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1	Genome integration and excision by a new Streptomyces		
2	bacteriophage, φJoe		
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20 Abstract

21 Bacteriophages are the source of many valuable tools for molecular biology and genetic 22 manipulation. In Streptomyces, most DNA cloning vectors are based on serine integrase site-23 specific DNA recombination systems derived from phage. Because of their efficiency and simplicity, serine integrases are also used for diverse synthetic biology applications. Here we 24 present the genome of a new *Streptomyces* phage, ϕ Joe, and investigate the conditions for 25 integration and excision of the \u03c6Joe genome. \u03c6Joe belongs to the largest Streptomyces phage 26 27 cluster (R4-like) and encodes a serine integrase. The attB site from S. venezuelae was used 28 efficiently by an integrating plasmid, pCMF92, constructed using the ϕ Joe *int/attP* locus. The attB site for \u03c6Joe integrase was occupied in several Streptomyces genomes, including S. 29 30 *coelicolor*, by a mobile element that varies in gene content and size between host species. Serine integrases require a phage-encoded recombination directionality factor (RDF) to 31 activate the excision reaction. The \$\phi_Joe RDF was identified and its function was confirmed in 32 33 vivo. Both the integrase and RDF were active in *in vitro* recombination assays. The \$dot Joe sitespecific recombination system is likely to be an important addition to the synthetic biology and 34 genome engineering toolbox. 35

36 Importance

Streptomyces spp. are prolific producers of secondary metabolites including many clinically 37 38 useful antibiotics. Bacteriophage-derived integrases are important tools for genetic 39 engineering as they enable integration of heterologous DNA into the Streptomyces chromosome with ease and high efficiency. Recently researchers have been applying phage 40 integrases for a variety of applications in synthetic biology, including rapid assembly of novel 41 42 combinations of genes, biosensors and biocomputing. An important requirement for optimal 43 experimental design and predictability when using integrases, however, is the need for multiple enzymes with different specificities for their integration sites. In order to provide a 44 broad platform of integrases we identified and validated the integrase from a newly isolated 45 Streptomyces phage, oJoe. oJoe integrase is active in vitro and in vivo. The specific 46

- 47 recognition site for integration is present in a wide range of different Actinobacteria, including
- *Streptomyces venezuelae*, an emerging model bacterium in *Streptomyces* research.

49 Introduction.

50 Over the past few decades, serine integrases have become widely established as tools 51 for genome engineering and synthetic biology (1, 2). Serine integrases are phage-encoded, DNA site-specific recombinases that mediate recombination between two short (<50 bp) 52 sequences. The integration reaction occurs during the establishment of lysogeny, during 53 which the integrase causes a single crossover between the attB site on the bacterial 54 55 chromosome and the attP site on the circularised phage genome leading to the integrated phage DNA flanked by the recombinant sites, attL and attR (1, 3). Integrase dimers bind to 56 the two *att* sites and produce double-strand breaks with 2 bp overhangs (3, 4); the cut ends 57 are then exchanged and the DNA backbone is re-ligated to produce the recombinant products 58 (5). The attL and attR sites each contain reciprocal halves of the attP and attB sites (6). As 59 integrases are unable to use *attL* and *attR* as substrates without an accessory protein, the 60 recombination directionality factor (RDF), the integrated phage genome is stable until the 61 RDF-encoding gene is expressed during prophage induction (3). Recombination between attL 62 63 and *attR* is the excision reaction and is essentially the reverse of integration, releasing the phage genome and reforming attP and attB. Whilst only integrase is required to mediate 64 integration, excision requires both integrase and the RDF. Genome engineers have exploited 65 66 these systems to integrate genes of interest into a specific site on the chromosome, which can 67 either be the endogenous attB or an introduced attB or attP used as a docking site (1). The 68 simplicity of the serine integrase mediated site-specific recombination systems means that 69 they are reliably portable to heterologous hosts where DNA can be integrated stably and in single copy. 70

The simple requirements of serine integrases make them amenable to a wide variety of applications. The earliest examples of this were to integrate an *attP* plasmid into a target genome containing the cognate *attB* (or *vice versa*) (7), allowing stable delivery of genes into diverse species, including bacteria (6, 8–10), mice (11), mosquitos (12) and humans (13). More complex genetic engineering approaches use integrases in *in vitro* ordered assembly of

76 multiple DNA fragments (14, 15). In vivo genome manipulations can also be achieved either by iterative rounds of recombination (16, 17) or multiplexing orthogonal integrases/att sites 77 (18). Integrase mediated DNA rearrangements can also be used to provide permanent genetic 78 memory in novel types of biosensors (19, 20). Some applications, such as post factum 79 80 modifications (15) or biocomputing (19, 21), need controlled excision and this requires integrase and its cognate RDF. The RDF binds directly to the integrase protein and is thought 81 to induce a conformational change that allows attL and attR to be used as recombination 82 substrates whilst inhibiting recombination of attB and attP (22, 23). 83

84 A limiting factor for the use of serine integrases for complex, multiplexed applications is the number of well-characterized integrases and, perhaps more pressingly, RDFs. Only 85 seven integrase/RDF pairs have been characterized to date (from phages TP901-1 (24), ϕ C31 86 (22), ϕ BT1 (25), Bxb1 (23), ϕ Rv1 (26) and SPBc (27), and from the excisive element of 87 Anabaena and Nostoc cyanobacteria species (28)), but many more integrases have been 88 studied without their RDFs (1, 2, 29–31). Integrase genes are easily identified by comparative 89 90 sequence analysis and, when the integrase is prophage encoded, the attachment sites can also be predicted (31). RDFs, however, are far more difficult to predict because known 91 examples share little sequence homology, vary markedly in size and also differ in gene 92 93 location in phage genomes (1). Expansion of the available arsenal of serine integrases and 94 RDFs is desirable to enable advanced synthetic biology applications.

Phages that encode serine integrases are prevalent in Gram-positive bacteria, and in particular in Actinobacteria. Here, we describe a newly isolated *Streptomyces* phage, ¢Joe, and its serine integrase (Int) that is only distantly related to characterized integrases. ¢Joe Int is active *in vivo* in *Streptomyces* and *E. coli*, the integrase protein is readily purified and is able to carry out efficient *in vitro* recombination. We also describe the ¢Joe RDF, a 6.8 kDa protein that is able to promote excisive recombination and inhibit integration.

101

102 Materials and Methods

103 Growth media

Escherichia coli strains were generally grown in LB, except where otherwise noted. Antibiotics were added for selection where appropriate (apramycin: 50 μg/ml, chloramphenicol: 50 μg/ml, kanamycin: 50 μg/ml, ampicillin: 100 μg/ml). Preparation of competent cells and transformation of *E. coli* were performed as described in Sambrook *et al.*, 2001 (32). *Streptomyces* strains were grown on Mannitol Soya agar (33) supplemented with 10 mM MgCl₂ for plating conjugation mixtures and antibiotics, where required (apramycin: 50 μg/ml, nalidixic acid: 25 μg/ml).

