# Chromosome Transfer in *Rhodobacter sphaeroides*: Hfr Formation and Genetic Evidence for Two Unique Circular Chromosomes

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A 600-bp oriT-containing DNA fragment from the Rhodobacter sphaeroides 2.4.1 S factor (ori $T_s$ ) (A. Suwanto and S. Kaplan, J. Bacteriol. 174: 1124–1134, 1992) was shown to promote polarized chromosomal transfer when provided in cis. A Km<sup>r</sup>-ori $T_s$ -sucR-sacB (KTS) DNA cassette was constructed by inserting ori $T_s$ -sacR-sacB into a pUTmini-Tn5 Km1 derivative. With this delivery system, KTS appeared to be randomly inserted into the genome of R. sphaeroides, generating mutant strains which also gained the ability to act as Hfr donors. An AseI site in the Km<sup>r</sup> cartridge (from Tn903) and DraI and SnaBI sites in sacR-sacB (the levansucrase gene from Bacillus subtilis) were employed to localize the KTS insertion definitively by pulsed-field gel electrophoresis. The orientation of ori $T_s$  at the site of insertion was determined by Southern hybridization analysis. Interrupted mating experiments performed with some of the Hfr strains exhibited a gradient of marker transfer and further provided genetic evidence for the circularity and presence of two chromosomal linkage groups in this bacterium. The genetic and environmental conditions for optimized mating between R. sphaeroides strains were also defined. The results presented here and our physical map of the R. sphaeroides 2.4.1 genome are discussed in light of the presence of two chromosomes.

One essential aspect of the biology of Escherichia coli which has contributed to the importance and usefulness of this organism was the discovery of gene transfer by Lederberg and Tatum in 1946 (21). In addition, the genetic system itself (17) has been used to deduce the circularity of the E. coli chromosome far in advance of any physical mapping techniques (32). For these and other specific reasons dealing with the complexity of genome organization in Rhodobacter sphaeroides 2.4.1 (35, 36), we set about the development of a genetic system equivalent to the Hfr system in E. coli which could be used in conjunction with the already available physical map of this α-purple, nonsulfur photoheterotrophic bacterium.

Several plasmids with a broad host range, such as IncP plasmids RP4 and RK2, have been investigated, but a low frequency of chromosomal transfer has limited their use (16, 31). High-frequency chromosome transfer in R. sphaeroides promoted by a broad-host-range plasmid carrying Tn501 has been reported and has allowed the first extensive genetic mapping of this bacterium (24). However, a lack of understanding about the initiation and orientation of transfer and the nature of RP1::Tn501 insertion into the R. sphaeroides genome has hampered the completion of this map (24). Blanco et al. (2) reported that a retrotrar sfer, i.e., chromosomal markers are transferred not only from the donor but also from the recipient, may add another complication to these broad-host-range-mediated mapping experiments. Nevertheless, this method has allowed construction of a circular linkage map of the R. capsulatus B10 chromosome (48).

An improved method to facilitate chromosome transfer in gram-negative bacteria was initiated by cloning the oriT or

mob region from RP4 or RK2 into Tn5 (13, 30). In a slightly different strategy, Johnson (18) cloned oriT and the entire transfer functions of RP4 into Tn5 and Tn1, yielding new transposons ca. 50 kb in size. In this system, the complete conjugal transfer sequences are provided in cis relative to oriT.

The genome of *R. sphaeroides* 2.4.1 consists of two unique circular chromosomes and five endogenous plasmids (10, 20, 36). The evidence which we have accumulated pointing to the presence of two chromosomes is as follows. (i) the existence of two large (~3,000- and ~900-kb) circular physical DNA structures, (ii) the presence of one *rrn* operon on the large chromosome and two *rrn* operons on the small chromosome, (iii) the presence of tRNA genes on each chromosome, (iv) the 1:1 stoichiometry between chromosomes, and (v) the finding that all strains of *R. sphaeroides* examined by us provided by laboratory collections around the world have two chromosomes comparable in size to the two observed in strain 2.4.1. However, these same strains have variable numbers (from one to six) and sizes of plasmids.

At least two of five endogenous plasmids of R. sphaeroides 2.4.1, i.e. the S factor and plasmid D, are transmissible (37). A 427-bp oriT-containing DNA fragment has been isolated from the S factor  $(oriT_S)$  and been shown to confer high-frequency transfer between R. sphaeroides strains when  $oriT_S$  was cloned into an otherwise nontransmissible plasmid (37).

This study demonstrated the ability of oriT<sub>S</sub> to promote polarized chromosomal transfer in either orientation, the construction of Hfr-like strains by random transposition mutagenesis employing a Tn5 derivative containing oriT<sub>S</sub>, and the application of these developments for strain constructions, classical genetic mapping, and genetic linkage analysis that have been shown by physical analyses of genome structure in previous work (36).

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TABLE 1. Bacterial strains and plasmids

