Curing the Megaplasmid pTT27 from *Thermus thermophilus* HB27 and Maintaining Exogenous Plasmids in the Plasmid-Free Strain

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Stepwise deletions in the only plasmid in *Thermus thermophilus* HB27, megaplasmid pTT27, showed that two distantly located loci were important for maintenance of the plasmid. One is a minimum replicon including one gene, repT, coding a replication initiator, and the other encodes subunits of class I ribonucleotide reductase (RNR) for deoxynucleoside triphosphate (dNTP) synthesis. Since the initiator protein, RepT, bound to direct repeats downstream from its own gene, it was speculated that a more-downstream A+T-rich region, which was critical for replication ability, could be unwound for replication initiation. On the other hand, the class I RNR is not necessarily essential for cell growth, as evidenced by the generation of the plasmid-free strain by the loss of pTT27. However, the plasmid-free strain culture has fewer viable cells than the wild-type culture, probably due to a dNTP pool imbalance in the cell. This is because of the introduction of the class I RNR genes or the supplementation of 5′-deoxyadenosylcobalamin, which stimulated class II RNR encoded in the chromosome, resolved the decrease in the number of viable cells in the plasmid-free strain. Likewise, these treatments dramatically enhanced the efficiency of transformation by exogenous plasmids and the stability of the plasmids in the strain. Therefore, the class I RNR would enable the stable maintenance of plasmids, including pTT27, as a result of genome replication normalized by reversing the dNTP pool imbalance. The generation of this plasmid-free strain with great natural competence and its analysis in regard to exogenous plasmid maintenance will expand the availability of HB27 for thermophilic cell factories.

*Thermus* spp. are extremely thermophilic bacteria that can grow at high temperatures from 50 to 82°C. Their thermophilic enzymes, e.g., DNA polymerase for PCR, have been used in industrial applications, and their potential as cell factories to produce enzymes from other thermophiles has been demonstrated (1). The bacteria themselves have also been extensively investigated as model organisms for systems biology including structural genomics (1, 2), and their genomic DNAs, with G+C contents as high as about 70%, are an interesting target for synthetic biology (3). *Thermus*-related studies have been encouraged by the existence of an established genetic engineering system based on natural competence (4–6), with thermostabilized drug resistance markers (7–10) and other thermostable genetic tools (1, 11–16).

Complete genomic sequences of *T. scotoductus* SA-01, *T. oshimai* JL-2, *Thermus* sp. strain CCB_US3_U1, *T. aquaticus* Y51MC23, *T. thermophilus* strains HB8, HB27, SG0.5P17-16, and JL-18, and others are available online. *Thermus* spp. often harbor a megaplasmid larger than 100 kbp, whereas in some strains, the megaplasmid genes seem to be transferred to the chromosome (17, 18). Similar megaplasmids have been observed in other bacteria of the *Thermus-Deinococcus* group showing resistance to extreme stresses such as high temperature, radiation, desiccation, and oxidation. It has not been reported whether these megaplasmids, carrying hundreds of genes, are essential for cell growth in the bacterial group. Furthermore, the replication initiation mechanism remains to be analyzed, in contrast to that of smaller *Thermus* plasmids such as pTT8 (9.3 kbp) (19, 20), a 16-kbp unnamed plasmid from strain ATCC 27737 (21), pTA103 (2.0 kbp) (22), and pVV8 (81 kbp) (23).

Megaplasmids are often found in bacteria such as *Alphaproteobacteria*, *Streptococcaceae*, and *Planctomycetaceae*. Among them, the replication origins of alphaproteobacterial megaplasmids have been the most extensively studied (24–26). The three contiguous genes repABC are responsible for plasmid replication and segregation, and repAB and repC encode partitioning proteins and an initiation factor, respectively (24–28). An A+T-rich region observed in and around the repABC genes was speculated to be unwound for replication initiation. In *Paracoccus versatilis*, belonging to the *Alphaproteobacteria*, the pTAV3 megaplasmid curing has been reported to lead to a growth defect in minimal medium, raising the question of whether the replicon is a plasmid or a minichromosome (25). Similarly, a large replicon, pNRC100, in a halophilic archaean has been reported to harbor essential genes, and the potential of the replicon to evolve into a new chromosome has been mentioned, based on its resistance to curing (29). On what criteria should the issue of megaplasmid versus minichromosome be judged?

