Plasmid Conjugation from Proteobacteria Journal of as Evidence for the Origin of Xenologous Genes in Cyanobacteria Bacteriology David Encinas, M. Pilar Garcillán-Barcia, María Santos-Merino, Luis Delaye, Andrés Moya and Fernando de la Cruz J. Bacteriol. 2014, 196(8):1551. DOI: 10.1128/JB.01464-13. Published Ahead of Print 7 February 2014. Updated information and services can be found at: http://jb.asm.org/content/196/8/1551 These include: SUPPLEMENTAL MATERIAL Supplemental material REFERENCES This article cites 60 articles, 25 of which can be accessed free at: http://jb.asm.org/content/196/8/1551#ref-list-1 CONTENT ALERTS Receive: RSS Feeds, eTOCs, free email alerts (when new articles cite this article), more»

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Plasmid Conjugation from Proteobacteria as Evidence for the Origin of Xenologous Genes in Cyanobacteria

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Comparative genomics have shown that 5% of *Synechococcus elongatus* PCC 7942 genes are of probable proteobacterial origin. To investigate the role of interphylum conjugation in cyanobacterial gene acquisition, we tested the ability of a set of prototype proteobacterial conjugative plasmids (RP4, pKM101, R388, R64, and F) to transfer DNA from *Escherichia coli* to *S. elongatus*. A series of BioBrick-compatible, mobilizable shuttle vectors was developed. These vectors were based on the putative origin of replication of the *Synechococcus* resident plasmid pANL. Not only broad-host-range plasmids, such as RP4 and R388, but also narrower-host-range plasmids, such as pKM101, all encoding MPF_T-type IV secretion systems, were able to transfer plasmid DNA from *E. coli* to *S. elongatus* by conjugation. Neither MPF_F nor MPF_I could be used as interphylum DNA delivery agents. Reciprocally, pANL-derived cointegrates could be introduced in *E. coli* by electroporation, where they conferred a functional phenotype. These results suggest the existence of potentially ample channels of gene flow between proteobacteria and cyanobacteria and point to MPF_T-based interphylum conjugation as a potential mechanism to explain the proteobacterial origin of a majority of *S. elongatus* xenologous genes.

orizontal gene transfer (HGT) is an outstanding player of bacterial evolution (1). Among classical HGT mechanisms, natural transformation was demonstrated in several cyanobacteria, including Synechococcus elongatus PCC 7942 (referred to here as Se7942) (2), Synechococcus sp. strain PCC 7002 (3), and Synechocystis sp. strain PCC 6803 (4), while conjugative transfer among Anabaena strains was also reported (5). Although no experimental evidence for transduction has been reported, several marine phages that contain photosynthetic genes have been detected (6, 7). This fact could indicate that photosynthetic genes are also mobilized by transduction. Comparative genomic analysis of the Se7942 and other cyanobacterial genomes identified xenologous genes based on the combination of multiple approaches: best BLAST hit out of cyanobacteria, absence of the ubiquitous octanucleotide HIP1 motif (for highly iterated palindrome 1) (8), differences in codon usage, GC index and trinucleotide skews (9). Based on these criteria, a majority of these genes (162 out of 253) probably originated from the phylum proteobacteria. These data suggest that functional mechanisms of HGT must exist, to provide a genetic bridge between phyla proteobacteria and cyanobacteria.

Conjugation has been used as a tool for introducing shuttle vectors from *Escherichia coli* to both pluricellular (several strains of heterocyst-forming *Anabaena* [10], non-heterocyst-forming *Leptolyngbya* sp. strain BL0902 [11], akinetes, hormogonia, and heterocyst-forming *Fischerella muscicola* PCC 7414, and *Chlorogloeopsis fritschii* PCC 6912 [12]) and unicellular (Se7942 [13, 14], several strains of marine *Synechococcus* [60], *Prochlorococcus* strain MIT9313 [15], and *Synechocystis* sp. strain PCC 6803 [13]) cyanobacteria. These shuttle vectors were either based on the mobilization of the promiscuous plasmid RSF1010 or in a ColE1-like origin of transfer (16). Although conjugative plasmids of several incompatibility groups were tested as helpers to mobilize these shuttle vectors, only IncP1-MOB_{P11} plasmids, such as RP4 and R751, were successful at transferring DNA from *E. coli* to cyanobacteria (17). Thus, up to now, mobilizable shuttle vectors have

relied on IncP1 helper plasmids to be transferred to cyanobacterial recipients by conjugation (14, 16).

With few exceptions, proteobacterial conjugative plasmids can be grouped into five MOB families (MOB_P, MOB_F, MOB_O, MOB_C, and MOB_H) and three mating-pair formation types $(MPF_T, MPF_F, and MPF_I)$ (18). Natural combinations MOB_{P11} - MPF_{T} (present in IncP1 α plasmid RP4), MOB_{F11} -MPF_T (present in IncN plasmid pKM101 and IncW plasmid R388), MOB_{P12}- MPF_{I} (present in IncI α plasmid R64), and MOB_{F12} -MPF_F (present in IncFI plasmid F) were tested here to investigate the range of proteobacterial conjugative systems able to conjugate DNA to cyanobacteria. These plasmids were used as helpers to mobilize a series of BioBrick-compatible shuttle vectors containing a cognate MOB from E. coli to Se7942. Such vectors were helpful to define the functional replicon of plasmid pANL. The conjugation results showed that all tested MPF_T plasmids, regardless of their MOB type, were proficient for delivering DNA to cyanobacteria by conjugation, suggesting that plasmid conjugation from proteobacteria has contributed to the composition and evolution of Se7942 genome.