Phage Isolation. The procedures for isolation, plating and titre of phage with *Streptomyces* 111 as the isolation host are described in detail in Kieser et al., 2000 (33). Raw soil samples were 112 enriched for environmental phage using S. coelicolor M145 as a propagation host (34). Briefly, 113 3 g of soil was added to 9 ml Difco[™] nutrient (DN) broth (BD Diagnostics, Oxford, UK) 114 supplemented with 10 mM CaCl₂, 10 mM MgSO₄ and 0.5% glucose. Streptomyces spores 115 were added to a concentration of 10⁶ colony forming units/ml (cfu/ml) and incubated at 30°C 116 with agitation for 16 h. Soil and bacteria were removed by centrifugation and filtration through 117 118 a 0.45 µm filter. A dilution series of the filtrate in SM buffer (100 mM NaCl, 8.5 mM MgSO₄, 50 mM Tris-HCl pH 7.5, 0.01% gelatin) was plated with S. coelicolor spores to isolate single 119 plaques. Phage were recovered from single, well-isolated plaques by single plaque soak outs 120 in DN broth and re-plated with the host strain for three rounds of plague purification. A high 121 122 titre phage preparation was generated from plates inoculated with sufficient plaque forming units (pfu) to generate almost confluent lysis (33). The phage suspensions were filtered, 123 pelleted by ultracentrifugation and resuspended in 0.5 ml SM buffer (35). The concentrated 124 phage were further purified by caesium chloride isopycnic density gradient centrifugation (36). 125

Next Generation Sequencing. Phage DNA was extracted by phenol:chloroform purification
 (32) and the presence of pure phage DNA was confirmed by restriction digest. Phage DNA

128 was sequenced and assembled in collaboration with Dr Darren Smith at NU-OMICS (Northumbria University). DNA was prepared for next generation sequencing on the Ilumina 129 MiSeq platform using the Nextera XT library preparation kit (Illumina, Saffron Waldon, UK). 130 Samples were loaded and run using a 2 × 250 cycle V2 kit. DNA samples were diluted to 0.2 131 132 ng/µl, prior to normalization and pooling. Paired end sequencing reads were provided as FASTQ files (NU-OMICS, Northumbria University, Newcastle, UK) and subjected to 133 downstream analysis. ORF prediction and annotations were assigned using DNA master 134 135 (Lawrence lab, Pittsburgh, PA), Glimmer (37) and Genemark (38). The annotated genome 136 sequence was submitted to GenBank (accession number: KX815338).

Electron Microscopy. Purified phage were negatively stained with uranyl acetate (39) and
 imaged in a FEI Tecnai 12 G2 transmission electron microscope fitted with a CCD camera.

139 Mass Spectrometry. Whole phage samples were run into a 7 cm NuPAGE Novex 10% Bis-140 Tris gel (Life Technologies) at 200 V for 6 mins. The total protein band was excised and 141 digested in-gel with 0.5 µg trypsin, overnight at 37°C. Peptides were extracted, concentrated 142 and loaded onto a nanoAcquity UPLC system (Waters) equipped with a nanoAcquity Symmetry C₁₈, 5 µm trap (180 µm x 20 mm Waters) and a nanoAcquity HSS T3 1.8 µm C₁₈ 143 capillary column (75 µm x 250 mm, Waters). The nanoLC system was interfaced with a maXis 144 HD LC-MS/MS system (Bruker Daltonics) with CaptiveSpray ionisation source (Bruker 145 Daltonics). Positive ESI-MS and MS/MS spectra were acquired using AutoMSMS mode. 146 147 Instrument control, data acquisition and processing were performed using Compass 1.7 148 software (microTOF control, Hystar and DataAnalysis, Bruker Daltonics. The collision energy 149 and isolation width settings were automatically calculated using the AutoMSMS fragmentation table, absolute threshold 200 counts, preferred charge states: 2 - 4, singly charged ions 150 151 excluded. A single MS/MS spectrum was acquired for each precursor and former target ions 152 were excluded for 0.8 min unless the precursor intensity increased fourfold. Protein 153 identification was performed by searching tandem mass spectra against the NCBInr database

using the Mascot search program. Matches were filtered to accept only peptides with expectscores of 0.05 or better.

156 Plasmid Construction. Plasmids used in this study are listed in Table 1 and oligonucleotides in Table 2. General molecular biology techniques including plasmid DNA preparation, genomic 157 158 DNA preparation, restriction endonuclease digestion and agarose gel electrophoresis were performed as described in Sambrook et al., 2001 (32). In-fusion cloning technology (Clontech) 159 was generally used for construction of plasmids. Polymerase chain reaction (PCR)-amplified 160 161 DNA was generated using primers with Infusion tags for insertion into plasmid vectors, which 162 had been cut with restriction endonucleases. The
\$\phiJoe\$ integrating plasmid, pCMF92, was created by Infusion cloning of the \$\phiJoe int gene and attP region, obtained by PCR with Joe 163 Int-attP F/R primers and oJe genomic DNA as a template, into the 3.1 kbp EcoRI-SphI 164 fragment from pSET152. Plasmid pCMF91 was generated by inserting the amplified attP site 165 prepared using \$\phi_Joe genomic DNA as a template and primers Joe attP F/R into EcoRI 166 167 linearized pSP72. The integration sites in S. coelicolor were named attLsc and attRsc and were amplified from *S. coelicolor* gDNA using Joe *attB1* F/R and Joe *attB2* F/R. The *attB* site 168 from S. venezuelae (attBsv) was amplified using S. venezuelae gDNA with Joe attB Sv F/Joe 169 attB R primers. All three attachment sites were inserted into EcoRI-linearized pGEM7 to 170 produce pCMF90, 94 and 95, respectively. The reconstituted S. coelicolor attB sequence 171 (attBsc) was prepared from two complementary oligonucleotides, Joe attB Recon F and Joe 172 attB Recon R (Ultramer primers, IDT) that were annealed and inserted into EcoRI-linearized 173 174 pGEM7 to produce pCMF97. pCMF98 contains the \$\phiJoe attLsv and attRsv sites in head-totail orientation and was isolated by transformation of an in vitro recombination reaction 175 between pCMF91 (containing ϕ Joe *attP*) and pCMF95 (containing *attBsv*) into *E. coli*. The 176 attLsv and attRsv sites in pCMF98 were confirmed by Sanger sequencing (GATC Biotech Ltd, 177 178 London, UK). The recombination reporter plasmid pCMF116 was constructed by PCR amplification of *lacZa* using *E. coli* MG1655 gDNA (40) as a template and Joe BzP forward 179 180 and reverse primers encoding the ϕ Joe *attBsv* and ϕ Joe *attP*, respectively, resulting in the

181 *attBsv* and *attP* sites flanking the *lacZa* gene in head-to-tail orientation. The amplified DNA 182 was inserted into XmnI-linearized pACYC184. pCMF103 was constructed in the same way 183 as pCMF116 except that Joe LzR F/R primers containing the ϕ Joe *attLsv* and *attRsv* sites 184 were used.