pTT27, with a size of about 233 kbp, is the only plasmid in *T. thermophilus* HB27, and its complete sequence was the first released among the megaplasmids from *Thermus* spp. (30). *Thermus-Deinococcus* bacteria are usually pigmented yellow or red because of carotenoid production (31), and HB27 is no exception. The terminal steps of the synthesis of this yellow pigment are encoded on pTT27 (30). Likewise, pTT27 encodes terminal biosynthesis steps for cobalamin (30), which serves as a cofactor for cellular enzymes, as a ligand for DNA riboswitches in the control of gene expression (32), and as a light-sensing chromophore to...
TABLE 1 PCR primer sequences to amplify ca. 5-kbp regions for homologous recombination with pTT27

<table>
<thead>
<tr>
<th>Locus</th>
<th>Location on pTT27 (nt)</th>
<th>Length of region (bp)</th>
<th>Primer sequencea</th>
</tr>
</thead>
</table>
| 1     | 231441–3800            | 4,965                 | 5′-AACTGGCCGGTTGAG
|       |                        |                       | GAATTCTCATCTGAGCC|
| 2     | 59961–63394            | 5,434                 | 5′-AACTGGCCGGTTGC|
|       |                        |                       | AGTAATGTCGAGCT|
| 3     | 119941–125680          | 5,740                 | 5′-AACTGGCCGGTTG|
|       |                        |                       | AAGTAATGTCGAGC|
| 4     | 179931–185270          | 5,340                 | 5′-AACTGGCCGGTTG|
|       |                        |                       | AAGTAATGTCGAGC|

a The underlined, boldface, and italic sequences indicate NotI, EcoRI, and SalI sites, respectively.

MATERIALS AND METHODS

Cells, plasmids, and materials. *T. thermophilus* HB27 was described previously (36). The construction of an HB27 recA-null mutant and of plasmids pUC-Hm and pUC-YH1 was in which a hygromycin resistance (Hm') gene was cloned at the NdeI site of pUC19 and the replication origin of the intrinsic plasmid pVV8 from *T. thermophilus* H88 was inserted into pUC-Hm, which was performed as described previously (23). Plasmid pUC-TH8, in which the replication origin of pTT8 from H88 (19) was inserted into pUC-Hm, was constructed in this study. A kanamycin resistance (Km') gene (7) and the Hm' gene (9) for *T. thermophilus* were kindly donated by Seiki Kuramitsu (Osaka University) and Yoshinori Koyama (AIST), respectively, whereas a bleomycin resistance (Bm') gene (10) was chemically synthesized by TaKaRa Bio. *E. coli* Rosetta2(DE3) and the plasmid pET-22b were purchased from Merck Millipore, while plasmid pSTV29 for *E. coli* was from TaKaRa Bio. Oligonucleotides with a 6-carboxyfluorescein (FAM) label at the 5′ ends were synthesized by Eurofins Genomics K.K. 5′-Deoxyadenosylcobalamin (AdoCbl) (coenzyme [vitamin] B12) was purchased from Sigma-Aldrich.

Growth conditions and transformation. *T. thermophilus* was grown at 70°C in TR medium, and 1.5% gelan gum with 1.5 mM CaCl2 and 1.5 mM MgCl2 was added to the medium for the plates (6). Km at 500 μg/ml, Hm at 100 μg/ml, Bm at 20 μg/ml, and/or AdoCbl at 0.1 μg/ml was added to the medium when needed. Transformation was carried out as described previously (6). The recA-null mutant was grown at 58°C as described previously (37). Synthetic medium (38) was used as the minimal medium.

Initial deletion of pTT27 region in HB27. Fragments of approximately 5 kbp for recombination with pTT27 in *T. thermophilus* HB27 cells were amplified by PCR with the primers listed in Table 1 and La Taq with GC buffer (TaKaRa Bio). One fragment digested by EcoRI and NotI was inserted into an EcoRI-SalI region of pUC-Hm with another fragment digested by NotI and SalI (Fig. 1A, panel i). The resultant plasmid was linearized by NotI digestion and then transferred to HB27. When all colonies appeared white on the TR plate containing Hm, it was judged that the expected deletion of the pTT27 region had been achieved in the transformants. The deletion was checked by restriction enzyme digestion of the extracted plasmid in addition to colony PCR.

Two 5-kbp fragments were cloned into an EcoRI-SalI region of pSTV29 in a similar manner, and subsequently the Kn' gene was inserted into the NotI site between the fragments (Fig. 1A, panel ii). The resultant plasmid was linearized by XbaI and transferred to the pTT27-miniaturized HB27. The transformants were selected on a Kn-containing TR plate.

