by 35 cycles of denaturation at 95°C, 30 s, annealing at 55°C, 30 s, and elongation at 72°C, 30 s. Following the last cycle, a melt-curve analysis was performed to verify that a single product was produced for each primer pair. The resulting data was analyzed using Excel and plotted using Prism4.

DETERMINATION OF SPONTANEOUS MUTATION RATES
A fluctuation test based on the Luria and Delbrück assay (Luria and Delbrück, 1948; Rosche and Foster, 2000) was used to determine the spontaneous mutation rates of 10 S. pyogenes strains as previously described (Scott et al., 2008). Strain NZ131 was used as a SpyCI-free control for comparison to the SpyCI containing strains. A single colony of each strain was used to inoculate a THY broth culture and grown overnight at 37°C. The overnight culture then was diluted in THY to less than 1000 CFU/ml and dispensed as 41 ml aliquots into culture tubes. Cultures were incubated at 37°C overnight. One tube was used to determine total CFU/ml by diluting in phosphate buffered saline (PBS) and plating dilutions of 10⁻⁴ and 10⁻³ CFU/ml on THY agar plates in triplicate. The 40 remaining cultures were each mixed with 3 ml of melted soft agar (0.6% agar in water at 45°C) and poured onto THY plates containing 10× the previously determined minimum inhibitory concentration of the gyrase inhibitor ciprofloxacin (2 µg/ml; Scott et al., 2008). For some experiments, 500 µg/ml streptomycin was used for selection of spontaneous antibiotic resistant mutants. Plates were incubated at 37°C for 6–17 days to allow for growth of ciprofloxacin or streptomycin resistant colonies. Plates were examined for colonies each day and were counted on the day when colonies first appeared. Since strain Manfredo never produced antibiotic resistant colonies at this population density, the protocol was modified to use 5 ml replicate cultures for overnight growth that were concentrated by centrifugation and resuspended in 1 ml of THY before plating onto selective medium or diluting for CFU determination. The mutation rate with confidence limits was calculated for each strain using the algorithm of Ma et al. (1992), combined with the technique of maximum likelihood estimation (Stewart, 1994), as implemented by the software package ft (Shaver and Sniegowski, 2003). All experiments were performed at least three times for each strain and the data collected was averaged to determine mutation rate in mutations/generation (µ).

UV IRRADIATION KILLING ASSAY
The sensitivity of S. pyogenes strains to killing by UV irradiation from a calibrated 254 nm germicidal lamp (120 µW/cm²) was done as previously described (Scott et al., 2008). UV irradiation of cultures was carried out in a darkened room to avoid photoreactivation. Assays were conducted a minimum of three times per strain to confirm results.

FLUORESCENT PRIMER EXTENSION
Overnight cultures of S. pyogenes strains started from a single colony were grown in 50 ml THY broth at 37°C for 16 h. Cultures were then diluted 1:20 in 11 THY broth and incubated at 37°C, and growth was monitored by absorbance. At A600 = 0.6, samples were collected and quick-frozen using a dry ice-ethanol bath. RNA isolation was performed and treated with DNase as described above. RNA integrity was evaluated at the University of Oklahoma Health Sciences Center Laboratory for Genomics and Bioinformatics using an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). Fluorescent primer extension was carried out in RNase-free 0.5 ml tubes using 0.2 µM 5’-6-FAM labeled primers. Primers were made to map the 5’ transcriptional start sites of mutS and mutL; accordingly, primers were designed to anneal so that the 3’ end of the primer would extend toward the 5’ end of the gene. The primers for mutS, mutL, and SpyCIM5 (ϕMan5,5 int) were designed to anneal to a region of RNA between 180 and 200bp from the predicted start codon for each gene (Table 2). Primer extension was performed by combining 1 µl of diluted FAM labeled primer with 20 µg RNA and DEPC-treated water to a total volume of 20 µl. Samples were heated at 70°C for 5 min, cooled on ice for 10 min, incubated at 58°C for 20 min, and left at room temperature for 15 min. The following reagents were added