Strain or plasmid	Relevant characteristic(s) <sup>a</sup>	Source or reference
R. sphaeroides 2.4.1		
strains:		S. Dryden and S. Kaplan
L188A	rrnA::Sp/Sm <sup>r</sup>	34
PUHA1	puhA::Km <sup>r</sup>	37
ΔS	2.4.1 cured of S	37
MS2III-48	Green S::Tn5 Nx <sup>r</sup>	This work
48ΔS	MS2III48 cured of S; Nx <sup>r</sup>	This work
PUHA14S	PUHA1 cured of S L188A cured of S	This work
LΔS ΔSR	ΔS, rrnB::Te <sup>τ</sup>	This work
48ΔSR	48ΔS, <i>rrnB</i> ::Τc <sup>r</sup>	This work
HΔSA	PUHA1 $\Delta S$ , puf:: $2\pi i T_S$ - $Sp/Sm^r$ (A orientation)	This work
HΔSB	PUHA1\Delta S. puf::oriTs-Sp/Smr (B orientation)	This work
HΔSAX	HΔSA × 48ΔSR exconjugant; green Tc <sup>r</sup> Km <sup>r</sup> Sp/Sm <sup>s</sup> PS <sup>-</sup>	This work
HΔSBX	HΔSB × 48ΔSR exconjugant; green Tc <sup>r</sup> Km <sup>r</sup> Sp/Sm <sup>s</sup> PS <sup></sup>	This work
Hfr1	ΔS, KTS (I, 861, cw) Aux PS <sup>+</sup>	This work
Hfr2	48ΔS, KTS (1, 2,650, cw) PS <sup>+</sup>	This work
Hfr3	ΔS, KTS (1, 2,078, ccw) PS <sup>+</sup>	This work
Hfr4	48ΔS, KTS (II, 743. cw) PS <sup>+</sup>	This work
Hfr5	ΔS, KTS (II, 3, ccw) PS <sup>+</sup>	This work
L9	LAS, KTS (1, 739, ccw) PS <sup>+</sup>	This work This work
L1	LAS, KTS (I, 2,795, cw) PS+	This work
L <sub>D</sub> Sf5	LAS × Hfr5 exconjugant; Spr Smr KMr PS+	This work
48∆Sf5	L $\Delta$ Sf5 × 48 $\Delta$ SR exconjugant; Nx <sup>r</sup> Kr1 <sup>r</sup> Tc <sup>8</sup> PS <sup>+</sup>	THIS WOLK
E. coli strains:		20
DH5α	$F^-$ lacZ $\Delta$ M15 recA1 hsdR17	29
HB101	Res Mod recall Smr	29 30
S17-1	Pro Res Mod recA integrated plasmid RP4-Tc::Mu-Km::Tn7	14
S17-1 (λ <i>pir</i> )	Same as S17-1, lysogenized with $\lambda pir$ bacteriophage	17
Plasmids		,
pRK2013	ColE1*replicon; Tra+ of RK2, Km <sup>r</sup>	6
pSUP203	pBR325 derivative; mob <sup>+</sup> Tc <sup>r</sup> Cm <sup>r</sup> Ap <sup>r</sup>	30
pSUP202	Same as pSUP2)3, but Mob is located between Cmr and Apr genes	30
pHP45Ω	Source of Sp/Sm <sup>r</sup> cassette	25 25
pHP45Ω-Tc	Same as in pHP45Ω; Tc <sup>r</sup> instead of Sp/Sm <sup>r</sup>	
pUC4K	Source of Km <sup>r</sup> cartridge and intermediate vector	45 29
pBR322	pMB1 replicon; Apr Tcr	42
la	Source of sacR-sacB; single internal EcoRI site in sacB eliminated	14
pUC18Not	pUC18 with No'I sites flanking its multiple cloning sites	14
pUTmini-Tn5 Km1	Mini-Tn5 Kml in plasmid pUT	S. Dryden and S. Kapla
pUC35	5.7-kb EcoRI fragment containing rri B operon cloned into pUC19 HindIII fragment of Sp/Sm <sup>r</sup> cassette inserted into pSUP202-HindIII	S. Dryden and S. Kapla
pSUPssΩ	4.25-kb SphI fregment containing gene Q pufKBALMX cloned into pUC19 such	B. de Hoff and S. Kapla
pUI908	that the direction of pufBA transcription opposes that of lacP	D. do Hon and D. Lorp.
III200	1.9-kb EcoRI-A/wNI fragment of pUC19 ligated with 0.82-kb EcoRI-A/wNI	A. Varga, S. Dryden, ar
pU1389	fragment of pUC18	S. Kaplan
pUI1001	BamHI fragment of Sp/Sm <sup>r</sup> cassette from pHP45 $\Omega$ cloned into pUI389-BamHI	E. Neidle and S. Kaplar
pUC35-TcA	5.5-kb <i>Eco</i> RI fragment containing Tc <sup>r</sup> gene from pMH1701 cloned into pUC35-	This work
pocssien	EcoRV (A orientation)	
pAS404A	11.2-kb EcoRI fragment of pUC35-T:A cloned into pSUPssΩ-EcoRI	This work
pAS303	Cointegrate of pAS302 into S::Tn5: 'fc' Km' Ap'	37
pAS321	~14-kb BelII-XbaI fragment (including internal BelII site) of pAS303 cloned into	This work
p. 135 2 2	nRK415 with Km <sup>r</sup> (BamHI fragment) inserted in outside BglII site	
pAS323	~5.5-kb SalI fragment of pAS321 (1 SalI site derived from Km <sup>r</sup> cartridge in	This work
P	pAS321) clored into pUC4K-Sall	
pAS328	~2.0-kb PstI fragment of Sp/Sm <sup>r</sup> car ridge from pUI1001 inserted into pAS323-PstI	This work
pAS332	~3 9-kh Sall fragment of pAS328 cloped into pUC4K-Sall	This work
pAS341A	2.1-kb PstI fragment of Sp/Sm <sup>r</sup> cartridge from pUI1001 inserted into pAS332-PstI	This work
	(A orientation)	This work
pAS403	3.2-kb AseI fragment of pUI908 cloud into pSUP203-AseI	This work
pAS348A	6-kb EcoRI fragment of pAS341A inserted into the XhoI site in pAS403 by blunt-	This work
	end ligation (A orientation)	This work
pAS348B	Same as pAS348A but in B orientation (opposite of A orientation)	This work This work
pAS346	3.2-kb Smal-EcoRI fragment of pAS341A (including Sp/Sm <sup>r</sup> ) cloned into pUC19-	TIII2 MOLK
	Smal-EcoRI	This work
pAS354	1.3-kb Sall fragment of pAS346 cloued into pUC4K-Sall	TIII2 MOIK

TABLE 1-Continued

Strain or plasmid	Relevant characteristic(s)"	Source or reference
pAS375A	2 kb of sacB gene (BcmHI filled in) from p asmid Ia cloned into pAS354-Stul- Smal (A orientation)	This work
pAS377	2.7-kb XbaI-SstI fragment (sacB-oriT <sub>S</sub> ) of pAS375A cloned into pUC18NOT-XbaI-SstI	This work
pAS378	2.7-kb Not1 fragment (sacB-oriT <sub>S</sub> ) of pAS3 <sup>17</sup> cloned into pUTmini-Tn5 Km1-Not1 such that the orientation is KTS	This work
pAS380	2-kb Smal fragment containing ΩTc from pHP45Ω-Tc inserted into pAS378- Scal	This work
pAS384	3.4-kb <i>Dral-SspI</i> fraginent of pBR322 (containing <i>oriV</i> and Tc <sup>r</sup> gene) inserted into pAS378-Scal	This work

<sup>&</sup>quot;The location and orientation of KTS are indicated in parentheses following KCS. I and II indicate insertion in the large and small chromosomes, respectively, and the number in the middle indicates the relative distance (in kilobases) from  $criT_S$  to puf (in the large chromosome) or rrnC (in the small chromosome). PS, photosynthetic growth; Aux, auxotroph.

compare or contrast the strains under study physically. Thus, as stated above, the AseI schizotype of strain  $\Delta S$  is identical to that of strain 2.4.1.