Stepwise deletions of pTT27 region in *E. coli*. The Kn'-labeled pTT27 derivative (pUC27H4) is a shuttle vector between *T. thermophilus* and *E. coli* with a size of about 40 kbp. Additional deletions of the pTT27 region were carried out by commonly used methods with *E. coli*. The replication ability of the reduced pTT27 regions was verified by transformation of the recA-null mutant of HB27 harboring the intact pTT27. When the region possesses replication ability in *T. thermophilus*, colonies on the selection plate containing Hm are white because of the loss of intact pTT27 resulting from plasmid incompatibility.

DNA preparation and gel electrophoresis. The alkali-SDS method was meticulously applied to the preparation of the megaplasmid and its derivatives from *T. thermophilus*, followed by ultracentrifugation in a CsCl-ethidium bromide gradient (39) especially for large-scale preparation. Intact genomic DNA was prepared in an agarose plug by a method described previously (40), followed by in-gel digestion with restriction enzymes. The plasmid or genomic DNA was analyzed (Fig. 1B, panel iv; see Fig. 3B) by contour-clamped homogeneous electric field (CHEF) gel electrophoresis in a 1.0% agarose gel in TBE buffer (50 mM Tris-borate [pH 8.0] and 1.0 mM EDTA) at 15°C. Gels were stained with ethidium bromide and visualized under UV light.

Curing of pTT27 from HB27. In plasmid pUC27H11, three genes (the ribonuclease reductase [RNR] α and β subunit genes and rep T) were cloned into pUC-Hm. When the Kn' gene replaces the Hm' gene on the

mediating light-dependent gene regulation (33). The closely related strain HB8 possesses a similar, 24-kb-larger megaplasmid with the same name, but synteny between the two pTT27 megaplasmids is less conserved than that between two chromosomes (34). Large plasmids, as described above, seem to be present at low copy numbers in each cell (24–26), while the copy number of pTT27 is estimated to be four or five per cell in HB8 (35). *T. thermophilus* is a polyplid, and the pTT27 copy number is equal to the chromosomal one (35). We focused on the sole plasmid pTT27 in HB27 to survey its replication origin and to determine which genes are essential for cell growth. If the plasmid had been smaller than 100 kb, allowing it to be easily handled for plasmid preparation and transformation, fragments of digested pTT27 would have been subcloned in *Escherichia coli* to identify the replication origin and essential genes, as performed previously in pVV8, which can be replicated by a repV gene alone (23). At the start of the present study, however, preparation of the intact high-purity pTT27 from HB27 was very difficult because of insufficient skill in manipulating the megaplasmid. Therefore, a strategy of pTT27 miniaturization in the HB27 cell was employed repeatedly until a size permitting easy handling was obtained.

In this study, a minimum replicon of the megaplasmid pTT27 was identified as a 2.3-kb region including a single gene encoding an initiator protein. During the process, a plasmid-free substrain of HB27 was generated as a result of pTT27 curing. This plasmid-free strain, with superior natural competence, has great potential as a host strain for DNA engineering and thermophilic cell factories. Findings on the stability of exogenous plasmids in the strain as presented here will contribute greatly to the development of DNA manipulation with *T. thermophilus* HB27 as well as its plasmid-free substrain.
plasmid, the plasmid is referred to as pUC27K11. *T. thermophilus* HB27 was initially transformed by pUC27H11. Subsequently, the obtained Hm-resistant white HB27 was transformed by pUC27K11, and transformants were selected in the presence of both Km and Hm. After a transformant resistant to both drugs was grown in TR medium in the absence of the antibiotics for 3 days, the cells were spread on an antibiotic-free TR plate. Among 300 colonies analyzed for drug resistance, 13 exhibited sensitivity to both drugs for 3 days, the cells were spread on an antibiotic-free TR plate. The plasmid pUC27H11 contains the *repT* gene and *RNR* genes on the pUC-Hm vector. A PvuII fragment containing the genes from the plasmid was digested by XhoI and NcoI, and the resultant fragment, was ligated into pUC-Hm as a homologous region for recombination (dotted lines) with pTT27. The plasmid constructed in *E. coli* was transferred to *T. thermophilus* HB27 after linearization by NotI digestion. The transformants were selected by Hm resistance from pUC-Hm. Black and white stars indicate the positions of the NdeI (5'-TTTTGCGGCCGCGCCGGGGA) and NotI (5'-TTTTTCATATGGCCCGCAAGCGCAAGCAGGACGCCACCCC) primer and 5'-GCCCTC-3'.