\(^{1}\)http://frodo.wi.mit.edu
**Table 2 | Plasmids and PCR primers.**

| Plasmid | Description | Ref. |
|---------|-------------|------|
| pMV158GFP | Mobilizable, Gram-positive vector encoding gfp | Nieto and Espinosa (2003) |
| pSMART-LCKan | Low copy, transcription-free cloning vector | Lucigen Corp., Middleton, WI, USA |
| pGEM-T Easy | PCR cloning vector | Promega Corp., Madison, WI, USA |
| pDL276 | Shuttle vector for low %G+C Gram-positive bacteria | Dunny et al. (1991) |
| pCAMP17 | Phage T12 derived integration vector under the control of the CAMP promoter | Gase et al. (1999) |
| pWM448 | SpyCIM1 int clone into the CAMP promoter in pCAMP17 | This work |
| pWM481 | SpyCIM5 int pseudogene and upstream (primers M5int5 and M5int4) fused to gfp in pSMART-LCKan | This work |
| pWM483 | Insert from pWM481 cloned into pDL276 | This work |
| pWM485 | SpyCIM5int pseudogene (primers M5int1 and M5int4) fused to gfp in pSMART-LCKan | This work |
| pWM486 | SpyCIM5 int pseudogene subregion (primers M5int2 and M5int4) fused to gfp in pSMART-LCKan | This work |
| pWM487 | SpyCIM5 int pseudogene subregion (primers M5int3 and M5int4) fused to gfp in pSMART-LCKan | This work |
| pWM488 | Insert from pWM486 cloned into pDL276 | This work |
| pWM489 | Insert from pWM485 cloned into pDL276 | This work |
| pWM490 | Insert from pWM487 cloned into pDL276 | This work |

**Sequence Primer | Gene target | Product size (bp)**

**PRIMERS FOR PCR**

| Primer | Gene target | Product size (bp) |
|--------|-------------|-------------------|
| 5’AGCAGGATTCACCAGCCTTA | attB-L | 655 (strain SF370) |
| 5’CTCAATTTTATCAACAGTACG | attB-R | 694 (strain Manfredo) |
| 5’CTTGGCTGGTAAACTGCTTTG | attP-L | 461 |
| 5’CAAGCGCTTGAAGTGTTGGTGC | attP-R | 406 |
| 5’AGCTTTTATGGTTAACTGTTCG | mutS-L | 413 |
| 5’AGCTTTTATGGTTAACTGTTCG | mutS-R | 413 |
| 5’ATGTAAGAGGGAGTACAGTTT | gfp-L | 717 |
| 5’TATTGGATATGTTATCATCATT | gfp-R | 717 |
| 5’ACAGACGACAGGAAGAAC | M5int1 | Various lengths |
| 5’ATAAGATAACCCTAGTTGATA | M5int2 | |
| 5’TATTGGTACTTGTTGAA | M5int3 | |
| 5’ATTTGGTTTTAACCATTCTCAAC | M5int4 | |
| 5’TCTACTCCCCATGGGAACAA | M5int5 | |

**PRIMERS FOR qRT-PCR**

| Primer | Gene target | Product size (bp) |
|--------|-------------|-------------------|
| 5’ACTGCCAATCATCAACGTCAAT | SpyCIM1 attP | 96 |
| 5’TGGAGGTGTTAAACCTAATGAAACC | SpyCIM5attP | 112 |
| 5’CAATACCCATGAAAACACTGCAAAT | attB | 65 |
| 5’CCCGAACTCAATGTATTTTTTCAT | mutS | 108 |
| 5’CCCGAACTCAATGTATTTTTTCAT | mutL | 111 |
| 5’CACTCCTCCCTTGTGCATT | 16S rRNA | 126 |
to each sample after primers were allowed to anneal: 4 μl dNTPs (10 μM each), 4 μl DTT (0.1 M), 8 μl 5 × first strand buffer, 4 μl Superscript II Reverse Transcriptase (all reagents from Invitrogen).

cDNA was synthesized by incubating samples for 2 h at 42°C and then precipitated by the addition of 1 μl glycogen (Ambion), 4 μl 3 M sodium acetate, and 100 μl of 100% ethanol. Samples were stored overnight at −20°C and harvested by centrifugation at 4°C at 13,000 × g for 20 min. Pellets were washed with 70% ethanol and collected by centrifugation as before for 5 min. Pellets were dried to completion on medium heat in a Savant DNA120 concentrator (Thermo Scientific) and submitted to The University of Oklahoma Health Sciences Center Laboratory for Genomics and Bioinformatics for DNA fragmentation analysis on an ABI 3730 capillary sequencer (Applied Biosystems, Foster City, CA, USA). Data was analyzed using Peak Scanner Software (Applied Biosystems).