185 The integrase expression plasmid for protein purification, pCMF87, was constructed by insertion of a PCR fragment encoding the \$\phiJoe int gene, amplified from \$\phiJoe gDNA using 186 primers Joe H6-Int F/R, into Ncol-linearized pEHISTEV expression vector. ϕ Joe g52, encoding 187 188 the RDF, was PCR-amplified from ϕ Joe gDNA using primers Joe MBP-g52 F/R and inserted 189 into pETFPP 2 MBP-tag expression vector linearized by PCR with CleF/R to create pCMF96. For in vivo recombination assays the integrase expression plasmid, pCMF107, was 190 constructed by insertion of a PCR fragment encoding the \$\phiJoe int gene, amplified from \$\phiJoe int gene, 191 192 gDNA using primers Joe pBAD Int F/R, into Ncol-linearized pBAD-HisA expression vector. A φJoe gp52 and integrase co-expression plasmid, pCMF108, was created by amplification of 193 each gene using Joe pBAD gp52 F/R and Joe pBAD Int Co-Ex F/Joe pBAD Int R primers, 194 respectively, and insertion of both PCR products simultaneously into pBAD-HisA. The co-195 expression insert from pCMF108 was subsequently PCR-amplified using Joe H6-gp52 F/Joe 196 H6-Int R primers and transferred to Ncol-linearized pEHISTEV to produce an alternative 197 expression vector, pCMF117. 198

199 Conjugation and integration of plasmids in Streptomyces. Transfer of plasmids into Streptomyces strains was performed according to the procedures described by Kieser et al., 200 201 (2000) (33). Conjugation donors were produced by introduction of plasmids into the nonmethylating *E. coli* strain, ET12567, containing an RP4 derivative plasmid (pUZ8002), by 202 203 transformation. Recipient Streptomyces spores were used at a concentration of 10⁸/ml, mixed 204 with the *E. coli* donors, plated onto mannitol soya agar supplemented with 10 mM MgCl₂ with no antibiotic selection and incubated at 30°C overnight. Plates containing the donor cells were 205 overlaid with 1 ml water containing 0.5 mg nalidixic acid (for E. coli counterselection) and 206 antibiotic for selection of exconjugants (apramycin) before further incubation of all plates at 207

30°C for three days. Integration efficiency was calculated as the number of apramycin resistant colonies/10⁸ cfu (8).

210 Protein Purification. E. coli BL21(DE3) containing the relevant expression plasmid were grown (37°C with agitation) in 500 ml 2YT medium (1.6% w/v tryptone, 1.0% w/v yeast extract, 211 0.5 w/v NaCl) to mid-exponential growth phase. The cultures were rapidly chilled on ice for 15 212 min, IPTG was added (final concentration 0.15 mM) and the cultures were further incubated 213 (17°C, 16 h, with agitation). Cells were harvested by centrifugation, resuspended in 20 ml lysis 214 buffer (1 M NaCl, 75 mM Tris pH 7.75, 0.2 mg/ml lysozyme, 500 U Basemuncher 215 216 Endonuclease; Expedeon Ltd.) and incubated on ice (30 min). The cells were lysed by sonication and debris was removed by centrifugation (18,000 g, 5 min, 4°C). The supernatant 217 was applied to a 5 ml HisTrap FF crude column that had been pre-equilibrated with binding 218 buffer (20 mM sodium phosphate, 0.5 M NaCl, 20 mM imidazole, pH 7.4) on an ÄKTA pure 219 25 chromatography system (GE Healthcare). Bound, his-tagged protein was eluted with a step 220 gradient of binding buffer containing 125 mM and 250 mM imidazole. Imidazole was removed 221 222 from the eluted fractions by pooling the fractions containing the desired protein and applying the pooled solutions to a HiPrep 26/10 desalting column (GE Healthcare) equilibrated with 223 imidazole-free binding buffer. Finally, the protein extracts were subjected to size exclusion 224 225 chromatography on a HiLoad 16/60 Superdex column. Purified protein fractions were 226 concentrated in a Vivaspin sample concentrator (GE Healthcare) and quantified by 227 absorbance at 280 nm on a Nanodrop spectrophotometer (Thermo Scientific). Protein analysis 228 was performed by denaturing acrylamide gel electrophoresis using pre-made gels (4-12% 229 gradient acrylamide; Expedeon Ltd.); gels were stained with InstantBlue (Expedeon, Ltd.). For storage, an equal volume of 100% glycerol was added to protein samples before freezing at -230 231 80°C.

In vitro Assays. Recombination reactions (final volume of 20 μl) were carried out in φC31
RxE buffer (10 mM Tris pH 7.5, 100 mM NaCl, 5 mM DTT, 5 mM spermidine, 4.5% glycerol,
0.5 mg/ml BSA) (41), Bxb1 RxE buffer (20 mM Tris pH 7.5, 25 mM NaCl, 1 mM DTT, 10 mM

235 spermidine, 10 mM EDTA) (23) or TG1 RxE (as Bxb1 RxE plus 0.1 mg/ml BSA) (42). Integrase and RDF proteins were added at the concentrations indicated for each experiment. 236 Plasmids containing the recombination substrates were used at 100ng per reaction. Reactions 237 were either incubated at 30°C for 2 h (to reach steady state) or for specified times. Reactions 238 239 were stopped by heat (10 min, 75°C), the buffer was adjusted to be compatible with restriction enzymes and the plasmids were digested with XhoI (NEB). The linearized reaction mixtures 240 were run on a 0.8% agarose gel and the relative band intensities were measured to assess 241 activity. Recombination efficiencies were calculated as intensity of product band(s)/sum 242 243 intensity of all bands.

Bioinformatics. The \$\phi Joe genome was visualized using DNAplotter (43). The attB DNA 244 alignment and logo consensus sequence were created with Jalview (44). Protein sequence 245 alignments for visual presentation were produced using the Clustalw (45) program within the 246 Bioedit suite (46). Protein alignments for phylogenetic analysis were produced using Clustal 247 Omega (47) and maximum likelihood trees were created in Mega6 (48). The BLOSUM62 248 249 similarity matrix was used for protein alignment and annotation (49). Structural alignment of the small RDF proteins was carried out with Promals3D (50). Band densities for in vitro assays 250 were measured using the FIJI GelAnalyzer module (51). Accession numbers for all sequences 251 used here are provide in Table S2. 252

253 Results and Discussion.

254 Isolation of actinophage ϕ Joe and genome sequence. Raw soil samples were enriched 255 for environmental phage using S. coelicolor strain M145 as a propagation host. The phage chosen for further analysis, \$Joe, is a siphovirus with a capsid diameter of 46.5 nm (SD 1.6 256 nm, n=9) and a long flexible tail of 199.5 nm (SD 12 nm, n = 8) with clear striations visible in 257 most images (Fig. 1). ϕ Joe is able to plaque on a broad range of Streptomyces hosts, 258 259 producing lytic infection of seven out of nine species tested (Table 3). Saccharopolyspora erythraea (formerly Streptomyces erythraeus) and Streptomyces venezuelae were resistant 260 261 to infection.