Preparation of RepT recombinant protein. The plasmid pET-RepT, for the overexpression of the C-terminally His-tagged RepT protein, was constructed by ligating a DNA fragment containing the repT gene to the Ndel-NotI site of pET-22b. The DNA fragment was amplified by PCR using the pTT27 plasmid as a template. The primer sequences for PCR were 5'-TTTTCTATATGCGCCGCAGCGCAAGCGCAAGCAGGACGCCACCCC' for the 5' primer and 5'-TTTTTCATATGGCCCGCAAGCGCAAGCAGGACGCCACCCC' for the 3' primer, where the underlined bases show the positions of the Ndel (5' primer) and NotI (3' primer) sites. Subsequently, plasmid construction, overproduction, and purification of the C-terminally His-tagged RepT fused to the maltose-binding protein (MBP) were performed as described in a previous report (23). To separate the His-tagged RepT from MBP, the purified proteins were treated with PreScission protease (GE Healthcare Bio-Sciences) and applied to a column of HisTrap HP (GE Healthcare Bio-Sciences) as described previously (23). The eluted proteins were concentrated, dialyzed against 20 mM Tris-HCl (pH 8.0) containing 1 mM EDTA, 0.5 M NaCl, and 3 mM β-mercaptoethanol, and used for further analyses. The protein concentration was determined by the Bradford assay and by measuring UV absorption using a theoretical molar extinction coefficient of 1.13, which was calculated from ε values of 1,576 M⁻¹ cm⁻¹ for Tyr and 5,225 M⁻¹ cm⁻¹ for Trp at 280 nm (41).

Gel retardation assay. The plasmid pUC27H11 carries repT and RNR genes on the pUC-Hm vector. A PvuII fragment containing the genes from the plasmid was digested by XhoI and Ncol, and the resultant fragment.

FIG 1: Large-scale deletion in megaplasmid pTT27. (A) Deletion approaches used in this study. Panel i, two PCR-amplified 5-kbp fragments, represented by the arrows, were inserted into pUC-Hm as a homologous region for recombination (dotted lines) with pTT27. The plasmid constructed in *E. coli* was transferred to *T. thermophilus* HB27 after linearization by NotI digestion. The transformants were selected by Hm resistance from pUC-Hm. Black and white stars indicate essential gene(s) and the replication origin, respectively. Panel ii, similar 5-kbp fragments were inserted into pSTV29 with Km as indicated. The resultant plasmid linearized by XbaI was transferred to the Hm-resistant HB27 shown in panel i. The transformants were selected in the presence of Km. (B) Actual large-scale deletions in pTT27. Panel i, deletion trials are schematically shown. The top line indicates the intact pTT27 of 232,605 bp, and the numbers above are positions on the megaplasmid. Red arrows 1, 2, 3, and 4 and blue arrows a, b, c, and d show the loci for homologous recombination (Table 1) and the target regions for colony PCR in panel iii, respectively. The lower lines represent the putative residual region after deletion, and the numbers at both ends display the end locations of the region. The replication ability of each region is shown by (+) or (−) on the left side, and the region exhibiting the replication ability, as shown by (+), is emphasized by a bold line. The leftmost bold numbers match the lane numbers in panel iv and Fig. 2C. Panel ii, colonies on the Hm-containing plate. The deletion trial of the region from locus 4 to locus 2 resulted in all colonies being white, owing to the lack of the carotenoid biosynthesis-related genes on pTT27. Panel iii, the existence or nonexistence of the four regions shown by blue arrows in panel i in the colonies of panel ii was checked by colony PCR with the primers listed in Table 2. The deletion from locus 4 to locus 2 was supported by the lack of amplification of regions a and d. Lane M, size marker. Panel iv, the intact pTT27 (lane 1) and pUC27H2 (lane 2) prepared from HB27 were digested by both NdeI and SphI, followed by CHEF gel electrophoresis. Lane M, size marker of concatemeric λ DNA and the HindIII-digested one. Running conditions were 3 V/cm with a 12-s pulse time and 16-h running time at 15°C.
TABLE 2 Primer sets for colony PCR to check deletions in pTT27