**CLONING AND EXPRESSION OF THE SpyCIM1 INTEGRASE**

The integrase ORF from SpyCIM1 from strain SF370 was amplified by PCR and cloned into the commercial vector pGEM-T Easy (Promega, Madison, WI, USA) to create plasmid pWM447 (Table 2). This PCR reaction introduces a unique BamHI site into the 5’ end of the ORF, allowing gene fusion to the S. pyogenes CAMP promoter (Gase et al., 1999). Following digestion of pWM447 with BamHI and PstI to recover the cloned integrase gene, the fragment was ligated to pCAM17 (Gase et al., 1999) to create plasmid pWM448. This plasmid contains the phage T12 integrase and attachment site part of the plasmid and will mediate site-specific integration at an unrelated site, the tmRNA gene (McShan et al., 1997). The plasmid pWM448 was introduced in strain Manfredo by electrotansformation (McLaughlin and Ferretti, 1995). For detection of integrase-mediated excision, DNA or RNA was isolated from strains Manfredo, SF370, or Manfredo (pWM448) during early logarithmic growth. PCR specific for attB was used to detect excision of the SpyCIM5 in strain Manfredo following introduction of pWM448.

**CONSTRUCTION OF SpyCIMS INTEGRASE-REPORTER FUSIONS**

Various lengths of the Manfredo SpyCIM5 int pseudogene and upstream region were amplified using PCR and fused to the green fluorescent protein (GFP) to create reporter fusions (primers and plasmids listed in Table 2). The longest PCR product (1458 bp, primers M5int5 and M5int4) included the entire upstream region, the pseudogene, and the downstream region to the beginning of the mutL ORF, and was cloned to create pWM483. Increasingly shorter products were amplified covering only the pseudogene or portions of the pseudogene and the downstream region as shown in Figure 7 (using primers M5int1 + M5int4, M5int2 + M5int4, and M5int3 + M5int4 to create the inserts for plasmids pWM489, pWM488, and pWM490, respectively). The GFP gene (gfp) was amplified by PCR from plasmid plasmid pMV158GFP (Nieto and Espinosa, 2003). PCR products were synthesized with phosphorylated primers and ligated using standard protocols (Sambrook and Russell, 2001). Following ligation, the desired fusion product was amplified using the downstream gfp primer and the desired SpyCIM5 int pseudogene primer. The amplified fragments were cloned into the pSMART-LCKan vector (Lucigen, Middleton, WI, USA). The cloned insert was removed by digestion with EcoRI and ligated into the EcoRI site of shuttle vector pDL276 (Dunny et al., 1991). This plasmid was introduced by electrotansformation into S. pyogenes strains Manfredo or NZ2131. For fluorescent microscopy, streptococcal strains were grown overnight in BHI supplemented with 5% heated horse serum and 200 μg/ml kanamycin. The cultures were harvested by centrifugation and washed once with PBS. Cells were resuspended in 0.1 volume of PBS, and GFP expression was visualized on a Leica model DM4000B fluorescent microscope using a SPOT digital camera and software package (v. 4.7; Diagnostic Instruments, Inc., Sterling Heights, MI, USA). Mitomycin C treatment, an overnight culture grown in BHI supplemented with 5% heated horse serum and 200 μg/ml kanamycin was diluted 1.20 into fresh media and grown at 37°C until A600 = ~0.1. The culture was divided into two aliquots, and mitomycin C (0.2 μg/ml) was added to one aliquot. Incubation was continued for 1 h, the cells were washed and resuspended in PBS as above, and microscopy was performed.

**BIOINFORMATICS**

The genome sequences for the SpyCI (annotated as prophages) integrated between mutS and mutL were extracted from the Genbank annotated bacterial genomes of SF370, MGAS10270, MGAS10750, MGAS6180, MGAS10394, and Manfredo (Table 1). The sequences were aligned and a phylogenetic tree was created using Geneious (Drummond et al., 2012), and the tree was further analyzed by neighbor-network analysis by SplitsTree using the Equal Angle algorithm (Huson, 1998). Promoter predictions were done using neural network promoter prediction software (Reese, 2001) available at the Berkeley Drosophila Genome Project website2. Prediction of rho-independent transcriptional terminators in the SpyCI genomes was done using TransTermHP (Kingsford et al., 2007).