puhA is located ca. 31 kb from pufBA (35), so that the presence of the Kmr insertion containing an Asel site in PUHA1 resulted in digestion of the 73-kb AseI fragment present in  $\Delta S$  into 31- and 42-kb Asel fragments in PUHA1AS (35). The Sp/Smr cartridge has two Asel sites flanking the gene for Sp/Sm<sup>r</sup> (25), so insert on of oriT<sub>S</sub>-Sp/ Smr into pufM, depending on the orientation, should yield a predictable AseI schizotype for strains HΔSA and HΔSB, as depicted in Fig. 1B. AseI schizotyping of strains  $\Delta S$ ,  $H\Delta SA$ , and HASB, in conjunction with Southern hybridization analysis using a 1.3-kb SalI fragment of pAS354 (Table 1) containing  $oriT_S$ , unambiguously demonstrated that H $\Delta SA$ and H $\Delta$ SB contained the recombined DNA fragment ( $oriT_s$ -Sp/Sm<sup>r</sup>) in opposite orientations (Fig. 2). The hybridization signals at 19.5 (H $\Delta$ SA) and 5.6 (H $\Delta$ SB) kb revealed the presence of  $oriT_S$  in opposite orientations in these recombinant strains. The hybridization signal at 31 kb was due to the similarity between DNA sequences in the S factor (including the  $oriT_{\rm S}$  region) and plasmid D (10, 37). The 31- and 63-kb AseI fragments are derived from plasmid ) (see plasmid fingerprinting in reference 37). A weak hybridization signal also detected at the 110-kb AseI fragment corresponds to pRS241a (37).

Recipient strain construction. A 5.5-kb EcoRI fragment containing the  $Tc^r$  gene from pMH1701 (37) was inserted into the EcoRV site in the rrnB operon (8), which is located on chromosome II. The entire DNA fragment containing rrnB:: $Tc^r$  (ca. 11.2 kb) was excised as an EcoRI fragment and inserted into the EcoRI site of pSUPss $\Omega$  to yield pAS404A, and the altered rrnB operon was ntroduced into the small chromosome of strains  $\Delta S$  ( $Crt^+$ ) and  $\Delta S$  ( $Crt^-$ ) as previously described (37; Table 1) to yield strains  $\Delta SR$  and  $\Delta SR$ , respectively (Table 1).

Chromosome transfer from strain H $\Delta$ SA or H $\Delta$ SB into strain 48 $\Delta$ SR. The relevant characteristics of these donor and recipient strains are indicated in Table 1, and the structures are shown in Fig. 1A. Since H $\Delta$ SA and H $\Delta$ SB were not able to grow photosynthetically, mating was performed under LB-DMSO conditions as described in Materials and Methods. The results are summarized in Table 2.

Mating between H $\Delta$ SB and 48 $\Delta$ SR consistently yielded at least 10-fold more exconjugants than that of H $\Delta$ SA and 48 $\Delta$ SR. In fact, under short-term mating conditions (8 h), we were unable to detect any Km<sup>r</sup> exconjugants from H $\Delta$ SA  $\times$  48 $\Delta$ SR. From these results, the orientation of  $oriT_S$  can be

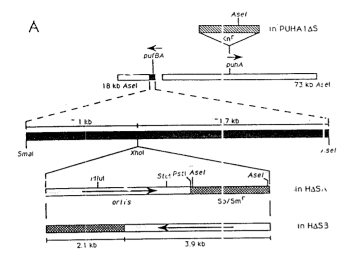
deduced such that  $Km^r$  in H $\Delta SB$  will be one of the earliest markers (ca. 30 kb from  $oriT_S$ ) and  $Km^r$  in H $\Delta SA$  will be one of the last markers (ca. 3,000 kb from  $oriT_S$ ) to be transferred into the recipient (Fig. 1A).

 ${\rm H}\Delta{\rm SB} \times 48\Delta{\rm SR}$  matings yielded 70% green and 30% red exconjugants. All of the green exconjugants were  ${\rm Sp/Sm^s}$ , and 90% of the red exconjugants were  ${\rm Sp/Sm^r}$ . These genetic results suggested that the *crt* marker was more closely linked to  ${\rm Sp/Sm^r}$  (inserted into the *puf* operon) than to *puhA*. These genetic results were supported by the actual physical distances of the *crt* marker, which are approximately 11 kb from *puf* and 20 kb from *puhA* (38).

HΔSA × 48ΔSR mating yielded 99% red, Sp/Smr exconjugants and was the result of the orientation of  $oriT_S$  in this donor. The Kmr marker in this strain would be one of the last markers to enter the recipient (ca. 31 kb before the point of chromosome recircularization), so that the exconjugants were more likely generated from the transfer and stabilization of all of chromosome I, as was the case in 27% of the exconjugants of H $\Delta$ SB  $\times$  48 $\Delta$ SR. This result also helps to explain why, in the short period of mating (LB-DMSO for 8 h), we were unable to observe any exconjugants from the  $H\Delta SA \times 48\Delta SR$  mating, since in this mating we demanded the transfer of nearly all of chromosome I. The generation time of R. sphaeroides grown anaerobically in the dark in LB-DMSO is approximately 9 to 12 h or three times longer than when it is grown photosynthetically at 10 W/m2, so that the overall metabolic activity, including chromosome transfer, would be expected to be slower than that observed during optimum photosynthetic mating conditions (see below).

To determine whether introduction of  $oriT_S$  from H $\Delta$ SA and H $\Delta$ SB into strain 48 $\Delta$ SR could induce plasmid cointegration or rearrangement mediated by plasmid D (37), representative colonies among the exconjugants from H $\Delta$ SA or H $\Delta$ SB matings with 48 $\Delta$ SR were subjected to plasmid fingerprinting analysis (37). The results showed that all of the representative exconjugants had normal plasmid profiles. Despite the homology between plasmid D and  $oriT_S$ , no recombination or rearrangements were observed in these exconjugants. These representative exconjugants were further examined by schizotyping as described previously (35). The results showed the expected schizotypes. Similarly, the exconjugants displayed the anticipated phenotypes.