MATERIALS AND METHODS

Strains and culture conditions. Strains used are detailed in Table 1. The original Se7942 strain we used was already cured of the endogenous plasmid pANS. Se7942 was cultured at 30°C in BG11 medium (19) by bub-

Received 20 December 2013 Accepted 1 February 2014 Published ahead of print 7 February 2014 Address correspondence to Fernando de la Cruz, delacruz@unican.es. D.E. and M.P.G.-B. contributed equally to this article. Supplemental material for this article may be found at http://dx.doi.org/10.1128 /JB.01464-13. Copyright © 2014, American Society for Microbiology. All Rights Reserved. doi:10.1128/JB.01464-13

TABLE 1 Bacterial strains used in this study

Strain	Description and/or relevant characteristics ^a	Source or reference ^b
β2150	Δ dapA::(erm-pir) thrB1004 pro thi strA hsdS lacZ Δ M15 (F' lacZ Δ M15 lacI ^q traD36 proA ⁺ proB ⁺); Em ^r Sm ^r	39
BW27783	$lacI^{q}$ rrnB3 $\Delta lacZ4787$ hsdR514 DE(araBAD)567 DE(rhaBAD)568 DE(araFGH) $\Phi(\Delta araEp P_{CP8}-araE)$; Nx ^r	52
DH5a	F^- endA1 glnV44 thi-1 recA1 relA1 gyrA96 deoR nupG Φ 80dlacZ Δ M15 Δ (lacZYA-argF)U169, hsdR17($r_k^- m_k^+$) λ^- ; Nx ^r	53
Synechococcus elongatus PCC 7942	Wild-type strain lacking plasmid pANS (NC_007604 plus NC_004073); also known as Anacystis nidulans R2-SPc; classified into the cyanobacterial section I	54; PCC
GRPS1	S. elongatus PCC 7942 with a mutation in rps12-R43; Sm ^r	55
Leptolyngbya sp. strain PCC 7410	Wild-type strain; classified into the cyanobacterial section III	PCC
Plectonema boryanum	Wild-type strain; classified into the cyanobacterial section III	PCC
Anabaena variabilis ATCC 29413	Wild-type strain (NC_007413 plus NC_007410 plus NC_007411 plus NC_007412); classified into the cyanobacterial section IV	ATCC
Nostoc punctiforme PCC 73102	Wild-type strain (NC_010628 plus NC_010631 plus NC_010632 + NC_010630 plus NC_010633 plus NC_010629); classified into cyanobacterial section IV	PCC
Nostoc punctiforme ATCC 29133	Wild-type strain; classified into the cyanobacterial section IV	ATCC

^{*a*} Em^r, erythromycin resistance; Nx^r, nalidixic acid resistance; Sm^r, streptomycin resistance.

^b PCC, Pasteur Culture Collection; ATCC, American Type Culture Collection.

bling 1% CO₂ with continuous light at 60 µmol of photons m⁻² s⁻¹. *Leptolyngbya* PCC 7410, *Anabaena variabilis* ATCC 29413, *Plectonema boryanum*, *Nostoc punctiforme* PCC 73102, and *Nostoc punctiforme* ATCC 29133 were cultured in BG11 at 25°C using 20 µmol of photons m⁻² s⁻¹, 10 rpm, and atmospheric CO₂ conditions. The *E. coli* strains used were BW27783, DH5 α , and β 2150. They were grown at 37°C under shaking in Luria-Bertani medium (LB). Strain β 2150 was supplemented with 30 µM diaminopimelic acid (DAP30). The antibiotics used for selecting cyanobacteria were neomycin at 5 or 25 µg/ml (Neo5 or Neo25), chloramphenicol at 5 or 10 µg/ml (Cm5 or Cm10), and streptomycin at 10 or 50 µg/ml (Sm10 or Sm50). The antibiotics used for selecting *E. coli* were kanamycin at 50 µg/ml (Km50), rifampin at 50 µg/ml (Nx20), and streptomycin at 300 µg/ml (Sm300).

Construction of vectors. The plasmids and oligonucleotides are given in Table 2 and Table S1 in the supplemental material, respectively. The steps for the construction of the shuttle vectors are depicted in Fig. S1 in the supplemental material, while dislodging vectors are described in Fig. S2 and S3 in the supplemental material. Details on the construction procedures are summarized in the supplemental material.