**RESULTS**

SpyCI and Mutator Phenotypes in S. pyogenes

In S. pyogenes and other Gram-positive bacteria, MMR requires the participation of two proteins, MutS and MutL, to ensure the fidelity of the genome replication. The loss of MMR function results in a mutator phenotype and can promote homologous or non-homologous recombination in both prophages and eukaryotes (Glickman and Radman, 1980; Ressaygier et al., 1989; Selva et al., 1997). In group A streptococci, a single polycistronic mRNA is transcribed that includes mutS and mutL as well as the downstream genes napP, napA, and tag; the promoter for this operon is immediately upstream of mutS (Figure 1). Frequently, a prophage-like chromosomal island (SpyCI) is integrated between mutS and mutL in S. pyogenes genomes, separating mutS and the other downstream genes from the promoter upstream of mutS. The integration of SpyCI leads to inactivation of the downstream genes, including mutL, rendering the bacteria defective for MMR. We recently have shown that in strain SF370, SpyCI integration is dependent on the growth phase of the bacterium (Scott et al., 2008). When cells are rapidly dividing during logarithmic growth, the SpyCI excises from the bacterial chromosome and allows both

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2http://www.fruitfly.org/seq_tools/promoter.html
mutS and mutL to be transcribed; however, in stationary phase the SpyCI re-integrates and silences the genes downstream of mutS (Figure 1). Through this novel system of gene regulation, the SpyCI regulates the streptococcal mutation rate in response to nutrient availability or other environmental signals.

Nine of the 16 sequenced S. pyogenes genomes contain a related prophage-like SpyCI between mutS and mutL, whose presence could result in a complex mutator phenotype. Six of the genome strains were available for this study, and using a modified Luria and Delbrück fluctuation assay (Luria and Delbrück, 1948; Rosche and Foster, 2000) the rate of spontaneous mutation was determined for these S. pyogenes genome strains (SF370, Manfredo, MGAS6180, MGAS10270, MGAS10394, and MGAS10750). Genome strain NZ131, lacking a SpyCI, was used as a wild type control as was done previously (Scott et al., 2008). Additionally, clinical M49 strains containing a SpyCI integrated into mutL (270fer, 3470-04, and 5474-00) were used as serotype-matched strains to NZ131.

The mutation rate was estimated to be $2.19 \times 10^{-8}$ mutations/generation ($\mu$) for the non-SpyCI containing strain NZ131 (Figure 2 and Table 3), a value in agreement with our previous report (Scott et al., 2008). Consistent with the hypothesis that a SpyCI located between mutS and mutL prevents competent MMR, thereby increasing an organism’s mutation rate, the mutation rates for seven of the SpyCI containing strains were determined to be at least one order of magnitude higher than that of NZ131 (MGAS10750, 5474-00, 3470-04, 270fer, SF370, MGAS6180, and MGAS10270). Strain MGAS10394 was found to have a mutation rate of $3.66 \times 10^{-8} \mu$, over 160 times higher than NZ131 (Table 3). The range of observed mutation rates may reflect the variations in the controlling repressor-operator modules in the SpyCI (discussed below) as well as other differences in genetic backgrounds in these strains. Further, the SpyCIM49-containing clinical M49 strains (5474-00, 3470-04, and 270fer) showed an increased mutation rate as compared to NZ131, showing that the phenotype was serotype-independent (Table 3). These studies are highly suggestive, but confirmation of the role of SpyCI in increasing the mutation rate of its host will require the analysis of isogenic strains (manuscript in preparation). The one exception to these findings was that the SpyCIM5 containing strain Manfredo had the lowest mutation rate ($1.0 \times 10^{-11} \mu$), suggesting that the presence of the SpyCI did not interfere with MMR in this strain. The mutation rates clustered into three different groups (Table 3): non-mutator with a mutation rate $<10^{-8} \mu$ (Manfredo and NZ131); mutator with a mutation rate of $\sim 10^{-7} \mu$ (MGAS10750, 5474-00, 3470-04, 270fer, SF370, MGAS6180, and MGAS10270); hypermutator with a mutation rate of $10^{-6} \mu$ (MGAS10394).