Construction of a suicide vector carrying Km<sup>r</sup>-oriT<sub>S</sub>-sacR-sacB (KTS). Since we were able to demonstrate polarized chromosome transfer directed by oriT<sub>S</sub>, we sought to develop a mechanism by which we could readily construct



## B The expected Asel-schizotypes of:

2.4.145	PUHA14S	Hasa	наѕв
<del></del>	42	42	42
	31	31	31
1.6	18	195*	16.3
			56*
		2 1	2.1
		1 7	

FIG. 1. Construction of donor strains containing  $oriT_S$  inserted in two different orientations. (A) A 6-k3 EcoRI DNA cassette consisting of oriT<sub>S</sub> (3.9 kb) and the gene for Sp/Sm<sup>r</sup> (2.1 kb) was excised from pAS341A. This fragment was inserted into a unique XhoI site in pAS403 by blunt-end ligation, yielding recombinant suicide plasmids each with a 6-kb oriT<sub>S</sub>-Sp/Sm<sup>r</sup> gene cassette in different orientations (pAS348A and pAS348B). Introduction of each of these plasmids into strain PUHA12S generated two kinds of Hfr donor strains (HASA and HASB) with respect to the relative orientation of the DNA fragment containing oriT<sub>S</sub>. (B) Diagram of the expected AseI schizotypes of HΔSA and HΔSB in compar son with those of  $\Delta S$  and PUHA1 $\Delta S$ . Other AseI fragments larger than 73 kb in these four strains are identical and are not depicted. A Km<sup>r</sup> cartridge in strain PUHA1 AS results in digestion of the 73-kb AseI fragment into 42- and a 31-kb AseI fragments upon digestion with AseI (35). These 42- and 31-kb AseI fragments should be conserved in strains HASA and HASB. The Spr Smr cassette has two Asel sites (25); therefore, insertion of a 6-kb DNA cassette containing or iTs-Sp/Smr in the 18-kb AseI fragment generates three new AseI fragments with molecular sizes corresponding to the relative o ientation of oriT<sub>S</sub>-Sp/Sm<sup>r</sup> in the chromoson e. AseI fragments which carry  $oriT_S$  sequences are indicated by asterisks (see also Fig. 2).

additional Hfr-like strains by random insertion of  $oriT_S$  into the R. sphaeroides 2.4.1 genome. Moreover, the  $oriT_S$ -containing DNA fragment which we used was only 600 bp, while that used to construct H $\Delta$ SA and H $\Delta$ SB was 3.9 kb.

Plasmid pUTmini-Tn5 Km1 (14, 22) was chosen as a vehicle for construction of an  $oriT_S$  delivery system. Modification of this plasmid by insertion of  $oriT_S$ , sacR-sacB, and the Tc<sup>r</sup> gene yielded recombinant suicide plasmid pA 3380 (Fig. 3A). This plasmid (Fig. 3B) has a  $\gamma$  origin of replication from plasmid R6K; therefore, it can replicate only in a host providing a Pir protein, such as SM10 $\lambda$ pir or S17-1 $\lambda$ pir (14). Transfer into the recipient was achieved by utilizing RP4

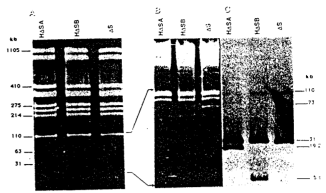


FIG. 2. Genome analysis of strains H $\Delta$ SA and H $\Delta$ SB. (A) The Asel schizotypes of DNA fragments larger than 73 kb from strains H $\Delta$ SA and H $\Delta$ SB are identical to that of the reference strain ( $\Delta$ S). (B) The Asel schizotypes of the smaller-molecular-size fragments ( $\leq$ 73 kb) of strains H $\Delta$ SA and H $\Delta$ SB showed the expected results. as depicted in Fig. 1B. (C) Confirmation of the restriction patterns in panel B by Southern hybridization analysis using a 1.3-kb Sall fragment containing  $oriT_S$  from pAS354 as a probe. The strong signal at the 31-kb Asel fragment and the weak signal at the 110-kb Asel fragment are due to cross-hybridization of the probe with sequences from Asel-digested plasmids D and pRS241a, respectively (37). The TAFE conditions for panel A were as follows: stage 2, 50-s pulse. 6 h; stage 3, 23-s pulse, 8 h; stage 4, 7-s pulse, 4 h; stage 5, 4-s pulse, 2 h. Those for panel B were as follows: stage 2, 8-s pulse, 8 h; stage 3, 4-s pulse, 6 h; stage 4, 2-s pulse, 3 h.

mob, which can be driven by the products of the tra genes of plasmid RP4, which are provided in trans, such as in E. coli S17-1 (30). Transposition, as opposed to plasmid cointegration, was indirectly ascertained by replica patching on Tc-containing medium. The gene for Km<sup>r</sup> (from Tn903) has a unique AseI site, while the sacB gene contains SnaBI and DraI sites. These restriction enzyme sites facilitate precise localization of oriT<sub>S</sub> (or KTS in its entirety) in the R. sphaeroides genome (35).

In addition, we also constructed a derivative of pAS378 which can replicate independently of the Pir protein by inserting a 3.4-kb *DraI-SspI* fragment from pBR322 containing both *oriV* and the Tc<sup>r</sup> gene into the *ScaI* site of pAS378. This chimeric plasmid (pAS384) is essentially identical to

TABLE 2. Analysis of exconjugants from H $\Delta$ SA  $\times$  48 $\Delta$ SR and H $\Delta$ SB  $\times$  48 $\Delta$ SR matings

Mating"	Frequency <sup>b</sup>	Description
PUHA1ΔS × 48ΔSR	<10-9	Chromosomes not able of transfer by themselves
$H\Delta SB \times 48\Delta SR$	$10^{-6} - 10^{-7}$	70% green Sp/Sm <sup>s</sup> , 27% red Sp/Sm <sup>r</sup> , 3% red
$\dot{\text{H}}\dot{\Delta}\text{SA} \times 48\Delta\text{SR}$	$10^{-7} - 10^{-8}$	Sp/Sm <sup>s</sup> 99% red Sp/Sm <sup>r</sup> , 1%  green Sp/Sm <sup>s</sup>

<sup>&</sup>quot; Mating was performed in the dark in liquid LB-DMSO medium at 32°C for 4 h.

<sup>b</sup> Calculated as the number of exconjugants (Tc<sup>r</sup> Km<sup>r</sup>) per donor (Km<sup>r</sup> Sp/Sm<sup>r</sup>) obtained from the average of three separate experiments.

 $<sup>^{\</sup>circ}$  Sp/Sm $^{\circ}$  and Sp/Sm $^{\circ}$  phenotypes were calculated from replica patches of 100 randomly picked colonies of the H $\Delta$ SB  $\times$  48 $\Delta$ SR mating and from all of the colonies which appeared from the H $\Delta$ SA  $\times$  48 $\Delta$ SR mating (20 to 50 colonies). All exconjugants were PS $^{-}$ , since selection for Km $^{\circ}$  concomitantly generated a strain with inactive puhA (see the text for details).