Conjugation assays between E. coli and cyanobacteria and between E. coli. Biparental assays were used to conjugate DNA from E. coli to cyanobacteria. They were performed at 30°C for Se7942 and at 25°C for other cyanobacterial genera. For each conjugation, a Se7942 culture sample equivalent to 15 μ g of chlorophyll (around 6 \times 10⁸ cyanobacterial cells/µg of chlorophyll) was mixed with 100 µl of serial dilutions of a 109-cells/ml E. coli culture. Conjugative mixtures were placed on top of a nitrocellulose filter that was in turn placed on a BG11 plate supplemented with 5% LB plus DAP30 (LB-DAP30) for 1 h in the dark. The conjugative mixture was incubated for 24 h in the presence of light (60 µmol of photons m⁻² s⁻¹). Filters were later changed to fresh BG11 plates, incubated for an additional 24-h period, and finally transferred to BG11-Neo25 under the same conditions. Transconjugant colonies became visible after 7 to 14 days of incubation. Conjugation between E. coli cells was performed as previously described (20). Strains β 2150 or BW27783 were used as donors, while DH5a was used as a recipient strain. Conjugation frequencies were expressed as the number of transconjugants per donor cell and calculated as described previously (21).

Natural transformation. Se7942 was transformed with plasmid pDEP30 according to a protocol described earlier (22). Transformation

mixtures were deposited onto nitrocellulose filters (Millipore) and incubated in BG11 plates at 30°C with continuous light for 24 h. Transformants were selected in BG11-Cm10.

Dislodging assays and analysis of cyanobacterial transconjugant colonies. Dislodging vectors pDEP21 and pDEP23 were introduced in Se7942 by conjugation from *E. coli* using RP4 as helper plasmid. Individual transconjugant Se7942 colonies, carrying either pDEP21 or pDEP23, were grown in 250 ml of BG11-Neo5. Once cultures reached an optical density at 750 nm (OD₇₅₀) of 2, 1.0 ml was transferred to 250 ml of BG11-Neo5. Serial dilutions were repeated for 42 or 64 generations of growth (for pDEP23 or pDEP21, respectively), when the axenic condition of these cultures was confirmed. The presence or absence of pANL was checked by PCR using the primer pairs 31/32, 33/34, and/or 35/36 when appropriate.

Transconjugant colonies of Se7942 strain GRPS1 carrying pDEP23 were serially replicated in BG11-Neo5 plates to remove E. coli cells. Axenic cultures of GRPS1 carrying pDEP23 were transformed with pDEP30 plasmid DNA. The latter plasmid contains a Synechocystis sp. strain PCC 6803 gene *rps12* under the control of the *psbA1* gene promoter and a *cat* gene flanked by two 1-kb fragments located at both sides of the pANL maintenance region (see Materials and Methods and Fig. S3 in the supplemental material). Thus, a double crossover between pANL and pDEP30 should remove pANL maintenance region. Transformant colonies were grown in BG11-Cm5 plates to improve the segregation of the mutation, which was checked by PCR (30 cycles of 94°C for 60 s, 50°C for 30 s, and 72°C for 30 s) using the primer pairs 37/38 and 39/40 (see Fig. S4 in the supplemental material). The colonies were grown in liquid medium BG11-Neo5-Sm10 to favor displacement of the pANL derivative lacking the maintenance region (pDEP32) by pDEP23. Individual colonies recovered from BG11-Neo5-Sm50 plates were analyzed by PCR (30 cycles of 94°C for 60 s, 50°C for 30 s, and 72°C for 30 s) using the primer pairs 33/34 and 35/36 to check for the absence of the pANL derivative.

To analyze the cyanobacterial transconjugants, single colonies were streaked out twice in BG11-Neo5 plates to remove *E. coli* donors. Transconjugants were then grown in BG11-Neo5 up to an OD₇₅₀ of 1 to 1.5. The axenic condition of these cultures was tested in LB-DAP30. Plasmid DNA was isolated using GenElute TM Plasmid Miniprep kit (Sigma-Aldrich) and used to transform *E. coli* DH5 α by electroporation. Kanamycin-resistant (Km^r) *E. coli* transformants were analyzed by electrophoresis of plasmid DNAs with EcoRI and PstI in 1% agarose gel run in