We previously reported that strain SF370 had increased susceptibility to damage from UV irradiation when compared to SpyCI-free strains (Scott et al., 2008). As shown in Figure 1, SpyCI insertion separates mutL, lmrP, tag, and ruvA from their promoter upstream of mutS. The RuvA protein acts as a sliding collar to prevent the unwinding of Holliday junctions during DNA recombination (Kaplan and O’Donnell, 2006). RuvA then recruits RuvB to the junction to promote branch migration, and the complex subsequently recruits RuvC to resolve the Holliday junction into two discrete DNA duplexes (Parsons et al., 1995; Eggleston et al., 1997; Eggleston and West, 2000). Loss of ruvA expression renders a cell sensitive to DNA damage caused by UV irradiation.
FIGURE 2 | Mutation rate and UV sensitivity of S. pyogenes strains. The grouping of the S. pyogenes strains by mutation rate ($\mu$) is shown as in Table 3. Next to the calculated mutation rate, the sensitivity to UV irradiation is shown for each strain. UV exposure time in seconds is indicated at the top and all strains were plated at the same dilution. All experiments were performed at least three times for each strain.

Table 3 | Calculated mutation rates from fluctuation test.

| Strain       | Mutation rate in mutations/generation ($\mu$) | Fold difference in mutation rate* | Phenotype |
|--------------|---------------------------------------------|----------------------------------|-----------|
| Manfredo     | $1.00 \times 10^{-11}$                      | $4.6 \times 10^{-4}$             | Non-mutator |
| NZ131        | $2.19 \times 10^{-8}$                       | 1.0                             | Non-mutator |
| MGAS10750    | $1.04 \times 10^{-7}$                       | 4.7                             | Mutator    |
| 5474-00      | $1.04 \times 10^{-7}$                       | 4.7                             | Mutator    |
| 3470-04      | $1.65 \times 10^{-7}$                       | 7.5                             | Mutator    |
| 270fer       | $2.26 \times 10^{-7}$                       | 10.3                            | Mutator    |
| SF370        | $2.41 \times 10^{-7}$                       | 11.0                            | Mutator    |
| MGAS6180     | $2.47 \times 10^{-7}$                       | 11.3                            | Mutator    |
| MGAS10270    | $3.79 \times 10^{-7}$                       | 17.3                            | Mutator    |
| MGAS10394    | $3.66 \times 10^{-6}$                       | 167.1                           | Hypermutator |

*As compared to strain NZ131.

The mutation rate with confidence limits was calculated as described in the methods. All experiments were performed at least three times for each strain and the data collected was averaged to determine mutation rate in mutations/generation ($\mu$). The mutation rates, as determined by the fluctuation test, clustered into three different groups: non-mutator with a mutation rate $< 10^{-8} \mu$ (Manfredo and NZ131); mutator with a mutation rate of $10^{-7} \mu$ (MGAS10750, 5474-00, 3470-04, 270fer, SF370, MGAS6180, GAS10270); and hypermutator with a mutation rate of $10^{-6} \mu$ (MGAS10394).

(Miyazaki et al., 1989). As seen in Figure 2, the 10 strains showed a wide range of sensitivities following UV irradiation. In general, the presence of SpyCI in a strain correlated with an increased sensitivity to UV-induced cell death as compared to SpyCI-free strain NZ131. Genome strains SF370, MGAS6180, MGAS10394, and MGAS10750 were at least 100-fold more sensitive to UV killing under the test conditions, with strain MGAS10394 being the most sensitive of all. The SpyCI containing M49 clinical strain 5474-00 and 3470-04 were also sensitive to UV irradiation. Not all of the SpyCI strains were equally sensitive, however. Strain MGAS10270 showed only a slight increase in sensitivity (~10-fold) while genome strain Manfredo and clinical strain 270fer were at least as resistant to killing as NZ131. Since little is known of the genetic background of strain 270fer, we can only hypothesize that some other mechanism must compensate for the loss of $ruvA$. However, Manfredo again was divergent from the other genome strains and was as UV irradiation resistant as the wild type strains. These unexpected results necessitated further investigation into strain Manfredo.