Genomic DNA was extracted from high titre ϕ Joe suspensions (>10¹⁰ pfu/ml) and 262 sequenced on the Illumina MiSeq platform with 2,542x coverage. The phage genome is 263 48,941 bases (Accession: KX815338) with a GC content slightly lower than the host bacteria; 264 65.5% compared to ~72% for most Streptomyces species. BLASTn was used to measure 265 266 nucleotide identity for the closest relatives to \$\$ Joe; the generalized transducing phage \$\$ CAM 267 (52) and two newly sequenced Streptomyces phages, Amela and Verse (Fig. S2), are 73, 76 and 76% identical, respectively, in global alignments. The \$\phi Joe genome contains 81 predicted 268 open reading frames (Fig. 2), the majority of which have similar amino acid sequences to the 269 270 three phages above and the well characterized R4 phage (53). Notably, similarity to ϕ Joe integrase (gp53) is absent from each of the closest genome matches but is instead present in 271 several more distantly related phages (Fig. 3), indicative of phage mosaicism (54). Specifically, 272 φJoe integrase is homologous to the uncharacterized integrases from five complete phages -273 274 Lannister (78% amino acid identity), Zemyla (74%), Danzina (73%), Lika (73%) and Sujidade (73%) (Fig. 3). Comparison to known integrases suggests that the catalytic serine is likely to 275 be at position 46 in the protein sequence (VRLSVFT). 276

Purified phage particles were submitted for shotgun LC-MS/MS analysis to determine
 the structural proteome. At least one peptide match was detected from fourteen \$\phiJoe\$ gene

products, five of which have predicted functions – Portal, Capsid, Tail Tape Measure, Scaffold,
Head-Tail Adaptor (Figure 2, Table S1). The remaining nine gene products have no known
function but all cluster close to the predicted structural genes within a region of the genome
spanning ~21 kbp.

283 integration systems, the attP site lies adjacent to the int gene encoding the integrase. The attP 284 285 sites for serine integrases are characteristically about 45 to 50 bp in length and contain inverted repeat sequences flanking a spacer of approximately 20 bp (3, 55). Examination of 286 the ϕ Joe genome identified a candidate *attP* site located 18 bp upstream of the *int* gene. A 287 plasmid, pCMF92, was constructed by replacing the ϕ C31 *int/attP* locus from the widely used 288 integrating vector pSET152, with the ϕ Joe *int/attP* locus (Fig.S1). Integration of pCMF92 would 289 290 confirm whether the integrase is functional, the nature of the *attP* site and, by rescuing the DNA flanking the integrated plasmid, the identity of the *attB* site could be deduced (Fig. 4). 291 pCMF92 was introduced into S. coelicolor J1929 and S. lividans TK24 by conjugation and 292 293 apramycin resistant colonies were obtained, but the frequencies were low (10⁻⁴ to 10⁻⁵ exconjugants/cfu) compared to other integrating vectors (10^{-2} to 10^{-3} exconjugants/cfu) (9, 294 18). To test whether integration was site-specific, four *S. coelicolo*r:pCMF92 cell lines were 295 amplified from independent exconjugants and the genomic DNAs were analysed by Southern 296 297 blotting using a probe derived from the \$\phi Joe int gene. In the four cell lines pCMF92 had 298 integrated into one of two different integration sites, as revealed by hybridisation of the probe 299 to two different restriction fragments (data not shown).

We then sought to characterize the two integration sites for pCMF92 in *S. coelicolor* by rescuing the integrated plasmids along with flanking DNA into *E. coli*. In pCMF92 there is 3.9 kbp of DNA between the ¢Joe *attP* site and the PstI cleavage site that contains the plasmid origin of replication and the apramycin resistance gene (Fig. S1). Genomic DNA from two *S. coelicolor*:pCMF92 cell lines, each containing pCMF92 integrated into one of the two different integration sites, was digested with PstI endonuclease, self-ligated and introduced into *E. coli*

306 DH5 α by transformation. The rescued plasmids were sequenced over the recombination sites 307 to validate the nature of the ϕ Joe *attP* site and to identify the chromosomal positions of the 308 two *S. coelicolor* integration sites. The ϕ Joe *attP* site was confirmed to be \leq 50 bp and the 5' 309 GG dinucleotide at the centre of an imperfect inverted repeat is predicted to be where the 310 crossover occurs (Fig. 4A).

The two S. coelicolor integration sites for pCMF92 are located 3.9 kbp apart, separated 311 by an apparent mobile genetic element comprising sco2603, encoding a putative serine 312 integrase with 68% identity to \$\$ Joe integrase and two further genes (Fig. 4B). Its product, 313 SCO2603, is 68% identical to ϕ Joe integrase. We hypothesized that the ϕ Joe integrating 314 plasmid is inefficient in S. coelicolor because an ancestral and optimal attB site is occupied by 315 the SCO2603-encoding element. The two integration sites for pCMF92 in S. coelicolor were 316 therefore called *attLsc* and *attRsc* to reflect the provenance of the sites containing the mobile 317 element. To test this hypothesis, the sequence of the ancestral *attB* site, *attBsc*, was predicted 318 by removing the sequence between *attLsc* and *attRs*c, including the *attP* moieties that would 319 320 have originated from the inserted mobile element (Fig. 4C). The reconstituted attBsc was used to interrogate the GenBank *Streptomyces* database for closely related extant sequences. 321 Three species were chosen from the top ten hits returned (S. avermitilis, S. albus and S. 322 323 venezuelae, Fig. 4D) and assayed for in vivo integration efficiency. S. venezuelae was the 324 only host to support highly efficient integration after conjugation with pCMF92, 160-fold greater frequency than S. coelicolor and 1,600-fold greater than S. lividans (Fig. 5A). The integration 325 frequencies for pCMF92 into S. venezuelae are similar to those reported for other 326 characterized serine integrases (9, 18) and we demonstrate below that the attB site from S. 327 venezuelae, attBsv, is indeed used efficiently by \$Joe integrase. Plasmid pCMF92 could 328 329 therefore be used as a new integrating vector for use in this newly emerging model system for Streptomyces research. 330

The *S. venezuelae attBsv* site was used as a BLASTn query to estimate the prevalence
 of potential \$\$\phi\$Joe insertion sites in sequenced species. In many instances, each half of the

333 query sequence matched separate locations in the target genome, suggesting that ϕ Joe-like attB sites are frequently occupied by either a prophage or a similar mobile element to that 334 observed in S. coelicolor J1929. Hits were subsequently filtered for matches of at least 80% 335 coverage with an e-value of $<1 \times 10^{-10}$ and a bit score >75, which revealed numerous apparently 336 337 unoccupied \$\phiJoe attB sites in diverse Streptomyces, Kitasatospora and Dermacoccus species (Fig. 4D). Generally, the *attB* site for ϕ Joe and the SCO2603 integrase-encoding elements is 338 located 74bp from the end of an ORF encoding a SCO2606-like predicted B12 binding 339 340 domain-containing radical SAM protein. Insertions this close to the end of an ORF may not 341 necessarily cause loss of function of the gene product and this could explain the prolific number of mobile elements that use this locus as an insertion site. Other than the 342 343 recombination genes, the genetic content of the mobile elements located here varies markedly in different bacterial species (Fig. S2). Some *Streptomyces* strains have an almost identical 344 SCO2603-containing genetic element to S. coelicolor J1929 (e.g. WM6391), others have no 345 genes other than the recombination genes (e.g. NRRLF-5123) and some contain up to 40 kbp 346 between the predicted attL and attR sites (Fig. S2). 347