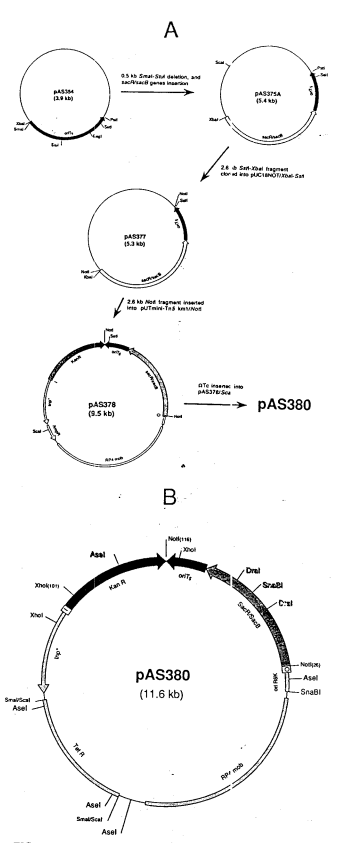


FIG. 3. Construction and physical structure of pAS380. (A) Construction of pAS380. A 0.5-kb SmaI-StuI fragment was deleted from plasmid pAS354 (Table 1) and replaced vith a 2-kb BamHI fragment containing the sacR and sacB genes from plasmid la which was made blunt ended by Klenow DNA polymerase, to yield

pAS380, except that it carries a pBR322 (pMB1) origin of replication (Table 1).

Random KTS insertion into the R. sphaeroides 2.4.1 genome. S17-1(pAS380) was mated with strains L $\Delta$ S, 48 $\Delta$ S, and  $\Delta$ S (Table 1). The exconjugants were selected on LB-Sp-Sm-Km for L $\Delta$ S and on Sis-Km for 48 $\Delta$ S and  $\Delta$ S. S17-1(pAS380) cannot grow on Sis minimal medium, since S17-1 is a proline auxotroph. Km<sup>r</sup> exconjugants were obtained at a low frequency ( $10^{-7}$  to  $10^{-8}$  per donor), despite several repeated matings. This may have been due to the inherent properties of this engineered transposon, in which  $tnp^*$  has been displaced outside the I and O ends (7), since similar transposition utilizing S17-1(pMH1701) (37) routinely yields high-frequency transposition ( $10^{-5}$  to  $10^{-6}$  per donor) (38).

Kmr exconjugants were replica patched onto Tc-containing medium to screen for exconjugants generated from cointegration of all of pAS380 with R. sphaeroides genomic DNA. From 80 to 90% of the Kmr exconjugants were Tcs. Some of these colonies were further analyzed to determine the location and relative orientation of oriT<sub>s</sub>. The precise location of KTS in the exconjugants was determined by digestion of total genomic DNA with restriction endonucleases AseI, SnaBI, and DraI, and then each of the schizotypes was compared with those of the parental strains (i.e.,  $\Delta S$ , 48 $\Delta S$ , and L $\Delta S$ ). The relative orientation of  $oriT_S$  in each strain was determined by Southern hybridization analysis using an oriT<sub>s</sub>-containing DNA fragment as a probe. This approach generated Hfr-like donors with different points of origin within the R. sphaeroides 2.4.1 genome, including the four remaining endogenous plasmids. Several representative Hfr strains with  $oriT_S$  located in the chromosomes are described in Fig. 4 and Table 1.

A gradient of chromosomal marker transfer directed by  $oriT_S$ . Strains L1 and L9 possess a KTS insertion at about 11 and 3 o'clock, respectively, in chromosome I (Fig. 4). Each of these strains also has an Sp/Sm<sup>r</sup> marker in rrnA (at 5 o'clock) and dominant normal Crt<sup>+</sup> (red) pigmentation as an additional genetic marker located at 12 o'clock (36). The orientation of  $oriT_S$  in L9 is counterclockwise (ccw), while in L1 it is clockwise (cw).

To demonstrate the gradient of transfer of the Sp/Sm<sup>r</sup>-red-Km<sup>r</sup> markers in both L1 and L9, we performed an interrupt-ed-mating experiment using strain 48\Delta SR as the recipient. Strain 48\Delta SR, as already described (Table 1), has a Tc<sup>r</sup> marker in chromosome II and a recessive Crt<sup>-</sup> (green) pigmentation.

The conditions and results of L1  $\times$  48 $\Delta$ SR and L9  $\times$  48 $\Delta$ SR matings are described in Fig. 5. As a control, L $\Delta$ S

plasmid pAS375A. A 2.6-kb Sstl-Xbal fragment of pAS375A was cloned into the Sstl-Xbal sites in pUC18NOT (Table 1) and yielded pAS377. A Notl fragment containing sacR-sacB and  $oriT_s$  from pAS377 was excised and inserted into a unique Notl site in pUT-mini-Tn5 Km-1. A 2-kb Smal fragment containing an  $\Omega$ Tc cassette from pHP45 $\Omega$ Tc was inserted into the Scal site of pAS378 to generate pAS380. (B) Physical structure and some relevant genetic loci on pAS380. The arrowheads in  $oriT_s$ , the gene for Km<sup>r</sup>,  $tnp^*$ , and sacR-sacB indicate their relative orientations.  $tnp^*$ , ori R6K, the gene for Km<sup>r</sup>, and RP4 mob were described previously by Herrero et al. (14). I and O denote the I and O ends of IS50, which also indicate the border of the transposed elements, i.e., KTS. The Asel, Dral, and SnaBl restriction sites in KTS, used to localize the KTS insertions, are highlighted (shadowed). Other Dral sites may be present in the vector DNA.

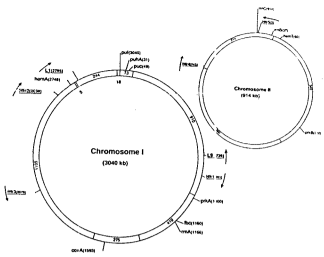


FIG. 4. Location and orientation of  $oriT_S$  in some representative Hfr donor strains generated by KTS insertion. The size of each AscI fragment (36) is shown inside or below the fragment. pufBA and rrnC are arbitrarily assigned the 12 o'clock position on the physical map of chromosomes I and II, respectively. The number in parertheses following a gene or Hfr designation indicates the relative distance (in kilobases) of that particular gene or  $oriT_S$  in each Hfr strain clockwise to either pufBA or rrnC. The arrow by each Hfr strain indicates the  $oriT_S$  orientation in that H r strain. For example,  $oriT_S$  in L9 was located ca. 739 kb from pufBA (or at 3 o'clock) in a ccw direction from pufBA.

(the parent strain of L1 and L9) was majed with 48 $\Delta$ SR and no exconjugants (<10<sup>-9</sup> per donor) were detected after 7, 9, and 20 h of mating. These results are completely consistent with the physical distances of Sp/Sm<sup>r</sup> from the origin of transfer, which are 400 kb in L9 and 1,600 kb in L1.