STRAIN MANFREDO IS DEFECTIVE FOR SITE-SPECIFIC INTEGRATION

The only SpyCI containing S. pyogenes genome strain that did not have a mutator phenotype or show enhanced sensitivity to UV irradiation was Manfredo. This result was paradoxical since the integrase gene in SpyCIM5 [annotated as $\phi$Man.5 (Holden et al., 2007)] has a deletion of 128 bp that results in not only loss of coding potential but also introduces a frameshift into this gene (Figure 3); in fact, it is annotated as a pseudogene (Holden et al., 2007). Further, the int promoter region in Manfredo is quite divergent from SF370 SpyCIM1 and the other genome strain SpyCI (Figure 3). Since SpyCIM1 integration and excision are crucial to MMR regulation in SF370, the loss of integrase activity in Manfredo could potentially disrupt this growth-dependent process and thus create a permanent mutator phenotype instead of a conditional one as in SF370. However, since such a mutator phenotype was not observed (Figure 2), the potential activity of the integrase gene was investigated.
FIGURE 3 | Alignment of the promoter and 5' ORF of the integrase genes from SpyCIM1 and SpyCIM5. The sequence alignment of the promoters and 5' portion of the int coding regions of SpyCIM1 and SpyCIM5 is shown (labeled M1 and M5, respectively). The predicted promoter for SpyCIM1 int is double underlined, and the single line indicates the probable beginning of the coding region. The SpyCIM5 promoter has significantly diverged from SpyCIM1 and the other SpyCI-like chromosomal islands. The alignment shows that 128 bp have been deleted from the SpyCIM5 int, resulting in not only a loss of genetic information but also the formation of a frameshift. Thus, computational analysis predicts that the Manfredo SpyCIM5 integrase is a non-functional pseudogene.
Strain Manfredo was grown to different stages of growth as determined by spectrophotometry, and mRNA was harvested and converted into cDNA. As can be seen in Figure 4A, transcripts for both mutS and mutL were present throughout growth, and qRT-PCR confirmed that expression was constitutive (not shown). Further, neither mRNA corresponding to the episomal SpyCI attP nor the chromosomal attB which appear following SpyCI excision could be detected: attP is an indication of phage integrase gene activity following excision while attB is contained on the wild type polycistronic mRNA containing both mutS and mutL (see Figure 1). Neither sequence could be detected in strain Manfredo (Figure 4A, attP and attB) nor could the corresponding encoding regions in the chromosomal DNA be detected by PCR (not shown). The absence of these regions resulting from SpyCI excision is in contrast to strain SF370 where these events are linked to the expression of mutL during early logarithmic growth (Scott et al., 2008).

To determine whether the lack of excision was due to the apparent defect in the integrase gene or some other confounding factor, the active integrase gene from SpyCI1 was cloned into vector pCAMP17 where it was placed under the control of the constitutively expressed CAMP promoter (Gase et al., 1999). This vector also contains the S. pyogenes bacteriophage T12 integrase and phage attachment site, mediating integration into the S. pyogenes tmRNA gene (McShan et al., 1997). The resulting plasmid, pWM448, would introduce the active SpyCI1 integrase into the Manfredo chromosome at a distant location and under the control of a promoter that would be unaffected by any SpyCI5 regulation. Introduction of pWM448 into strain Manfredo led to detectable expression of the SpyCI1 integrase gene by qRT-PCR (Figure 4B) with resulting appearance of the attP sequence resulting from excision of SpyCI5 (Figure 4C). Thus, the lack of SpyCI5 excision in strain Manfredo is the result of the deletion in the integrase gene and not from any defects in the attachment sites. No significant difference in the mutation rate or degree of UV sensitivity was observed after introduction of pWM448 (not shown); this was not surprising since Manfredo was essentially wild type for MMR before manipulation. However, mitomycin C treatment, which results in enhanced SpyCI1 excision in SF370 (Scott et al., 2008), inhibited excision of SpyCI5, which suggested that other factors such as DNA binding proteins might influence the ability of the integrase to mediate site-specific recombination in Manfredo (Figure 4C).

**TRANSCRIPTION OF mutL IN SF370 AND MANFREDO**

The lack of detectable SpyCI excision coupled with mutL expression and normal MMR phenotype in strain Manfredo suggests that an alternate promoter, possibly of SpyCI5 origin, controls expression of this gene. Transcriptional analysis by PCR showed mutL is transcribed transiently in SF370 and constitutively in
Manfredo (Figure 5). Treatment of SF370 with mitomycin C leads to intense expression of this transcript by population-wide SpyCI excision. In neither mitomycin C treated nor untreated SF370 is any mRNA detected once the cells reach deep stationary phase (overnight incubation), confirming the silence of the \textit{mutL} gene in this period of the cell’s life. In Manfredo, transcription of \textit{mutL} is constitutive and the amount of mRNA present gradually decreases as growth slows, but some mRNA is still detectable after overnight incubation (Figure 5). Treatment of Manfredo with mitomycin C inhibits transcription of this mRNA, possibly due to the induction of an antirepressor or other DNA binding protein, but appears to increase mRNA levels in the overnight sample compared to the untreated cells, which may result from the induction of damage repair in these cells. These results, coupled with the lack of detection of SpyCI excision, strongly suggest that \textit{mutL} expression in Manfredo is being promoted via some promoter within the \textit{SpyCIM5} that includes at least part of the integrase gene and that regulation of this transcript is fundamentally different from that seen in SF370 SpyCIM1.