348 to have broad appeal as a bioengineering tool it must be functional in heterologous hosts. As 349 350 a proof of principle, we tested the activity of ϕ Joe integrase in *E. coli* by cloning the integrase 351 gene into an arabinose-inducible expression vector, pBAD-HisA, to produce pCMF107. Meanwhile, we constructed a reporter plasmid, pCMF116, containing the *E. coli* $lacZ\alpha$ gene 352 353 flanked by \$\phiJoe attBsv and attP sites in head to tail orientation (Fig. S3). Both plasmids were introduced into E. coli TOP10 cells (Invitrogen) by co-transformation and plated on selective 354 agar plates containing 0.2% L-arabinose and 80 µg/ml X-Gal. pBAD-HisA lacking an insert 355 356 was used as a negative control. All of the transformants were white in the presence of ϕ Joe int, indicating efficient recombination between the attBsv and attP sites leading to loss of the 357 *lacZa* gene (Fig. 5B & S3). ϕ Joe integrase and its cognate *attBsv* and *attP* sites are, therefore, 358 active in *E. coli*. 359

360 Another key application for serine integrases is for *in vitro* combinatorial assembly of genes for optimising expression of metabolic pathways (14, 15). In this application different 361 integrases are used to join (by recombination) specific pairs of DNA fragments tagged with 362 their cognate attachment sites. In theory this procedure can be multiplexed to assemble many 363 364 DNA fragments together using different, orthogonally acting integrases. The aim is to generate artificial operons with defined or random order. To test the suitability of \$\$ Joe Int for in vitro 365 366 recombination reactions, the integrase gene was cloned into the His-tag expression vector 367 pEHISTEV and purified after overexpression in E. coli. In vitro recombination assays were carried out with ϕ Joe attP (pCMF91) versus each of attBsc, attLsc, attRsc and attBsv 368 (pCMF97, pCMF90, pCMF94 and pCMF95, respectively) and using a range of \$\phi_Joe integrase 369 370 concentrations. Successful recombination between attachment sites produces a co-integrant 371 plasmid, which can be distinguished from the substrate plasmids by a restriction digest and 372 agarose gel electrophoresis (Fig. S3). In this assay, recombination was undetectable when attLsc (pCMF90) or attRsc (pCMF94) were used with attP (pCMF91) as substrates. A small 373 374 amount of recombination was observed ($\leq 2\%$, Fig. S4) when the reconstituted *attBsc* (pCMF97) was used with attP (pCMF91). However, consistent with the observations in E. coli 375 376 and in Streptomyces, the S. venezuelae attBsv site (pCMF95) was a highly efficient substrate for recombination with the ϕ Joe *attP* site. ϕ Joe integrase was effective over a broad range of 377 concentrations (50 - 1000 nM) (Fig. 5C & S4). Using 200 nM integrase, detectable 378 recombination product was produced after ~10-15 min, and after 2 h approximately 70% of 379 the substrate molecules were converted to product (Fig. 5C & D). 380

There are only 6 bp that differ between *attBsc* and *attBsv*, and all the differences are on the left-hand arm of the *attB* sites (Fig. 4C). Previously, a mutational analysis of the ϕ C31 *attB* site showed that mutationally sensitive bases occur 2, 15 and 16 bases to either side of the crossover dinucleotide (56). As two of the differences between *attBsc* and *attBsv* are also 2 and 16 bases from the putative crossover 5'GG (Fig. 4C), these base pair differences might account for the poor activity of *attBsc* in the *in vitro* assays.

387 **Identification and validation of the** ϕ **Joe RDF protein, gp52.** Although there are dozens of serine integrases that have been described in the literature, there are only seven published 388 RDFs for serine integrases (ϕ C31 gp3 (22), ϕ BT1 gp3 (25), Bxb1 gp47 (23), TP901 ORF7/Xis 389 (24), Anabaena/Nostoc Xisl (57), SPBc SprB (27), and $\phi Rv1 Rv1584c/Xis$ (26)). The Bxb1 390 391 and ϕ C31 RDFs are amongst the largest of these RDF proteins (approximately 27.5 kDa, 250 amino acids) and their genes are located in proximity to the phage DNA replication genes. 392 Both RDFs have functions during phage replication in addition to acting as RDFs but they are 393 evolutionarily unrelated (25, 58). The RDFs from ϕ BT1 and another ϕ C31-like phage, TG1, 394 are close relatives of the ¢C31 RDF at the sequence level (85% and 59% identical, 395 respectively); furthermore, the ϕ BT1-encoded RDF acts on ϕ C31 integrase and vice versa 396 (25). The ϕ Rv1 and SPBc RDFs are located within 1 or 2 ORFs of the *int* gene, a feature which 397 398 is reminiscent of the xis genes that act with tyrosine integrases. $\phi Rv1$, SPBc, TP901 and 399 Anabaena/Nostoc RDFs are much smaller proteins than ϕ C31 gp3 or Bxb1 gp47 (58 and 110 400 amino acids). Given the variation in RDF size, sequence and genomic location, there are no 401 sound generalizations yet for identifying new RDFs in phage genomes.

A list of four candidate genes (g40, 43, 49 and 52) for the \$\$Joe RDF was drawn up 402 based on comparable size to known, small RDFs and genomic location (i.e. not located 403 404 amongst the late/structural genes) (Fig. 2). One of the potential RDF genes (g52) is adjacent 405 to *int* in the ϕ Joe genome, but it is transcribed divergently, with the *attP* site situated between int and g52 (Fig. 2). Unlike the other candidate RDFs, gp52 homologues are only found in 406 those phages with ϕ Joe-like integrases (Fig. 3), and phylogenetic analysis of gp52 and the 407 integrase indicated that both proteins have followed a parallel evolutionary path (Fig. S5). 408 409 Pairwise alignment of the 6.8 kDa (62 amino acids) gp52 protein with other known small RDFs 410 revealed homology with $\phi Rv1 RDF$ (25.7% identity and 35.1% similarity; Fig. 6A). Also, examination of the mobile elements that have inserted into the attB sites in S. coelicolor and 411 other Streptomyces spp, revealed that they also contain a gene encoding a gp52 homologue 412 in a similar genetic context i.e. the int and g52 genes are adjacent to the attL and attR sites, 413

respectively, and would flank *attP* after excision (Fig. 4B & S2). The predicted secondary structure of \$\phi\$Joe gp52 contains an alpha-helix in the N-terminal region, a beta-sheet in the Cterminal region and an unstructured region in between (Fig. S6). Alignment of the \$\phi\$Joe-like RDFs found in intact phages and the RDFs found in the SCO2603-encoding mobile elements indicated that both of the structured regions are well conserved, particularly the putative alphahelix, but the centre of the protein is variable (Fig. S6).