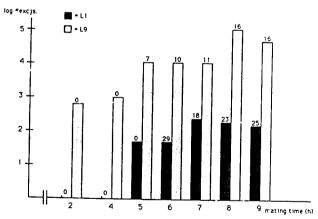


FIG. 5. Histogram of interrupted-mating experiments of L1  $\times$  48 $\Delta$ SR and L9  $\times$  48 $\Delta$ SR. Donors and recipients were pregrown photosynthetically in Sis-10% LB until the concentration was approximately 2.0  $\times$  10° viable cells per ml, which corresponds to a 16- to 18-h incubation time. One milliliter each of the donor and recipient were mixed in 5-ml Wheaton glass via s filled with LB and incubated photosynthetically. At the end of each mating period, 200 to 500  $\mu$ l of the mating mixture was withdrawn and agitated vigorously in a Vortex mixer. The exconjugants (excjs.) were calculated from the colonies which appeared on LB-Sp-Sm-Tc plates. The frequencies of total exconjugants a ter 7 h of mating for L1 and L9 were 2  $\times$  10<sup>-5</sup> and 3  $\times$  10<sup>-7</sup> per don r, respectively. The numbers above the bars are percentages of red colonies in the total number of colonies appearing on LB-Sp-Sm-Tc plates.

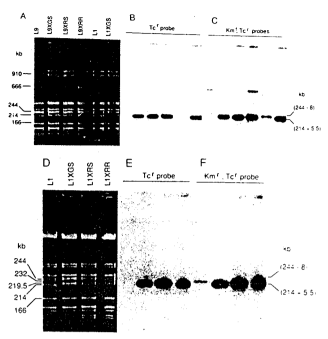


FIG. 6. Genome analysis of L1  $\times$  48 $\Delta$ SR and L9  $\times$  48 $\Delta$ SR exconjugants. (A) Asel schizotypes of the following strains: L9. L9XGS (L9 exconjugants, green, Kms), L9XRS (L9 exconjugants. red. Km<sup>s</sup>), L9XRR (L9 exconjugants, red. Km<sup>r</sup>), L1, and L1×GS (L1 exconjugants, green, Kms). (B) Autoradiogram of panel A with the gene for Tcr as a probe. (C) Autoradiogram of panel A with the genes for Tc<sup>r</sup> and Km<sup>r</sup> as probes. (D) Asel schizotypes of the following strains: L1, L1XGS, L1XRS (L1 exconjugants, red. Km'). and L1XRR (L1 exconjugants, red, Km<sup>r</sup>). Panels E and F are autoradiograms of panel D with the genes for Tcr and Tcr Kmr. respectively, as probes. The Tcr gene probe was a 5.5-kb EcoRI fragment containing the Tcr gene from pMH1701 (37). The Kmr gene probe was a BamHI fragment containing the Kmr gene from pUC4K. Hybridization signals appearing near the top edges of panels B, C, E, and F are due to the remnants of DNA which stayed in the wells of TAFE gels. The TAFE conditions for panel A were as follows: stage 2, 48-s pulse, 8 h; stage 3, 23-s pulse, 7 h; stage 4, 7-s pulse, 3 h. The TAFE conditions for panel D were as follows: stage 2, 25-s pulse, 9 h; stage 3, 20-s pulse, 5 h; stage 4, 10-s pulse.

Transfer of the red (crt) marker was detected after 5 h of mating and continued to increase until 9 h of mating. Because of the distance of the crt marker relative to the origin of transfer, the frequency of inheritance of red pigmentation was lower than that of Sp/Smr. Further analysis of the exconjugants indicated that all of the green Tcr Sp/Smr exconjugants were Kms, while the red exconjugants from L9 and L1 donors were 89 and 97% Km<sup>r</sup>, respectively. The genetic status of the exconjugants was further examined at the DNA level following schizotyping and Southern hybridization analysis as described in the legend to Fig. 6. The green Km<sup>s</sup>, red Km<sup>s</sup>, and red Km<sup>r</sup> exconjugants from L1 and L9 donors exhibited the expected AseI schizotypes. Strains L9 and L1 are both derived from strain LΔS, which has an Sp/Smr cassette inserted in rrnA (Table 1 and Fig. 4), so that AseI digests the 410-kb fragment normally present in strain ΔS (wild type) into new 244- and 166-kb AseI fragments because of the presence of AseI sites in the Sp/Smr cassette (25). In addition, KTS insertion in L9 results in cleavage of the 910-kb Asel fragment into new 244- and 666-kb Asel fragments because of the presence of AseI sites in the Km<sup>r</sup>

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gene, while KTS insertion in strain L1 generates new ca. 236- and 8-kb Asel fragments from the 244 kb Asel fragment present in the wild type. Therefore, upon digestion with Asel, strain L9 gives rise to 666- and 166-k5 fragments and a triplet of 244-kb Asel fragments, while strain L1 shows 236and 8-kb fragments and a doublet of 244-kb AseI fragments (Fig. 6A and 6D). The 8-kb AseI fragment was not detected under these pulsed-field gel electrophoresis conditions. Strain 48\Delta SR has a Tcr gene inserted into the 214-kb AseI fragment, and there is no AseI site present in this Tcr cassette, so the presence of this cassette can be detected because of the increased molecular size of the particular fragment into which Tcr was inserted.

The L9 green Km<sup>s</sup> and red Km<sup>s</sup> excon ugants showed a 5.5-kb Tcr insertion in the 214-kb AseI fragment, a doublet of 244 kb, and a 166-kb AseI fragment, which confirmed the presence of the Sp/Smr cassette in the 410-lib AseI fragment. Therefore, these exconjugants were new strains with genotypes different from those of their parental strains. The genotypes of these exconjugants are rinA::Sp/Smr and rrnB::1c<sup>r</sup>, with Crt<sup>-</sup> (green) or Crt<sup>+</sup> (red) resulting from chromosome I transfer from L9 to 48ASR.

The L9 red Kmr exconjugants exhibited an AseI schizotype identical to that of the Km<sup>s</sup> L9 exconjugants, with the only exception being that the 910-kb AseI fragment was digested into 244- and 666-kb Asel fragments. Digestion of the 910-kb AseI fragment in this strain indicated the presence of KTS. Thus, the genotype of these exconjugants is rrnA::Sp/Smr rrnB::Tcr Crt+ KTS insertio 1 at 9 o'clock on chromosome I in a ccw orientation. These results suggested that the Kmr exconjugants from this mating should be the result of transfer and recircularization of all of chromosome I into the recipient cells.