**TRANSCRIPTION OF **\textit{mutL}** BY A PROMOTER WITHIN THE **\textit{SpyCIM5} **\textit{int} PSEUDOGENE**

To identify the promoter activating \textit{mutL} in Manfredo, primer extension was used to map the start of \textit{mutS} and \textit{mutL} mRNA in strains NZ131, SF370, and Manfredo. In all three strains, \textit{mutS} transcription initiates from essentially the same location, with reverse transcription terminating 147 bp upstream of the primer start site in NZ131 and SF370 and 146 bp upstream of the primer start site in Manfredo (not shown). Thus, transcription is initiating about 30 bases downstream of the putative start codon as previously determined by \textit{in silico} analysis of all three genome annotations, placing the transcriptional start site at nucleotide 1,754,003 in NZ131, 1,790,110 in SF370, and 1,778,534 in Manfredo. Primer extension for \textit{mutL} in NZ131, SF370, and Manfredo did not produce any substantial, significant, or reoccurring peaks (not shown), suggesting that the reverse transcription is not terminating within the 500 bp window analyzed by the ABI 3730 capillary sequencer and consistent with a polycistronic mRNA initiating upstream of \textit{mutL}. The presence of this polycistronic message is expected in NZ131, which has no SpyCI, and in SF370, which has SpyCIM1 capable of excising from the chromosome (Scott et al., 2008). In Manfredo, where SpyCIM5 is unable to excise from the chromosome, these data support the hypothesis that transcription of \textit{mutL} is driven from a promoter within the \textit{int} pseudogene.

Primer extension of SpyCIM5\textit{int} cDNA revealed the presence of a promoter within the pseudogene that was completely distinct from the one predicted for \textit{int} in SpyCIM5 or the other genome SpyCI (Figure 6). The pattern of \textit{mutL} gene expression (Figure 5) and the non-mutator phenotype (Figure 2) strongly suggests that this promoter must be constitutively expressed; and this expression pattern is in contrast to SF370 where expression of \textit{mutL} only occurs at a specific point in the growth cycle or following DNA damage. This mapped promoter within SpyCIM5\textit{int} corresponded to an \textit{in silico} predicted promoter (Figure 6). Transcription from the promoter region upstream of the \textit{int} pseudogene could never be detected by primer extension (not shown).

To test whether the SpyCIM5 \textit{int} pseudogene region was capable of promoting expression of a downstream gene, the region of the Manfredo chromosome that includes the SpyCIM5 \textit{int} pseudogene, its upstream sequence, and downstream through the beginning of the first codon of \textit{mutL} was fused to the GFP gene (\textit{gfp}). This construct was cloned into Gram-positive shuttle vector pDL276 to create plasmid pWM483, which was introduced into strains Manfredo and SpyCIF-free strain NZ131. As shown in Figure 7, GFP was expressed in Manfredo but not in NZ131, and the expression varied by cell in Manfredo. Treatment of Manfredo (pWM483) with mitomycin C resulted in the inhibition of green fluorescence, mirroring the inhibition of \textit{mutL} mRNA shown in Figure 5. These results suggest that the activity of this promoter associated with the SpyCIM5 \textit{int} pseudogene is dependent upon the genetic background of the cell, possibly requiring some gene product encoded by the SpyCIM5 itself. Removal of the region upstream of the \textit{int} pseudogene ORF (pWM489) resulted in diminished expression in Manfredo but with some weak expression of GFP in NZ131. Sequentially shortening this region (pWM488 and pWM490) did not restore full expression in Manfredo; mitomycin C treatment had little effect since the cells only expressed low levels without treatment. NZ131 expression continued to be at low levels in these truncated constructs. The results suggest that an enhancer exists in Manfredo that promotes efficient transcription of \textit{mutL} from the \textit{int} pseudogene and that SpyCIM5 may encode this enhancer since SpyCI-free NZ131 only weakly expresses this region. Although the potential enhancer of \textit{mutL} transcription in Manfredo is not identified, treatment with
mitomycin C, which induces SOS repair, results in inhibition of GFP expression. Thus, the potential enhancer may be a DNA binding protein that is sensitive to activated RecA cleavages seen in other cells.