420 RDFs are able to influence integrase-catalysed recombination in two ways; they activate the attL x attR reaction to regenerate attP and attB (excision) and they inhibit the attB 421 x attP integration reaction (22, 23). We were unable to produce sufficient soluble gp52 protein 422 for *in vitro* assays when expressed with a simple histidine-tag; however, a maltose-binding 423 protein MBP-gp52 fusion protein was more soluble. We tested the ability of MBP-gp52 to 424 inhibit integration by titrating the protein against a fixed concentration of integrase at MBP-425 gp52:Int ratios of 1:2 to 22.5:1. When the MBP-gp52 was in excess integration was repressed 426 to less than 10%; however, at less than equimolar concentrations, recombination was 427 428 equivalent to the control in which no MBP-gp52 was added (Fig. 6B). These results are similar 429 to observations for ϕ C31 and Bxb1 integrases and their cognate RDFs, gp3 and gp47 (22, 23). 430

431 To test the ability of gp52 to activate an excision reaction, a plasmid containing the cognate *attLsv* and *attRsv* sites was produced, pCMF98 (Fig. S3). The MBP-gp52 protein was 432 unable to promote efficient excision under any conditions tested (not shown). Removal of the 433 MBP-tag, using 3c protease, increased excision activity but the reaction was still inefficient 434 after 2 h incubation (Fig. 6C). Longer incubations of 5 – 20 h further increased the amount of 435 substrates converted to product up to 45%, but also led to significant amounts of excision 436 products (10-20%) by the integrase alone. Thus, in comparison to the activity of other RDFs, 437 gp52 has rather poor activity; ϕ C31 gp3 activates approximately 60 to 80% conversion of the 438 439 attL x attR substrates to products (22) and similar results are obtained with other RDFs (23, 25, 26). 440

441 To test the excision ability of ϕ Joe gp52 *in vivo*, a *g52* and *int* co-expression operon was designed in which int and g52 were located directly downstream of the T7 promoter and 442 ribosome binding site (RBS) in the expression vector pEHISTEV to produce pCMF117. A 443 reporter plasmid, pCMF103, was produced containing the $lacZ\alpha$ gene flanked by ϕ Joe attLsv 444 445 and *attRsv* sites (Fig. S3). pCMF117 and pCMF103 were introduced into *E. coli* BL21(DE3) 446 cells by co-transformation and plated onto LB agar supplemented with 0.5 mM IPTG to induce 447 expression of the g52-int operon (30). The reporter plasmid was then extracted from the 448 BL21(DE3) transformants and introduced into *E. coli* DH5a to determine the percentage of plasmids that had undergone attLsv x attRsv recombination and had lost the $lacZ\alpha$ gene. As 449 controls, plasmids expressing either only integrase (pCMF87) or only gp52 (pCMF100) were 450 also introduced together with the reporter (pCMF103) into BL21(DE3) and the assay was 451 452 repeated using the same procedure. When \$\phiJoe\$ integrase alone was expressed, excision occurred at a frequency of 37.6% (SD=5.1%, n=5) but when co-expressed with gp52 the 453 454 frequency rose to 96.8% (SD=1.3%, n=5) (Fig. 6D). Expression of gp52 without integrase led to no detectable excision events (Fig. 6D). Although overall recombination in vivo was higher 455 than *in vitro*, the relative levels of *attLsv* x *attRsv* recombination by ϕ Joe integrase alone and 456 \$\phi Joe integrase with gp52 were comparable. Taken together, the *in vivo* and *in vitro* data 457 indicate that ϕ Joe gp52 has RDF activity. 458

459 The observation that ϕ Joe integrase has a basal level of excision activity in the absence of its RDF is highly unusual for a phage-encoded integrase and further study may provide novel 460 461 insights into the mechanism and evolution of the serine integrases. Streptomyces phage ϕ BT1 462 integrase was shown to catalyse bidirectional recombination, albeit at extremely low levels 463 (59). The archetypal ϕ C31 integrase is only able to mediate *attL* x *attR* recombination in the 464 absence of gp3 when certain mutations are introduced just upstream or within a motif, the 465 coiled coil motif, required for subunit-subunit interactions during synapsis of DNA substrates 466 (60). The coiled coil motifs are also thought to play a role in inhibiting recombination between 467 attL and attR in the absence of the RDF; the ϕ C31 IntE449K mutation or its RDF, gp3, relieves

468 this inhibition (55, 60–62). Three independent structural predictions indicate the presence of a coiled coil domain in the \$\phi_Joe Int C-terminal domain (A395-T453, Fig. S7). The high basal 469 excision activity of ϕ Joe integrase could be due to incomplete inhibition of synapsis by the 470 coiled coil motif when integrase is bound to attL and attR, reminiscent of the hyperactive ϕ C31 471 472 mutant IntE449K (60). Natural bidirectional, large serine recombinases include the transposases TnpX (63) and TndX (64) from clostridial integrated conjugative elements 473 474 475 recombinases and the highly directional recombinases such as ϕ C31 and Bxb1 integrases. 476 Our data show that, under the *in vitro* conditions used, gp52 was highly effective at inhibiting 477 integration by ϕ Joe integrase but only weakly activated excision. It remains to be seen whether 478 this system, with its unusual properties, is sufficiently robust to regulate phage genome integration and excision according to the developmental choices of ϕ Joe. 479

480 The properties of the ϕ Joe integrase and gp52 are compatible with some of the existing applications for serine integrases, but they could also present opportunities for new 481 applications. ϕ Joe integrase is highly efficient in integration assays in vivo and in vitro, and in 482 *vivo* excision when the RDF is present. In *attB* x *attP* integration assays, the yield of products 483 484 by ϕ Joe integrase was comparable to well established integrases such as those of ϕ C31 or 485 Bxb1. Furthermore, ϕ Joe integrase is active in buffers compatible with other characterized 486 integrases indicating that it could be used in DNA assembly procedures in combination with other integrases. Although yet to be tested, assemblies generated with \$\phi_Joe\$ integrase could 487 488 later be used as substrates for modification by ϕ Joe integrase in a single step. The innate excision activity of \$\$\phi_Joe integrase could excise a fragment flanked by attLsv or attRsv sites 489 490 and, in the same reaction, replace it via an integration reaction. ϕ Joe integrase could therefore provide a more streamlined tool than the existing requirement for two steps by the more 491 492 directional integrases such as those from ϕ C31 and Bxb1 (15). Furthermore, given that ϕ Joe Int can mediate basal levels of excision in the absence of RDF, integrating plasmids based on 493 494 \$\phi Joe int/attP may display a degree of instability. Selection for the plasmid marker would ensure

plasmid maintenance when desired but, if the plasmid is easily lost without selection, this trait
could be desirable if there is a need to cure the strain of the plasmid or during studies on
synthetic lethality.