Similarly, the L1 exconjugants showed an AseI schizotype and Southern hybridization analysis results which were entirely consistent with the interpretation of the conjugation data (Fig. 6D, E, and F). For example, Km<sup>r</sup> exconiugants exhibited a KTS insertion in the 244-kb Ase. fragment and an Sp/Smr insertion into the 410-kb AseI fragment, as in the donor strain (L1). However, the donor strain was unambiguously distinguished from the exconjugants by its lack of Tc<sup>r</sup> (Fig. 6E and F). Schizotyping in conjunction with Southern hybridization analysis in this experiment clearly demonstrated that the recipients acquired either an Sp/Smr gene or a Km<sup>r</sup> gene from the donors during mating to yield various exconjugants, as stated above. These combined data suggested that  $oriT_S$  is able to direct a gradient transfer of chromosomal markers, and all of chromosome I may recircularize into the recipient cells.

Genetic evidence of two unique circular chromosomes. Mating between strains Hfr5 (Table 1; Fig. 4) and LΔS was performed as described in Materials and Methods. The exconjugants were selected on LB-Km-Sp-Sm. The results showed that chromosome II was transferred into the recipient strain, since acquisition of Kmr by LAS was achieved only after transfer and recircularization of chromosome II from the donor (Table 3).

L $\Delta$ Sf5 is an exconjugant from the mating of L $\Delta$ S and Hfr5. The genotype of this strain is rrnA::Sp/Smr KTS (II, 3, ccw) (Table 1; Fig. 4). Schizotyping verified the genotype of this exconjugant (Fig. 7). To examine the donor property of LΔSf5, it was mated with strain 48ΔSR. Selection was made on LB containing Km-Nx, Km-Tc, or Km-Sp-Sm. As shown in Table 3, selection on Km-Nx yielded the highest frequency of exconjugants. Although the donor (L\Delta Sf5) itself can spontaneously mutate to Nxr, it is easily distinguished

TABLE 3. Transfer of chromosome II

Mating"	Selection	Frequency <sup>b</sup>	Description <sup>c</sup>	
$L\Delta S \times 48\Delta SR$	Sp-Sm-Tc	<10-9	No exconjugants	
$L\Delta S \times Hfr5$	Sp-Sm-Km	$2.0 \times 10^{-4}$	100% red	
$L\Delta Sf5 \times 48\Delta SR$ $L\Delta Sf5 \times 48\Delta SR$ $L\Delta Sf5 \times 48\Delta SR$	Km-Nx Km-Tc Sp-Sm–Tc	$2.0 \times 10^{-5}$ $1.0 \times 10^{-7}$ $1.0 \times 10^{-7}$	Tc <sup>s</sup> , 99% green <sup>d</sup> 100% green 50% green, 50% red	
$48\Delta Sf5 \times L\Delta S$	Km-Sp-Sm	$2.0 \times 10^{-6}$	100% red	

" Strain 48ΔSR or LΔS was the recipient in these Hfr matings.

Relevant phenotypes of exconjugants.

from 48\Delta SR because of its red pigmentation. The exconjugants of this mating (green Nxr Kmr) were designated 48ΔSf5. Replica patches of 48ΔSf5 indicated that all of these exconjugants (from 100 representative colonies) were Tcs. Furthermore, pulsed-field gel electrophoretic analysis of 48ΔSf5 showed a schizotype identical to that of Hfr5. Therefore, chromosome II::Km<sup>r</sup> from donor strain LΔSf5 was transferred into recipient strain 48ΔSR and concomitantly replaced the resident chromosome II::Tcr in 48\Delta SR.

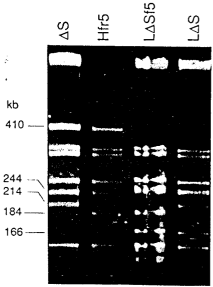


FIG. 7. Schizotype verification of strain LΔSf5. Strain LΔS has an Sp/Smr insertion in the 410-kb AseI fragment such that the AseI schizotype of this strain yielded new 244- and 166-kb Asel fragments. Thus, the 244-kb AseI fragment in LAS is a doublet. Strain Hfr5 has a KTS insertion in the 214-kb AseI fragment such that the 214-kb Asel fragment in this strain was digested into 184- and 30-kb Asel fragments. Lasf5 is strain Las which has undergone chromosome II displacement with chromosome II::Kmr from strain Hfr5. The Asel schizotype of this strain yielded a hybrid pattern inherited from its parental strains; i.e., both the 410- and 214-kb AseI fragments in the wild type ( $\Delta S$ ) were replaced by four new Asel fragments with molecular sizes of 244, 184, 166, and 30 kb. The 30-kb Asel fragment was not clearly visible with this TAFE condition. The TAFE conditions for this experi nent were as follows: stage 2, 30-s pulse, 8 h; stage 3, 23-s pulse, 3 h.

<sup>&</sup>lt;sup>b</sup> Calculated as the number of exconjugants divided by the number of donors. The results are averages from two separate experiments.

<sup>&</sup>lt;sup>d</sup> The red colonies (1%) in this mating were spontaneous Nx<sup>r</sup> derivatives of strain LaSf5 (see the text for details).

Consequently,  $48\Delta Sf5$  should be a third-generation donor strain generated from three successive matings (i.e.,  $Hfr5 > L\Delta Sf5 > 48\Delta Sf5$ ) as shown in Table 3.

Although chromosome II::Km<sup>r</sup> from L\DeltaSf5 successfully replaced the resident chromosome II::Tc<sup>r</sup> in 48\DeltaSR the pigmentation of 48\DeltaSf5 was still green, as was that of the parental recipient strain (48\DeltaSR). This result suggested that although all of chromosome II had been transferred the dominant crt marker (i.e., red) was not inherited by the recipient. Therefore, the crt marker should be located on another replicon which is different from chromosome II. In fact, from our physical mapping analysis (38) we knew that the crt marker is located between puf and puhA on chromosome I. These data also suggested that the Nx<sup>r</sup> marker is not located on chromosome II, since chromosome II::Km<sup>r</sup> replaced chromosome II::Tc<sup>r</sup> at a high frequency in the Nx<sup>r</sup> recipient (48\DeltaSR).

Selection of an LASf5 × 48ΔSR mating mixture on Km-Tc yielded fewer exconjugants (10<sup>-7</sup> per donor) than when selection was made on Km-Nx. Since both the Km<sup>r</sup> and Tc<sup>r</sup> markers are located on chromosome II, for inheritance of Km<sup>r</sup> Tc<sup>r</sup>, chromosome II::Km<sup>r</sup> from the donor must no just replace the resident chromosome II::Tc<sup>r</sup> from the recipient but must subsequently recombine with the resident chromosome II with the Km<sup>r</sup> and Tc<sup>r</sup> markers in the same replicon. Since the distance between Tc<sup>r</sup> (in rrnB) and KTS of 48ΔSf5 (near rrnC) is relatively short (~30 kb), we anticipate that this recombination event would be rather infrequent, thereby resulting in the low apparent transfer frequency upon simultaneous Km-Tc selection.