<table>
<thead>
<tr>
<th>Target region</th>
<th>Position on pTT27 (nt)</th>
<th>Amplified length (bp)</th>
<th>Primer sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>32391–33052</td>
<td>662</td>
<td>5'-GAGAGGACAGCCTGGGCGCATGGCTGGAGGAAGTCCCGG</td>
</tr>
<tr>
<td>b</td>
<td>92451–92950</td>
<td>490</td>
<td>5'-GAGAGGAGTTGGGAGGCAGCCTGGTCTCCACCGGAGTAC</td>
</tr>
<tr>
<td>c</td>
<td>162064–162504</td>
<td>441</td>
<td>5'-CCCTATGGGAAAGCCGCCACCTAGGTGCTCCACCTG</td>
</tr>
<tr>
<td>d</td>
<td>210037–210487</td>
<td>451</td>
<td>5'-CTGTTGCGCCACCAGGGGAGCTGTCCTCCCGGAGAAGAG</td>
</tr>
</tbody>
</table>

DNase I footprinting. A region from nucleotide (nt) 1 to 420 of a 699-bp XhoI fragment (see Fig. 6B) was amplified by PCR, for which one of the two primers was labeled with fluorescein at its 5' end, and used at 0.5 μM as a substrate for footprinting analysis. After the binding reaction in a total volume of 20 μl with 100 ng/μl poly(dC-dC)/poly(dC-dC) (Sigma) in exchange for λ DNA, the reaction mixture was incubated with DNase I (final concentration, 20 ng/μl) at 37°C for 30 min in 10 mM Tris-HCl (pH 8.0) containing 0.1 M NaCl, 1 mM diethiothreitol (DTT), 50 ng/μl acetylated bovine serum albumin (AcBSA), and 100 ng/μl λ DNA. The products were analyzed by ethidium bromide staining after electrophoresis with a 2.0% agarose gel. Substrates for Fig. 6C were amplified by PCR using the pUC27H11 plasmid as a template and used for the assay.

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**RESULTS**

Stepwise deletions in the pTT27 megaplasmid. Deletion of half of the pTT27 megaplasmid in *T. thermophilus* HB27 was attempted in order to identify the replication origin and the genes essential for cell growth. As shown in Fig. 1 and Table 1, four almost equally spaced loci (loci 1, 2, 3, and 4) on pTT27 were randomly selected. An approximately 5-kbp region in each locus was cloned for homologous recombination with the megaplasmid as described in Materials and Methods. Deletions of the regions from locus 3 to locus 1, from locus 4 to locus 2, from locus 1 to locus 3, and from locus 2 to locus 4 correspond to the remains of the regions from locus 1 to locus 3 (127 kb), locus 2 to locus 4 (125 kb), locus 3 to locus 1 (114 kb), and locus 4 to locus 2 (118 kb), respectively. These deletion trials were performed as shown in Fig. 1A, panel i. Although we had guessed that only the transformant in which the expected deletion was established would form colonies on the selection plate, colonies were observed in all four trials. In the case of the deletion from locus 3 to locus 1, all colonies on the plate were as yellow as those of the wild-type (WT) HB27 in Fig. 1B, panel ii. On the other hand, all of the colonies with the deletion from locus 4 to locus 2 were white. With the deletions from locus 1 to locus 3 and from locus 2 to locus 4, several colonies were white, but a large portion of the colonies were yellow. The yellow colonies link to the carotenoid biosynthesis-related genes on pTT27, and these genes are located upstream of locus 2 (Fig. 1B, panel i). This means that the deletions from locus 4 to locus 2 and from locus 1 to locus 3 must produce white colonies on the selection plate, and the result for the deletion from locus 4 to locus 2 suggests that the expected deletion was completed. The colonies from each deletion trial were exposed to colony PCR with the primers listed in Table 2 to check the deletion of the area sandwiched by the two 5-kb regions for homologous recombination. The results in Fig. 1B, panel iii, suggest that only the deletion from locus 4 to locus 2 could be established in a predictable way. This was supported by restriction enzyme digestion of the extracted plasmid (Fig. 1B, panel iv). As the largest two fragments from the intact pTT27 digested by both NdeI and SphI contain the region for homologous recombination, they were not observed in the plasmid with the deletion from locus 4 to locus 2. Instead, two truncated fragments, each of about 16 to 17 kb, would appear. In addition, three fragments of about 13 to 17 kb, as well as one 10.5-kbp fragment, one 3.3-kbp fragment, and two 2.3-kbp fragments, were lost in the deletion plasmid, but one 2.4-kbp fragment derived from pUC19 was added. The fragments around 13 to 17 kb were too intricate to assess, while the other changes were confirmed. Unlike the results for the deletion from locus 4 to locus 2, the expected digestion patterns were not observed in the other three deletion trials (data not shown). This halved pTT27 remaining from locus 2 to locus 4 is referred to as pUC27H2.