 TABLE 2 Plasmids used in this study

Plasmid	Description ^a	Source or reference
pRL443	Km ^s RP4 derivative; Ap ^r Tc ^r	51
R388	Su ^r Tp ^r	56
R64drd11	Sm ^r Tc ^r	57
pKM101	Ap ^r	58
pOX38	F derivative; Cm ^r	59
pSB1K3	Rep(pMB8); Km ^r ; backbone for BioBrick parts cloning	http://parts.igem.org/Part:pSB1K3
pSB1C3	Rep(pMB8); Cm ^r ; backbone for BioBrick parts cloning	http://parts.igem.org/Part:pSB1C3
pEXR91	$Rep(pMB8); Ap^r Km^r;$ containing gene <i>rps12</i> under the promoter of the <i>psbAI</i> gene 25	
pDEP5	pSB1K3::(<i>rep</i> _{pANL}); Km ^r	This study; see also Fig. S1
pDEP6	pDEP11:: (rep_{pANL}) ; Km ^r	This study; see also Fig. 2 and S1
pDEP7	$pDEP12::(rep_{pANL}); Km^{r}$	This study; see also Fig. 2 and S1
pDEP8	pDEP13:: (rep_{pANL}) ; Km ^r	This study; see also Fig. 2 and S1
pDEP9	pDEP14:: (rep_{pANL}) ; Km ^r	This study; see also Fig. 2 and S1
pDEP10	pDEP15::(<i>rep</i> _{pANL}); Km ^r	This study; see also Fig. 2 and S1
pDEP11	pSB1K3::($oriT_{RP4}$); Km ^r	This study; see also Fig. S1
pDEP12	$pSB1K3::(oriT_{pKM101}); Km^{r}$	This study; see also Fig. S1
pDEP13	$pSB1K3::(oriT_{R388}); Km^{r}$	This study; see also Fig. S1
pDEP14	pSB1K3::(<i>oriT</i> _{R64}); Km ^r	This study; see also Fig. S1
pDEP15	pSB1K3::(<i>oriT</i> _F); Km ^r	This study; see also Fig. S1
pDEP16	pSB1K3::(sepA1); Km ^r	This study; see also Fig. S2
pDEP17	pSB1K3::(<i>sepA2</i>); Km ^r	This study; see also Fig. S2
pDEP18	pSB1K3::(Ptac); Km ^r	This study; see also Fig. S2
pDEP19	pDEP17::(<i>sepA1</i>); Km ^r	This study; see also Fig. S2
pDEP20	pDEP19::(Ptac); Km ^r	This study; see also Fig. S2
pDEP21	pDEP6::(Ptac-sepA1-sepA2); Km ^r	This study; see also Fig. 1 and S2
pDEP22	pSB1K3::(<i>gap2-3</i>); Km ^r	This study; see also Fig. S2
pDEP23	pDEP21::(<i>gap2-3</i>); Km ^r	This study; see also Fig. S2
pDEP24	pSB1K3::(HS1); Km ^r	This study; see also Fig. S3
pDEP25	pSB1K3::(HS2); Km ^r	This study; see also Fig. S3
pDEP26	pSB1K3::(<i>cat</i>); Km ^r Cm ^r	This study; see also Fig. S3
pDEP27	pSB1K3::(<i>rsp12</i>); Km ^r	This study; see also Fig. S3
pDEP28	pDEP27::(<i>cat</i>); Km ^r Cm ^r	This study; see also Fig. S3
pDEP29	pDEP25::(<i>cat-rsp12</i>); Km ^r Cm ^r	This study; see also Fig. S3
pDEP30	pDEP29::(HS1); Km ^r Cm ^r	This study; see also Fig. S3
pDEP31	pDEP6-pANL cointegrate; Km ^r	This study; see also Fig. 3
pDEP32	pANL in which the maintenance region comprised between HS1 and HS2 has been replaced by <i>rps12-cat</i> ; Km ^r Cm ^r	This study; see also Fig. S4

^{*a*} Cm^r, chloramphenicol resistance; Tc^r, tetracycline resistance; Ap^r, ampicillin resistance; Sm^r, streptomycin resistance; Km^r, kanamycin resistance; Tp^r, trimethoprim resistance; Su^r, sulfonamide resistance.

 $0.5 \times$ TBE buffer (44.5 mM Tris borate, 44.5 mM boric acid, 1 mM EDTA [pH 8.2 to 8.4]). Gels were stained with Real-Safe (Real) and developed in a Gel Doc Imager (Bio-Rad).

Chromate resistance test. *E. coli* DH5 α was independently transformed with cointegrate pDEP31 or vector pDEP6 by electroporation. Saturated cultures from single transformant colonies, grown in LB-Km50, were used to inoculate 96-well plates containing 150 µl of LB-Km50 per well and different concentrations of K₂CrO₄. Plates were incubated at 37°C. Bacterial growth was monitored based on the OD₆₀₀ in a Victor3 plate reader (Perkin-Elmer). Generation times were calculated as ln(2)/*k*, where *k* represents the growth rate and corresponds to the slope of the exponential growth phase (three experiments, eight replicas per experiment).

RESULTS

Dislodging vectors to displace the indigenous Se7942 plasmid pANL. The 46.3-kb pANL plasmid, indigenous to Se7942, could potentially interfere with the conjugation or stability of other plasmids, hence our interest in attempting pANL plasmid curing. Several shuttle plasmids were built, all based on a 1,395-bp DNA segment containing pANL putative origin of replication (23). The replication region (here named *rep_pANL*) that was cloned to construct the pDEP vector series included an additional 18-bp fragment to complete the coding region of gene *anL57* (Fig. 1A).

pDEP21 was built as a dislodging vector containing the pANL replication region in order to remove plasmid pANL from Se7942 by vectorial plasmid incompatibility under selective pressure (24) (Fig. 1B and see Fig. S2 in the supplemental material). Since plasmid pANL encodes two toxin-antitoxin systems, both pANL antitoxin genes (sepA1 and sepA2) were also included in pDEP21 to avoid killing pANL segregants. The dislodging plasmid pDEP21 also contains oriT_RP4 to allow conjugation from E. coli to cyanobacteria. Transconjugant colonies were subcultured in liquid BG11-Neo5 for 64 generations. Ten individual colonies were analyzed by PCR using primers that specifically hybridized to pANL and not to pDEP21. They all rendered amplicons congruent with the presence of pANL. In addition, plasmid DNA isolated from four independent colonies, transformed into E. coli and subjected to restriction analysis, showed no differences in restriction pattern between the recovered plasmid DNAs and the original dislodging



FIG 1 pDEP21 autoreplicates in Se7942. (A) Genetic organization of the putative replication region of pANL. The replication sequence described previously (23) is indicated by a white box outlined in black, while the segment used in the present study as *rep_*pANL is indicated by a gray box outlined in gray. (B) Genetic map of the dislodging vector pDEP21. (C) Plasmid DNA recovered from Se7942(pDEP21) cultured in liquid BG11-Neo5 for 64 generations was restricted with EcoRI (E) and/or PstI (P). Plasmid DNA directly extracted from *E. coli* DH5 α (pDEP21) without any passage through Se7942 was used as a control in the restriction analysis.

vector pEDP21 (Fig. 1C), indicating that it could be autonomously maintained in Se7942 but was not able to completely displace pANL.