Selection of an LASf5 × 48ASR mating mixture on Sp-Sm-Km yielded a low frequency of exconjugants (Table 3). Since the Sp/Sm<sup>r</sup> marker is located on chromosome I, we did not anticipate finding a high frequency of exconjugants from this selection. Half of these exconjugants were red. This probably resulted from transfer of chromosome II::Km<sup>r</sup> into 48ASR following recombination with the resident chromosome II::Km<sup>r</sup> to gain Tc<sup>r</sup>, and then this chromosome II::Km<sup>r</sup> Tc<sup>r</sup> was retransferred into LASf5, yielding Km<sup>r</sup> Tc<sup>r</sup> Sp/Sm<sup>r</sup> exconjugants. The other exconjugants, which were given, indicated that both the Km<sup>r</sup> and Sp/Sm<sup>r</sup> markers were transferred into the recipient.

Putting all the data together, we have been able to demonstrate genetically that *R. sphaeroides* 2.4.1 contains two separate linkage groups (previously designated chromosomes I and II (36)). Unidirectional chromosome transfer from the Hfr donor strains was mediated by  $oriT_S$  inserted into either chromosome I or chromosome II. The exconjugants generated by either chromosomal displacement or KTS inheritance will behave as new Hfr donor strains.

#### DISCUSSION

This study demonstrated that a 600-bp oriT<sub>S</sub>-containing DNA fragment was able to promote chromosome transfer in an oriented manner when provided in cis. Thus far, outside of enteric bacteria, chromosome transfer mediated by endogenous plasmids in gram-negative bacteria has been reported only in Pseudomonas sp. (16), Agrobacterium tumefaciens (5), and this study. Under optimal mating conditions, a chromosomal marker located at a distance of 400 kb from the origin of transfer was mobilized into the recipient at a frequency of 10<sup>-5</sup> to 10<sup>-4</sup> exconjugants per donor in a 2- to 5-h mating. The Tra functions required for conjugal transfer in this system appear to be provided by plasmid D,

another transm ssible plasmid of *R. sphaeroides* 2.4.1 (37), although defini ive proof of this conclusion is lacking. The stable presence of plasmid D provides a simple approach to the construction of Hfr donors. However, since plasmid D is also present in recipient cells derived from strain 2.4.1 and if it is analogous to F or the F-like plasmids of *E. coli*, then surface exclusion mechanisms (47) and repression-derepression of plasmid transfer mediated by plasmid D may be involved in determining the ultimate transfer frequency. Further work will be required to verify this.

KTS insertion mutagenesis, in conjunction with schizotyping, has been used to locate  $oriT_S$  precisely within the genome of R. sphaeroides. These insertions appear to be stable, since the transposase gene  $(tnp^*)$  is not part of the transposed element (14). All of the resulting  $oriT_S$ -containing strains have been demonstrated to behave like Hfr donors, and some of these Hfr-like strains have been used to demonstrate a gradient of chromosomal transfer. By using a genetic approach, the presence of two unique circular chromosomes was revealed. This analysis supports the results of physical mapping (36) and the use of  $\gamma$  irradiation (46).

Although circular chromosomes are the most common chromosomal topology in bacteria (20), linear chromosomes have also been described (9, 20). Similarly, although bacterial plasmids are predominantly double-stranded circular DNA molecules, some linear DNA plasmids have been reported (9, 19). It is perhaps not a surprise that continued investigation of diverse groups of bacteria has laid to ruin the initial dogma surrounding the bacterial genome. Another element of that dogma has been the belief that bacteria have only a single chromosome, although in certain stages of growth or in certain bacteria multiple copies of the same chromosome can be present (27).

Several gram-negative bacteria which are phylogenetically related to R. sphaeroides (49), such as Rhizobium, Agrobacterium, Alcaligenes, Pseudomonas, and Paracoccus spp., harbor very large extrachromosomal replicons termed megaplasmids (11, 15, 33, 44). The sizes of some of these megaplasmids are much larger than the chromosomes of certain bacteria (20, 46). However, many of these very large replicons have been shown to be self-transmissible (1, 3, 5, 12, 26) and/or curable without an overall deleterious effect on the growth of the organism under all physiological conditions; i.e., loss of the replicon appears to affect only a single growth mode characteristic of the organism. Thus, self-transmissibility and size appear to be two of the criteria used to distinguish plasmids from chromosomes in bacteria. Invariably, when chromosome transfer is recognized it has been found to be mediated by acquisition of functions normally found associated with plasmids and transposons (4, 16, 17, 28, 43). The concept of essentiality has been used to distinguish chromosomes from plasmids. In many instances, this distinction is straightforward; i.e., plasmid loss results in loss of only a limited growth characteristic or ability, with the derivative being able to grow optimally under other growth conditions. Although many plasmids remain cryptic and difficult to cure, isolates of the same species or strain often show a high degree of variability as to plasmid content but not chromosomal profile.

From the standpoint of DNA size, either the 3,050-kb or the 914-kb circular DNA molecule of R. sphaeroides 2.4.1 is considered large enough to be a chromosomal DNA element, as described above, and neither is capable of spontaneous transfer. Additionally, these replicons have the following properties which make them more like bacterial chromosomes than plasmids. (i) Each carries critical housekeeping

genes, such as rrn (one on the large chromosome and two on the small chromosome) and tRNA genes, including f-met tRNA (8), hemA or hemT (41), cobA (36), gapB (39), and the transketolase gene (40), which have been found only in the "chromosome" of procaryotic organisms. Insertional inactivation of rrnB and rrnC or rrnA and rrnC genes resulted in an extremely slow growth rate of the mutants under all of the growth conditions attempted (8a). Inactivation of any one of the rrn operons yielded only a slightly diminished growth response under all of the growth conditions tested (8a). Thus, the presence of these genes appears to be important, at least to maintain normal growth in the laboratory under all of the conditions attempted. (ii) Each chromosome is present in a fixed stoichiometry relative to the others. (iii) All of the strains of R. sphaeroides examined by us have two chromosomes apparently similar in size to those observed for 2.4... (iv) On the other hand, all of the strains examined displayed extreme plasmid profile variability. As revealed here, each chromosome can behave genetically independently of the others. Thus, we are led to define a chromosome of a procaryotic organism as a DNA molecule of some "minimal" size which is non-self-transmissible and is a member of the genomic complement of an organism consisting of independent replicons, all of which are collectively necessary to provide optimal cell growth under all environmental conditions.

Here we have demonstrated how a combination of physical mapping techniques and pseudogenetics can be exploited to advance knowledge of the genomic structure of bacteria which are not yet considered in the mainstream of scientific development. As revealed here, schizotyping of recombinant organisms provides the necessary physical dimension to the development of a genetic system.

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