498 **Conclusions.** On the basis of sequence and genome organisation, phage Joe is a member of a large cluster of R4-like *Streptomyces* phages. Its closest relatives at the nucleotide level 499 are *Streptomyces* phages Amila and Verse with very high levels of nucleotide identity in the 500 501 regions encoding essential early and structural genes. However, Joe integrase is more closely related to the integrases from five other R4-like cluster phages - Lannister, Danzina, Zemlya, 502 Lika and Sujidade. At the present time the majority of Streptomyces phages belong to the R4-503 like cluster phages, but there is a continuum of relatedness throughout the cluster; for example 504 R4 is a more distant relative to ϕ Joe than any of the other phages mentioned above. 505

506 We identified the RDF for Joe integrase on the basis of its gene location, small size and distant similarity to another known RDF, Rv1584c. Although this identification was relatively 507 508 straightforward, it is not clear yet how general such an approach might be. The activity of ϕ Joe 509 integrase and RDF contributes to the growing number of complete serine integrase site-510 specific recombination systems that are available for use in synthetic biology applications. The ♦Joe int/attP plasmid, pCMF92, also adds to the number of useful integrating vectors for use 511 in *Streptomyces* species. However, and unusually for a phage integrase, ϕ Joe Int displays a 512 significant level of excisive recombination in the absence of its RDF while still being efficient 513 at mediating integration. This bi-directional property could be applied in new ways in future 514 applications of serine integrases. 515

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extracytoplasmic function sigma factor σ^E is required for normal cell wall structure in
Streptomyces coelicolor A3(2). J Bacteriol 181:204–211.
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- 725 726 Figures
- 727 Figure 1: A φJoe virion imaged by transmission electron microscopy.



Figure 2: Schematic of the φJoe genome.



Figure 3: Circos plot of the φJoe genome versus nine related phages.





Figure 4: φJoe attachment sites and integration sites.

Figure 5: Activity of ϕ Joe integrase in vivo and in vitro.





96.8% (SD=1.3%, n=5)

742 Figure 6: Identification of the φJoe RDF, gp52.

37.6% (SD=5.1%, n=5)

0.0% (SD=0.0%, n=5)

745 Figure Legends.

Figure 1: A \$\phi\$Joe virion imaged by transmission electron microscopy. Viral particles were
 negatively stained with uranyl acetate and this image was taken at 220,000x magnification.
 The scale bar represents 100 nm.

Figure 2: Schematic of the ϕ Joe genome. The genome is 48,941 bp in length. ORFs were 749 predicted using GeneMark and Glimmer then manually curated. ORFs are labelled and colour-750 coded based on their predicted function. Orange = recombination; cyan = metabolism and 751 DNA processing/replication; green = structural proteins; purple = lysis; black = regulatory; grey 752 = hypothetical proteins with no known function; red = candidate RDF genes. Genes marked 753 754 with an asterisk encode structural proteins that were detected by tandem MS:MS. The histogram below the genome contains purple bars to indicate below-average GC content 755 756 (65.5%) and green bars to indicate above-average GC content (1000 nt window size, 20 nt step). 757

Figure 3: Circos plot of the ϕ Joe genome versus nine related phages. A Blastn comparison was carried out for ϕ Joe, the five sequenced phage with a ϕ Joe-like integrase, the three closest whole genome matches and the well-characterized R4 phage. The E-value cut-off was set to 1×10^{-100} and the HSPs to 100, ribbons are coloured by genomic regions as defined in Figure 1 and depicted above the Circos plot. The histograms above each genome are coloured to reflect relative homology to the ϕ Joe sequence based on Blast score (Red>Orange>Green>Blue).

Figure 4: **\$\phi Joe** attachment sites and integration sites. **A.** Diagram of \$\phi Joe attP showing 765 766 the central dinucleotides (Purple) and imperfect inverted repeats (Orange and arrows). B. 767 Schematic of the genomic context of the two S. coelicolor integration sites (attLsc and attRsc, red boxes) used by the \$\phiJoe integrating plasmid pCMF92. The location of the PstI sites used 768 for identification of the att sites are shown. The DNA between the attLsc and attRsc sites is 769 an apparent mobile genetic element with homologous integrase and RDF genes (orange 770 771 arrows) to those of ϕ Joe. C. Alignment of S. venezuelae attB (attBsv) with the two S. coelicolor att sites (attRsc and attLsc) and the reconstituted attB site (attBsc) that would be produced by 772 excision of the DNA between *attRsc* and *attLsc*. **D.** Alignment of closely related *attB* sites 773 774 identified by a Blastn search against the non-redundant Genbank database. Hits were first filtered for matches of at least 80% and then for an e-value of $<1 \times 10^{10}$ and a bit score >75. 775 Nucleotide positions in C and D are shown as distance from the crossover dinucleotides (XX). 776

Figure 5: Activity of ϕ Joe integrase *in vivo* and *in vitro*. A. Conjugation efficiency of an 777 integrating vector, containing \$\phi Joe int and attP, into five recipient species - Streptomyces 778 coelicolor (Sc), S. lividans (SI), S. venezuelae (Sv), S. albus (Sal) and S. avermitilis (Sav). 779 Levels of significance for S. venezuelae versus all other species in a one-way ANOVA was p 780 = <0.001 (3 asterisks), all other comparisons were non-significant (n.s.). Error bars are 781 standard deviation (Sc n=5, Sv and SI n=3, Sal and Sav n=2). B. Representative image of an 782 *in vivo* integration assay to assess *attBsv/attP* recombination by ϕ Joe integrase (pCMF107) 783 and a negative control (pBAD-HisA). Recombination leads to deletion of an intervening *lacZa* 784 785 gene and white colonies, inactivity produces blue colonies. Integration efficiency is shown in brackets (n=3). C. Representative image of *in vitro* recombination of two substrate plasmids, 786 attP (pCMF91) and attBsv (pCMF95), to produce the co-integrant plasmid pCMF98. The 787

concentration of \$\overline\$Joe Integrase and incubation time for each reaction is indicated above the
 gel. **D.** Time-course for the integration reaction shown in part C.

Figure 6: Identification of the ϕ Joe RDF, gp52. A. Alignment of ϕ Joe and RV1 RDFs, 790 coloured using the BLOSUM62 scheme. B. Representative agarose gel showing in vitro 791 792 inhibition of integration by ϕ Joe RDF. The concentration of ϕ Joe Integrase and RDF for each reaction is indicated above the image. Reactions were stopped after 2 h and linearized using 793 Xhol. C. Representative agarose gel showing *in vitro* excision reactions catalysed by ϕ Joe 794 795 Integrase and RDF. The concentration of ϕ Joe Integrase and RDF for each reaction is indicated above the image. Reactions were stopped after 2 h and linearized using Xhol. D. In 796 vivo excision assay to assess attLsv x attRsv recombination by ϕ Joe integrase alone, ϕ Joe 797 RDF alone and ϕ Joe integrase co-expressed with the RDF. Recombination leads to deletion 798 799 of an intervening $lacZ\alpha$ gene and white colonies, inactivity produces blue colonies. Expression 800 from the T7 promoter successfully achieved almost complete excision activity for \$Joe Int + 801 RDF.