Why was colony formation observed in all of the deletion trials? It was speculated that homologous recombination between the transferred DNA and pTT27 occurred and that the expected deletion was completed. The deletion would have applied to some, but not all, copies of pTT27 because its copy number is four or five per cell (35). If the Hm⁺-labeled deleted version of pTT27 contained its replication factors, it would replace the intact pTT27 through plasmid incompatibility in the presence of Hm, as shown in the white colonies caused by the deletion of locus 4 to locus 2. If the Hm⁺-labeled deleted version possessed no replication factors, on the other hand, it would never replicate in a cell. However, the deleted one must be incorporated into the intact pTT27 by a homologous recombination in the remaining region somewhere, since it has been implied that recombination occurs with great frequency in *T. thermophilus* cells (35, 37). This is why the Hm⁺-resistant colonies were formed in all the deletion trials. Thus, the results of the colony PCR for the deletion from locus 4 to locus 2 were homogeneous, while those for the other deletions were heterogeneous, probably because of recombination at unspecific regions. These heterogeneous results were also shown in restriction enzyme digestion analysis of the extracted plasmids (data not shown). At least, the deletion of the region from locus 4 to locus 2 is established, and the four randomly selected loci fortunately encourage screening of the replication factors of pTT27 by the ap-
pearence of the white colonies due to the loss of the carotenoid biosynthesis genes.

Further deletion trials, as indicated in Fig. 1B, panel i, showed that the remainder of the region from nt 75911 to 170970 in pTT27 made all the colonies white. The resultant plasmid, as shown in lane 3 of Fig. 2C, is referred to as pUC27H3. When the pTT27 region of the plasmid was shortened at one end by about 4 to 5 kbp (Fig. 1B, panel i), most of the colonies were yellow. This suggests that both ends are important for pTT27 replication. Therefore, a center portion of the pTT27 region in pUC27H3 was replaced by the Km\(^{\prime}\) gene, as shown in Fig. 1A, panel ii, and B, panel i. The resultant smaller plasmid (pUC27H4) was confirmed (Fig. 2C, lane 4), showing that two loci positioned distantly on pTT27 were important for plasmid maintenance.

The pUC27H4 plasmid is a shuttle vector between *Thermus* and *E. coli* with a size of about 40 kbp (Fig. 2A), which can be easily manipulated in *E. coli*. When the *Thermus thermophilus* HB27 recA-null mutant, which formed yellow colonies, was transformed by pUC27H4, all transformants on the selection plate were white as a result of the loss of intact pTT27 resulting from plasmid incompatibility with pUC27H4. This implies that all transformants of the recA-null mutant made by the pUC27H4 derivative and exhibiting replication ability would form white colonies on the selection plate. Therefore, additional shortening of the region required for pTT27 replication was performed in *E. coli*, and its replication ability was checked with the HB27 recA-null mutant in a fashion similar to that for pUC27H4. As shown in Fig. 2A, deletions of the pTT27 region were repeated in a stepwise fashion; finally, pUC27H10, which exhibited replication ability, was obtained (Fig. 2C, lane 10). The plasmid possesses two pTT27 regions of about 2.6 and 3.6 kbp, in which the single open reading frame (ORF) TT_P0085 and the three ORFs TT_P0160, TT_P0161, and TT_P0162 are carried, respectively. To judge the necessity of the three operon-like genes TT_P0160, TT_P0161, and TT_P0162 for pTT27 replication, a null mutant plasmid for each gene was constructed to maintain the polycistronic transcription (Fig. 2B). When the recA-null mutant was transformed by these plasmids, the plasmid with the loss of TT_P0160 showed replication ability (Fig. 2C, lane 11), but that with the loss of TT_P0161 or TT_P0162 did not. These ORFs are registered in the database as encoding the β subunit of RNR (RNR-β) for TT_P0161 and the α subunit of RNR (RNR-α) for TT_P0162. RNR, which converts ribonucleotides to deoxyribonucleotides, is a key enzyme for the *de novo* synthesis of deoxyribonucleotides. The RNR complex consisting of α and β subunits seems to be important for pTT27 replication. On the other hand, TT_P0085 is registered as encoding a replication initiation protein. Here, the TT_P0085 protein and its gene are referred to as RepT, the replication initiation protein for pTT27, and the repT gene, respectively. The TT_P0160-deleted version of pUC27H10 (Fig. 2C, lane 11) is called pUC27H11.