In another attempt to remove pANL, regions named *gap2* and *gap3*, described as essential in an earlier study (23), were incorporated into the dislodging vector, thus producing pDEP23 (see Fig. S2 in the supplemental material). This vector was introduced in Se7942 by conjugation. Transconjugant colonies were cultured for 42 generations in BG11-Neo5 and plated out again, and plasmid DNA was extracted from single colonies. Restriction analysis showed a cointegrate between plasmids pDEP23 and pANL. No instance of pANL curing was detected.

A third attempt to remove pANL was carried out by deleting its maintenance region, previously described to be essential for stable carriage of pANL in Se7942 (23). This region is composed of two toxin-antitoxin system cassettes (sepA1-sepT1 and sepA2-sepT2), a set of partition genes (parA and parB), and an orf encoding a putative nucleotidyltransferase (anL30). Deletion of this pANL segment (coordinates 20984 to 24487 under GenBank accession no. AF441790) was carried out by homologous recombination in the Se7942 strain GRPS1, which is used for the construction of gene-replacement mutants (25). Plasmid pDEP30 (see Fig. S3 in the supplemental material) was introduced into strain GRPS1(pDEP23). The flanking areas of the pANL maintenance region surround genes rps12 and cat in pDEP30, which confer a dominant streptomycin-sensitive (Sm^s), chloramphenicol-resistant (Cm^r) phenotype. Deletion of the maintenance region was confirmed by PCR (see Fig. S4 in the supplemental material), and the deleted pANL derivative was named pDEP32. To favor pDEP23 in the competition with pDEP32, GRPS1 was grown in Neo25-Sm50. No GRPS1 colonies free of pDEP32 were

detected. All Sm^r neomycin-resistant (Neo^r) colonies tested contained a cointegrate between pDEP23 and pDEP32 (data not shown). In conclusion, all attempts at curing pANL failed, suggesting that this plasmid contains some genes that are essential for Se7942 viability under the tested conditions.

Mobilization of shuttle vectors from *E. coli* **to Se7942 using different prototype proteobacterial conjugative plasmids.** To test for the ability of different MPF systems to transfer plasmid DNA by conjugation to cyanobacteria, we tested the mobilization of a series of plasmid *oriTs* by their cognate conjugative plasmids from *E. coli* to Se7942.

Five prototype conjugative plasmids were tested (RP4, pKM101, R388, R64 and F) for transfer from E. coli to Se7942 (Fig. 2). They represent five frequently found incompatibility groups (Inc) in gammaproteobacteria, including four MOB subfamilies and three MPF types, thus comprising a representation of the diversity of proteobacterial mobility systems (18). First, mobilization of the shuttle vectors was tested by E. coli intraspecies crosses. As shown in Table 3 (column 3), all shuttle vectors were mobilized by their cognate conjugative plasmid between E. coli strains at frequencies of roughly 10⁻¹ transconjugants/donor cell, indicating that the helper plasmids were efficient in promoting mobilization of their cognate mobilizable vectors. When mobilization of the same shuttle vectors was tested, but using Se7942 as a recipient (Table 3, column 4), vectors containing the oriTs of RP4, pKM101, and R388 produced cyanobacterial Neo^r transconjugants. RP4, pKM101, and R388 transconjugants were obtained at frequencies of $\sim 10^{-3}$ in the mobilization of their cognate pDEP vectors. On the other hand, plasmids R64 and F showed conjugation frequencies undistinguishable from background levels. The temperature conditions used for Se7942 growth and mating were



FIG 2 Shuttle vector series. Mobilizable vectors are based on plasmid pSB1K3. All contain the *rep* region of plasmid pANL. Each one includes the origin of transfer (*oriT*) from a different prototype conjugative plasmid for which the Inc group, relaxase MOB family, and MPF type are indicated.

critical, since pDEP6 mobilization drastically dropped to background levels when the mating temperature was shifted from 30°C to 35, 37, or 40°C (data not shown). To rule out natural transformation as the cause of Se7942 Neo^r colonies, a control assay was carried out by repeating the conjugation experiment, using as the donor an *E. coli* strain containing vector pDEP5. Since pDEP5 is devoid of *oriT*, transfer of its Neo^r marker can occur only by natural transformation. The results shown in Table 3 (column 5) indicate that natural transformation was extremely inefficient under these conditions, occurring at frequencies not exceeding 10^{-8} . This in turn suggests that the number of transconjugants obtained using pDEP9 and pDEP10 as shuttle vectors (respectively mobilized by R64 and F) was within the range of natural transformation efficiency. Finally, experiments to mobilize pDEP6 (the vector containing *oriT_*RP4) to other cyanobacterial genera using RP4 as a helper plasmid were carried out using a diversity of cyanobacterial recipients: *Leptolyngbya* PCC 7410, *Anabaena variabilis* ATCC 29413, *Plectonema boryanum, Nostoc punctiforme* PCC 7310, and *Nostoc punctiforme* ATCC 29133. No transconjugants were obtained in any of these cases (data not shown). Since RP4 could mobilize RSF1010 to different cyanobacteria (16), it can be assumed that plasmid pANL (or its derivatives) cannot replicate in those cyanobacteria.