802 Table 1. Plasmids used in this study

Plasmid	Description	Resistance	Reference
pSET152	φC31 <i>int</i> + <i>attP</i> integrating vector	Apra	(65)
pEHISTEV	Expression vector, T7 promoter, C-terminal HIS6, TEV cleavage site	Kan	(66)
pETFPP_2	Expression vector; HIS6-MBP-3c Cleavage Site	Kan	(67)
pBAD-HisA	Expression vector, araBAD inducible promoter	Amp	Invitrogen
pCMF87	pEHISTEV + ¢Joe <i>int</i> (gp53)	Kan	This Study
pCMF90	pGEM7 + <i>S. coelicolor attRsc</i> (274 bp)	Amp	This Study
pCMF91	pSP72 +	Amp	This Study
pCMF92	φJoe int + <i>attP</i> integrating vector; pSET152	Apra	This Study
pCMF94	pGEM7 + <i>S. coelicolor attLsc</i> (419 bp)	Amp	This Study
pCMF95	pGEM7 + <i>S. venezuelae attBsv</i> (462 bp)	Amp	This Study
pCMF96	pETFPP_2 +	Kan	This Study
pCMF97	pGEM7 + S. coelicolor reconstituted attBsc (152 bp)	Amp	This Study
pCMF98	øJoe attLsv/attRsv; pCMF91 integrated into pCMF95	Amp	This Study
pCMF100	pEHISTEV + ¢Joe RDF	Kan	This Study
pCMF103	pACYC184 + φJoe <i>attLsv-lacZα-attRsv</i>	Cm	This Study
pCMF107	pBAD +	Amp	This Study
pCMF108	pBAD +	Amp	This Study
pCMF116	pACYC184 + φJoe <i>attBsv-lacZα-attP</i>	Cm	This Study
pCMF117	pEHISTEV +	Kan	This Study
pGEM7	General cloning vector	Amp	Promega
pSP72	General cloning vector; Accession X65332	Amp	Promega
pACYC184	General cloning vector; Accession X06403	Cm	(68)
pUZ8002	Conjugation helper plasmid; RK2 derivative with defective oriT	Kan	(69)

805 Table 2. Primers used in this study

Primer	Sequence (5' – 3')
Joe Int-attP F	CCGTCGACCTGCAGGCATGCCGTTCCCGCAGGTCAGAGC
Joe Int- <i>attP</i> R	ACATGATTACGAATTCTGTGGATCAGAACGTCTCGG
Joe H6-Int F	TTTCAGGGCGCCATGATGAGTAACCGACTACATG
Joe H6-Int R	CCGATATCAGCCATGTCAGAACGTCTCGGCGAAG
Joe <i>attP</i> F	TACCGAGCTCGAATTAAGACCGTCTCAGCCAGG
Joe <i>attP</i> R	TATCATCGATGAATTTCAGTGAAGACGGACAGG
Joe <i>attB1</i> F	CCGGGGTACCGAATTTGTGACGTCAGCCACAGC
Joe <i>attB1</i> R	TAGACTCGAGGAATTGACAAGGAGTGGCTCTGG
Joe <i>attB2</i> F	CCGGGGTACCGAATTGACTGCGTGCCGTCAGCC
Joe <i>attB2</i> R	TAGACTCGAGGAATTCGTCGTGTCGTCTGTCAG
Joe attB Sv F	CCGGGGTACCGAATTACCAGGTGGTGGATGAGC
Joe attB Recon F	TAGACTCGAGGAATTACCTTGATCTCGGTGTCCATCGCCGGGCAGACG
	CCGCAGTCGAAGCACGG
Joe <i>attB</i> Recon R	CCGGGGTACCGAATTGACAAGGAGTGGCTCTGG
Joe MBP-gp52 F	TCCAGGGACCAGCAATGAACGGACAGATCCTGG
Joe MBP-gp52 R	TGAGGAGAAGGCGCGCTACACCCAGCGCACCGA
CleF	CGCGCCTTCTCCTCACATATGGCTAGC
CleR	TTGCTGGTCCCTGGAACAGAACTTCC
Joe H6-gp52 F	TTTCAGGGCGCCATGAACGGACAGATCCTGGAG
Joe H6-gp52 R	CCGATATCAGCCATGCTACACCCAGCGCACCGA
Joe pBAD Int F	GAGGAATTAACCATGAGTAACCGACTACATG
Joe pBAD Int R	TGAGAACCCCCCATGTCAGAACGTCTCGGCGAAG
Joe pBAD gp52 F	GAGGAATTAACCATGAACGGACAGATCCTGGAG
Joe pBAD Int Co-Ex F	AGTGGTAGGTTCCTCGCCATG
Joe pBAD gp52 R	GAGGAACCTACCACTCTACACCCAGCGCACCGA
Joe LzR F	GGGTGTCAGTGAAGTAGTTGTGGCCATGTGTCCATCTGGGGGCAGACG
	CCGCAGTCGAAGCACGGCGATTTCGGCCTATTGGT
Joe LzR R	CCTGCCACATGAAGCGGATGTGACCCCGTCTCCATCTGCCCGCAGATG
	GACACCCACATCCAGATAATACGCAAACCGCCTCT
Joe BzP F	GGGTGTCAGTGAAGTATCTGGATGTGGGTGTCCATCTGCGGGCAGACG
	CCGCAGTCGAAGCACGGCGATTTCGGCCTATTGGT
Joe BzP R	CCTGCCACATGAAGCGGATGTGACCCCGTCTCCATCTGCCCCCAGATG
	GACACATGGCCACAACTAATACGCAAACCGCCTCT
SPBc H6-sprA F	CCGATATCAGCCATGGAGTTAAAAAACATTGTT
SPBc H6-sprA R	TTTCAGGGCGCCATGCTTACTACTTTTCTTAGTGG
SPBc MBP-sprB F	TCCAGGGACCAGCAATGGAACCTTACCAACGT
SPBc MBP-sprB R	TGAGGAGAAGGCGCGAAGCTTACTCTGCCTTCC
SPBc LZR F	GGGTGTCAGTGAAGTAGTGCAGCATGTCATTAATATCAGTACAGATAAA
	GCTGTATATTAAGATACTTACTACATATCTACGATTTCGGCCTATTGGT
SPBc LZR R	CCTGCCACATGAAGCTGGCACCCATTGTGTTCACAGGAGATACAGCTTT
	ATCTGTTTTTTAAGATACTTACTACTTTTCTAATACGCAAACCGCCTCT

Table 2. φJoe Host Range.

Host Species	Lysis (pfu ⁸⁰⁹
Streptomyces albus J1074	(2x10 ⁹)
Streptomyces avermitilis	(2x10 ⁹) ⁸¹⁰
Streptomyces coelicolor J1929	(2x10 ⁸)
Streptomyces coelicolor M145	√ 811
Streptomyces griseus	(4x10 ⁸) ₀₁₂
Streptomyces lividans TK24	(7x10 ⁷) ⁰¹²
Streptomyces nobilis	(1x10 ⁴) ₉₁₂
Streptomyces scabies	(6x10 ⁷)
Streptomyces venezuelae	X 814
Streptomyces venezuelae VL7	X
Streptomyces venezuelae VS1	X 815
Streptomyces venezuelae 10712	X 015
Saccharopolyspora erythraea	X 816
Saccharopolyspora erythraea	X 816

- **^ Pfu/ml values quoted are illustrative of the relative plaquing efficiencies when**
- challenged with the same phage stock propagated on *S. coelicolor* J1929