**HB27 plasmid-free substrain.** In a previous study (36), we cured a 9.3-kbp plasmid, pTT8, in *T. thermophilus* HB8. In a similar way, pTT27 curing from HB27 was attempted in order to obtain a strain without a plasmid, as described in Materials and Methods and Fig. 3A. The absence of the plasmid in the obtained candidate strains was confirmed by the digestion of genomic DNA with both EcoT22I and Murn. There are four EcoT22I cleavage sites (but two sites are as close as 12 bases apart) and four MunI sites in the chromosome of HB27 and no EcoT22I sites and one MunI site in pTT27. As shown in Fig. 3B, the WT strain shows seven chromosome-derived bands and one for the linearized pTT27 with a size of 233 kbp, whereas the candidate lacks pTT27. For additional confirmation, the nonexistence of repT and RNR genes was checked by PCR (data not shown). One of the obtained
plasmid-free strains with white colonies, named PFW, was used after this step. Generation of the plasmid-free substrain means that all the megaplasmid-borne genes would be nonessential under this growth condition, and even the pTT27-borne RNR gene is no exception.

**Minimum replicon of pTT27.** As shown in Fig. 4, the megaplasmid-borne RNR genes were cloned in the chromosome of PFW. To determine whether the repT region is a replicon, a recA-null mutant of the resultant PFW I-RNR strain was transformed by pUC27H12, in which the RNR gene region of pUC27H11 was deleted and only the repT gene is present (Fig. 4). The transformants were obtained, and the plasmid from them was observed (Fig. 4, lane 12). The repT-containing region was found to be the true replication origin of pTT27, and it was shown that the RNR genes could serve in trans against the replicon. When the parental PFW was transformed by pUC27H12, a few colonies were formed on the selection plate with a remarkably low transformation efficiency of 2.0 × 10^9 CFU/μg, as shown in Table 3. However, the plasmid was not extracted from cultured cells of the colonies (data not shown). At this point, therefore, the RNR genes were thought to be an indispensable factor for pTT27 replication.

**Effect of AdoCbl on PFW growth.** The plasmid-free strain PFW grew in nutrient-rich TR medium, although its growth was clearly slower than that of the WT (Fig. 5A), showing that the megaplasmid was dispensable for growth in that medium. In synthetic medium, on the other hand, the PFW strain never grew, while the WT did (Fig. 5A), indicating the necessity of some plasmid factors for growth under this condition. Since the pTT27 megaplasmid carries cobalamin biosynthesis-related genes in the region from locus 1 to 2 (Fig. 1B, panel i) (30), AdoCbl at 0.1 μg/ml was added to the synthetic medium. This addition reversed the lack of growth of the PFW strain in the medium (Fig. 5A), confirming that the cobalamin biosynthesis genes on pTT27 were essential for the growth of *T. thermophilus* HB27 in the synthetic medium. In the TR medium, the addition of AdoCbl also improved the poor growth rates of PFW and PFW I-RNR but did not change that of the WT (Fig. 5A). When cultured in the TR medium with or without the AdoCbl supplementation, the number of viable cells at an optical density at 600 nm (OD600) of 0.15 was estimated by counting colonies formed on the TR plate. Although PFW had fewer viable cells than the WT, AdoCbl supplementation or the existence of pTT27-derived RNR seems to resolve the decrease in the number of viable cells in PFW (Fig. 5B).

**Stability of exogenous plasmid in the PFW strain.** The AdoCbl supplementation improved the poor growth rate of the PFW strain in TR medium (Fig. 5A). Therefore, transformation of the strain by pUC27H12 was tried by culturing in AdoCbl-supplemented TR medium. The AdoCbl supplementation encouraged colony formation, with much higher transformation efficiency than that without AdoCbl (Table 3), and the plasmid was observed in extract from the transformant cultured in the AdoCbl-supplemented liquid medium. When the same transformant was cultured without the AdoCbl supplementation, however, the plasmid was not observed (data not shown). These results suggest that the addition of AdoCbl would be required in order to maintain the pUC27H12 plasmid in the cell. When the above-mentioned few colonies obtained from the transformation without AdoCbl (Table 3) were actually cultured in the AdoCbl-supplemented liquid medium, the plasmid was observed, unlike the case in the normal medium (data not shown). In the WT and PFW I-RNR, on the...
other hand, the pUC27H12 plasmid was stably maintained without AdoCbl (data not shown), and further, the addition of AdoCbl had little effect on transformation efficiency (Table 3). The presence of the pTT27-originated RNR seems to have the same effect on plasmid maintenance and transformation as the AdoCbl supplementation.