**DISCUSSION**

The results presented here demonstrate that a SpyCI-associated mutator phenotype is not unique to M1 genome strain SF370. This report examined the effect of SpyCI integration between the MMR genes *mutS* and *mutL* in six genome strains of *S. pyogenes*. We determined the associated mutator phenotypes as measured by spontaneous mutation rate and sensitivity to UV irradiation to better understand the consequences of prophage-like SpyCI integration at this universally conserved site. Further, we presented evidence that in the M5 genome strain, Manfredo, a novel promoter within the SpyCIM5 integrase pseudogene is used to rescue the expression of *mutL* and the downstream genes following CI genome decay. Interestingly, this promoter is inhibited by mitomycin C (and presumably the induction of SOS repair). It is not yet known whether this apparent mechanism of gene regulation is the result of positive selection or merely a circumstantial byproduct of evolution of this compensatory promoter that rescues MMR expression in Manfredo. Collectively, these results provide a snapshot of how SpyCI control of DNA repair in *S. pyogenes* occurs by more than one mechanism, depending upon the integrity of the SpyCI genome integrative functions. These studies provide insight into the co-evolution of prophage-like CIs and their host *S. pyogenes* cells.

Most of the SpyCI containing strains exhibited a mutator phenotype and were at least moderately sensitive to UV irradiation.
Strains MGAS10750, 5474-00, 3470-04, SF370, MGAS6180, and MGAS10270 all fell into this category. Since the UV irradiation data is qualitative in nature and may be influenced by other genes in the cell, it is difficult to determine definitively how well the mutation rate correlates with the degree of UV sensitivity. For example, strain 3470-04 appears to be more sensitive to UV light than the rest of the strains in this category, despite having a moderate mutation rate by comparison (Figure 2). It is possible that 3470-04 may have a defect in another DNA repair pathway that is independent of the MMR operon, which contributes to the observed phenotype. Creation of isogenic strains differing only by the presence of SpyCI integrated into mutL will help to clarify the specific contribution made by these CIs to the host phenotype.

Of particular interest is strain MGAS10394 with a mutation rate at least a full degree of magnitude higher than all of the other strains. MGAS10394 also has the highest degree of UV sensitivity; only a few colonies survived after any exposure to UV irradiation. The high mutation rate and low tolerance to UV irradiation suggests that the SpyCIM6 may exist more frequently in the integrated state and less as an episome, which would shift the MGAS10394 population toward the mutator phenotype. One of the regions of genetic diversity in the SpyCI is the predicted
region controlling lysogeny (i.e., the predicted repressor, antirepressor, and intervening operator sequence). Phylogenetic analysis of this region suggests that this region may be undergoing rapid evolution (Figure 8). A phylogenetic tree built upon the DNA sequence of this lysogeny module reveals that recombination has acted upon this region to create a high level of diversity, and this variation could alter the affinity that the repressor and antirepressor have for the operator DNA sequence as well as their sensitivity to proteases that could lead to cleavage and induction of excision. Further studies are needed to test this hypothesis. The operator sequence between the predicted repressor and antirepressor shows that several groupings exist (Figure 8A). SpyCIM1, SpyCIM2, SpyCIM28, and SpyCIM53 all have nearly identical operator sequences; SpyCIM5 also belongs to this group but has
undergone a number of changes at the DNA level. An interesting side effect of tipping the regulation toward integration and the mutator phenotype is that the SpyCI itself would be subject to increased rates of mutation as a result of SpyCI integration, there are notable exceptions to this generality: Manfredo, 270fer, and MGAS10394 demonstrated either a wild type phenotype, a mixed mutator phenotype, or a hypermutator phenotype, respectively. These exceptions may represent cases of newly acquired CI that have yet to undergo rounds of selective pressure as well as severely decayed phages that have been optimally streamlined to accommodate a more permanent residence in the host chromosome. The evolution, dissemination, and host range of SpyCI will be important topics for future studies to help clarify the impact that they have on streptococcal survival and virulence.

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