Analysis of pEDP6 in Se7942 transconjugants. To test whether pDEP6 could be maintained as an autonomous replicon in Se7942, plasmid DNA was extracted from four transconjugant colonies, transformed to *E. coli* to amplify the amount of plasmid

TABLE 3 Conjugative frequencies between E. coli and from E. coli to Se7942

Plasmid contained		Mobilization frequency ^c :			
in donor strain ^{<i>a</i>}	Inc MOB MPF type ^b	Between <i>E. coli</i> strains	From E. coli to Se7942	Of $\Delta oriT$ derivatives ^d	
$RP4+pDEP6^{e}$ $pKM101+pDEP7$ $R388+pDEP8$ $R64drd11+pDEP9$ $F+pDEP10^{f}$	$\begin{array}{l} IncP1\alpha \ MOB_{P11} \ MPF_T \\ IncN \ MOB_{F11} \ MPF_T \\ IncW \ MOB_{F11} \ MPF_T \\ IncI1\alpha \ MOB_{P12} \ MPF_I \\ IncFI \ MOB_{F12} \ MPF_F \end{array}$	$\begin{array}{l} 4.8 \times 10^{-1} \left(8.6 \times 10^{-2} \text{-}2.7 \right) \\ 5.2 \times 10^{-1} \left(3.1 \times 10^{-1} \text{-}8.4 \times 10^{-1} \right) \\ 5.0 \times 10^{-2} \left(1.9 \times 10^{-2} \text{-}1.3 \times 10^{-1} \right) \\ 8.5 \times 10^{-2} \left(6.1 \times 10^{-2} \text{-}1.2 \times 10^{-1} \right) \\ 4.8 \times 10^{-1} \left(3.8 \times 10^{-1} \text{-}5.9 \times 10^{-1} \right) \end{array}$	$\begin{array}{l} 2.2\times10^{-3}~(5.0\times10^{-4}1\times10^{-2})\\ 1.2\times10^{-3}~(4.5\times10^{-4}3.3\times10^{-3})\\ 8.9\times10^{-3}~(2.5\times10^{-3}3.1\times10^{-2})\\ <10^{-9}\\ <10^{-9}\end{array}$	$\begin{array}{c} 5\times10^{-9}(9\times10^{-10}{-}2\times10^{-8})\\ 2\times10^{-9}(6\times10^{-10}{-}1\times10^{-8})\\ 1\times10^{-8}(6\times10^{-9}{-}2\times10^{-8})\\ 4\times10^{-10}(8\times10^{-11}{-}2\times10^{-9})\\ 1\times10^{-9}(3\times10^{-10}{-}7\times10^{-9}) \end{array}$	

^{*a*} Donor strains were derivatives of *E. coli* strain β 2150 containing the plasmids shown in the first column. Strain BW27783 was used as the donor in the case of plasmid R388, since β 2150 was inhibitory to R388 conjugation, due to an inhibitory effect of the integrated F-plasmid (39; our unpublished results).

^b For a description of MOB and MPF types, see reference 18.

^c Mobilization frequencies are the average of at least six experiments. They were calculated by the log conversion of frequencies (number of transconjugants per donor cell) to obtain the means and standard deviations (indicated in parentheses), which are expressed as the anti-log of the calculated figures.

^d The same conjugation experiment from *E. coli* to Se7942 was conducted using the helper plasmids in combination with plasmid pDEP5, a nonmobilizable vector, to determine the background level of Neo^T Se7942 colonies that could arise in the mating experiments due to natural transformation. Transformation frequencies were calculated as the number of transforming cells/number of donor cells.

^e pRL443, an RP4 derivative sensitive to kanamycin, was used as a helper plasmid.

^f pOX38, a Cm^r F derivative, was used as a helper plasmid.



FIG 3 Analysis of transconjugant plasmid DNAs produced by pDEP6 conjugation to Se7942. The figure shows the electrophoretic analysis of DNA bands obtained by restriction analysis of transconjugant plasmid DNA. (A) Plasmid DNA recovered from Se7942(pDEP6) transconjugants (i.e., pEDP31) was digested with EcoRI (E) and/or PstI (P) and developed by 1% agarose gel electrophoresis. Plasmid pDEP6 DNA (isolated from *E. coli* DH5 α) was used as a control in the restriction analysis. (B) *In silico* restriction analysis of pDEP6-pANL cointegrate pDEP31 assuming a single crossover across the pANL *rep* region. (C) Schematic representation of the single crossover leading to pDEP31 formation, according to the results shown in panels A and B. The position of the single PstI site present in pANL replication region, which is not present in pDEP6, helps to determine the direction of the pDEP6 insertion.

DNA, and analyzed with restriction enzymes. All plasmid preparations recovered from Se7942 rendered the same restriction pattern. It was different from the original pDEP6 plasmid and consistent with it being a cointegrate (named pDEP31), formed by homologous recombination between pDEP6 and pANL *rep* regions, as shown in Fig. 3. Since pDEP31 contains the sulfate/chromate uptake operon of pANL (*srp* operon [23, 26]), it should result in increased chromate sensitivity of *E. coli* cells containing cointegrate pDEP31. This was proven by analysis of chromate sensitivity (see Fig. S5 in the supplemental material), which showed increased sensitivity to increasing amounts of chromate in DH5 α (pEDP31) with respect to the control strain of DH5 α (pDEP6). This result further indicates that the *srp* operon of Se7942 is adequately expressed in *E. coli*.

DISCUSSION

The main purposes of the present study were to determine which of the main conjugative systems from proteobacteria were able to transfer DNA to cyanobacteria by conjugation, to compare their relative efficiencies, and to optimize the conjugation protocol. The finding that all MPF_T -type plasmids were efficient donors is probably the most relevant result. It indicates that conjugation of proteobacterial plasmids is a probable source of xenologous genes in Se7942, as suggested previously (9).

For a start, and in order to optimize Se7942 as a conjugation recipient, we attempted to remove the indigenous pANL plasmid from Se7942 by using a plasmid incompatibility strategy. This technique is based in the fact that, when two plasmids carrying the same origin of replication coexist in a cell, they become unstable due to interactions between their replication machineries. Vectorial incompatibility (one of the plasmids is lost with higher probability than the other [24]) was previously exploited to cure native plasmids from Agrobacterium tumefaciens (27, 28), Bacillus anthracis (29), or Yersinia pestis (30), among many other examples. To cure pANL, two dislodging vectors were built: pDEP21 and pDEP23. They contain the proposed minimal replication region of plasmid pANL, defined as discussed in the supplemental material. When pDEP21 or pDEP23 were mobilized to Se7942 by RP4, pANL was not cured. Not even a pANL derivative lacking its maintenance region could be displaced by the dislodging vectors by applying selection to favor them in the competition. Thus, either the native pANL contains one or more essential genes that we did not include in our constructs, or the origin of replication described earlier (23) is incomplete, and some additional functions provided by pANL are indispensable for replication or maintenance of the dislodging vector. DNA extraction of transconjugant colonies showed that, in some cases, the pDEP derivative was still an autonomous replicon (Fig. 1), while in others it formed a cointegrate with pANL (Fig. 3). This result could occur if the incoming plasmid is unstable (by incompatibility) but cannot completely dislodge the resident plasmid, as discussed above. In our experiments, the restriction pattern of plasmid pDEP21 was not altered, indicating that it can remain as an autonomous plasmid in Se7942. On the other hand, plasmids pDEP6 and pDEP23, which contain the same rep_pANL as pDEP21, were always found forming cointegrates with pANL. Their different behavior remains unexplained. In any case, pANL could not be dislodged so all conjugation experiments were carried out in a Se7942 containing pANL.

Interphylum conjugation from E. coli to cyanobacteria is carried out in the laboratory solely by using an RP4-based helper plasmid (10, 16). Conjugation has been used as an alternative to transformation for the insertion of foreign genes in the Se7942 chromosome (22, 31). Mobilizable shuttle vectors were based either on plasmid pBR322 oriT (while MOB and MPF functions were provided in trans by a ColE1-like plasmid and RP4, respectively) (32), or on RSF1010, which is mobilized by RP4 (31). pBR322-based vectors were used to study whether plasmids other than RP4 supported mobilization to Anabaena strains M-131 or PCC 7120 (17). Other IncP1 plasmids could mobilize such vectors, while IncW plasmids could not. It should be pointed out that the transfer efficiency of a given mobilizable plasmid depends on the conjugative plasmid used as helper, because the relaxosome provided by the mobilizable plasmid should make appropriate contacts with the coupling protein and the mating apparatus provided by the conjugative plasmid. For example, RP4 mobilizes ColE1 and RSF1010 between E. coli strains 2 and 4 log more efficiently, respectively, than the IncW plasmid R388 (33). Thus, it is not surprising that IncW plasmids were unable to sustain conjugation to Anabaena in the referred conditions (17).

In the present study we used plasmids RP4, pKM101, R388, R64, and F, which represent the diversity of conjugative systems in proteobacteria (18), as potential donors in conjugation from proteobacteria to cyanobacteria. The relevant shuttle vectors contained always the same oriT as the conjugative plasmid used as helper, to avoid the above-mentioned inefficiencies due to heterologous interactions. This fact allows attention to be focused on the relative proficiency of each MPF type, not being distracted by the interactions between MPF and MOB modules. Actual transfer of a conjugative plasmid is an indication of two capabilities: first, its ability to invade the host population and, second, its efficiency of replicating in that host. Even in the absence of replication, the invasive plasmid DNA can recombine with the host genome (given the appropriate recombination sites), thus integrating within the chromosome and becoming an integrative and conjugative element (34). The host range and therefore the promiscuity of a plasmid can be inferred from sequence data (35). This type of analysis established that IncP1 plasmids (such as RP4) show a broad host range, in accordance with the fact that RP4 conjugates to cyanobacteria (10) and even to yeast (36). The same analysis indicated that IncW (R388) plasmids have a conjugative range perhaps broader than IncP, IncN (pKM101) plasmids have an intermediate range, and IncI (R64) or IncF (F) plasmids are narrow host range. Ample experimental evidence confirms these assumptions (37, 38).

Results shown in Table 3 indicate that all three conjugative plasmids with an MPF_T-type transport channel (i.e., RP4, pKM101, and R388) were able to achieve interphylum conjugation from *E. coli* to Se7942. The other two plasmids, F and R64, which contain MPF_F- or MPF_I-type conjugation systems (Fig. 2), could not introduce DNA in Se7942. These results suggest that different MPF types exhibit different abilities to conjugate to cyanobacteria. All transfer systems are able to transfer DNA to a much wider range of recipients than the replication ability of the vector plasmids (hence, suicide vectors used for gene delivery by conjugation [39, 40] or the induction of SOS response toward

invading DNA [41]). Thus, our results suggest that neither MPF_F nor MPF_I could deliver DNA to cyanobacteria, whereas all three MPF_T plasmids tested, irrespective of their MOB type, could do it. Moreover, the fact that pANL-based conjugation can occur through a variety of conjugative systems from *E. coli* to Se7942, but not to other cyanobacteria, suggests that pANL has a narrow replication host range.

What is special about MPF_T ? It is known that MPF_T , MPF_I , and MPF_F share homology in some protein components but also contain specific components that are signatures for each group (18). Thus, one possibility is that these differences could be responsible for the increased promiscuity of MPF_T-type elements. Curiously, transkingdom transfer from bacteria to yeast and plants has only been reported for MPF_T conjugal types (42–44). There is scarce information on factors that determine the range of potential recipients of a given mating system. TraN, one of the MPF_F components of plasmid F, interacts with the outer membrane protein OmpA. This interaction results in stabilization of conjugation partners and is necessary for efficient mobilization (45). Adhesin PilV of the thin pilus encoded by plasmid R64 specifically interacts with the lipopolysaccharide of the recipient cells, determining recipient specificity (46, 47). Receptors similar to OmpA or specific lipopolysaccharide components might be lacking in cyanobacteria and thus prevent the interphylum conjugation of MPF_{F} (F) or MPF_I (R64) plasmids, this being a second alternative to explain MPF_T enhanced promiscuity. In any case, our results clearly demonstrate that conjugation to cyanobacteria is not limited to IncP1 plasmids but involves many MPF_{T} plasmids. These results broaden the number of conjugative systems that can be used for the genetic manipulation of cyanobacteria and explain the origin of Se7942 xenologous genes.

But genetic exchange between proteobacteria and cyanobacteria could occur in both directions. In fact, *E. coli* cells harboring a cointegrate between the shuttle vector pDEP6 and the endogenous Se7942 plasmid pANL, which contains the complete sulfurregulated region of pANL, exhibited increased generation time in the presence of chromate (see Fig. S5 in the supplemental material), as occurred in Se7942 (26). When encoded out of the context of the sulfur-regulated gene cluster, gene *srpC* conferred chromate resistance to *E. coli* (48). On the contrary, a Se7942 *srpC* deletion mutant exhibited a lower doubling time than the wild-type Se7942 in the presence of chromate (26). The reproduction of a Se7942 phenotype in *E. coli* as a consequence of the presence of the pANL genome suggests that the genetic flow between proteobacteria and cyanobacteria could be bidirectional.

Finally, it should be emphasized that several parameters of the conjugation protocol were optimized during the course of the present study to maximize conjugation frequencies to cyanobacteria. First, conjugation worked best at 30°C, with few or no transconjugants obtained at higher temperatures (35, 37, and 40°C were tested), although Se7942 growth rate is maximal at 41°C. Second, *E. coli* strain β 2150 was used as a donor. This strain is auxotrophic for DAP, which helped killing donor *E. coli* cells during the selection regime. Third, several donor/recipient ratios were assayed for each mating experiment in order to obtain countable colonies in one of each series of filters. Other barriers to transphylum conjugation might exist that affect the efficiency of transphylum conjugation, including restriction/modification systems and CRISPRs, among others. Se7942 contains both (49, 50; our unpublished data). In our conjugation assays, the incoming

plasmid DNAs were not methylated for Se7942. It would be interesting to know whether an appropriate modification, such as that devised for RP4-based conjugation to *Anabaena*, which was based on the methylases that protect DNA against restriction by AvaI, AvaII, and AvaIII (51), would result in still higher conjugation frequencies.

ACKNOWLEDGMENTS

Work in the FdlC laboratory was financed by the Spanish Ministry of Economy and Competitivity (MINECO) grant BFU2011-26608 and the European Seventh Framework Program (282004/FP7-HEALTH-2011-2.3.1-2). M.S.-M. is received a Ph.D. fellowship from MINECO. M.P.G.-B. received a JAE-Doc contract from CSIC, cofinanced by the European Science Foundation.

We thank Paula Tamagnini for allowing D.E. to conduct some conjugation experiments in her laboratory, the Pasteur Culture Collection (Institute Pasteur, Paris, France) for supplying Se7942, and K. Takahama for kindly sharing Se7942 strain GRPS1.

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