As shown in Table 3, similar results were observed with other plasmids, pUC-Vmini and pUC-TT8, in which the minimum replicons of pV8 and pTT8 from *T. thermophilus* HB8 were each cloned in pUC-Hm. In these transformants of PFW, the plasmid also was stable and unstable with and without the AdoCbl supplementation, respectively (data not shown). The plasmid instability and the effect of AdoCbl on plasmid maintenance in PFW seem to be not specific to the pTT27 derivatives but rather to be applicable to other exogenous plasmids.

**RepT-binding site.** It was speculated that, like other replication initiation mechanisms, the replication of pTT27 would be initiated by the binding of some proteins as initiators to a specific region of the plasmid as the replicator. As the RepT protein, encoded by the *repT* gene was thought to be an initiator, the recombinant protein was prepared as described in Materials and Methods. The replicator must be located in the pTT27 region included in pUC27H12, but that in pUC27H11 was used for the substrate for the RepT-binding assay. In the gel retardation assay (Fig. 6A), the disappearance/retardation of the 699-bp XhoI fragment was observed with the increase of RepT. When the region downstream

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**FIG 4** RNR can function in pTT27 replication in trans. To test whether the RNR acts in the megaplasmid replication in trans, the transposase-encoding TT_C0665 gene on the chromosome of the PFW strain was replaced by the RNR genes with Bm’ for the selection marker, followed by *recA* deletion to avoid incorporation of pUC27H11 as a control plasmid into the chromosome through homologous recombination. Substitutions of all the chromosomes in the polyploid were confirmed by PCR and Southern blotting (data not shown). The resultant strain was transformed by pUC27H12, in which the RNR part of pUC27H11 (lane 11) was lost. The transformants were obtained in the presence of Hm, and the plasmid was confirmed from them as shown in lane 12. Arrowheads indicate the undigested plasmids. Lanes 11 and 13, pUC27H11 and pUC27H13, respectively, prepared in the same fashion; lane M, size marker of HindIII-digested λDNA.
from repT was deleted from the EcoRV site (Fig. 2A), its replication ability was lost. It is likely that the RepT binding to the site downstream from the repT gene is important for megaplasmid replication.

The sequence of the 699-bp fragment is shown in Fig. 6B. Two characteristics of the sequence are that nine direct repeats of 5'-TCGGAAGACTTGGGG-3' lie downstream from the repT gene and that the second to ninth repeats seem to appear at almost the same intervals. In addition, the sequence from nt 188 to 262 framed by the orange lines is just the same as that from nt 301 to 375. Further downstream from these repeats, an A+T-rich region with a G+C content of 32.7% is located over a wide range from nt 448 to 545, while the T. thermophilus HB27 genome shows a much higher G+C content of about 70% (30). To examine the effect of the repeats on RepT binding, additional gel retardation assays were performed. In the left gel of Fig. 6C, RepT shows sufficient binding to the first seven repeats but no binding to only the first three. On the right, on the other hand, the protein exhibits sufficient binding to the last six repeats but no binding to only the last two. Because the bindings are also observed significantly in the third and fourth repeats, no binding with RepT was detected (data not shown), supporting the interpretation of Fig. 6C that RepT requires multiple repeats for binding.

The DNA duplex of the A+T-rich region downstream from the direct repeats was speculated to be unwound when RepT was bound to the direct repeats. Therefore, the necessity of the A+T-rich region was confirmed in vivo. The region downstream from the repT gene was deleted to the 5' or 3' side of the A+T-rich region in pUC27H12 (nt 445 or 560 in Fig. 6B), and the resultant plasmids were transferred to the PFW I-RNR. The plasmid including the A+T-rich region (pUC27H13) resulted in colonies on the selection plate and was extracted from the culture (Fig. 4, lane 13), whereas the plasmid without that region caused no colony formation. Hence, the A+T-rich region is indispensable for pTT27 replication.

### DISCUSSION

Replication initiation of megaplasmid pTT27. The repT gene and its adjacent regions, as in pUC27H13, would be the true minimum replicon for megaplasmid pTT27. The replication initiation protein RepT binds to the direct repeat sequences of 5'-TCGGAAGACTTGGGG-3' downstream from its own gene repT, and the DNA duplex of the further-downstream A+T-rich region is likely to be unwound. Exclusion of the A+T-rich region from pUC27H13 caused a replication deficiency, strongly indicating the importance of the region. Eight repeats (the second to the ninth) are placed in a regular manner as shown in Fig. 6B, leading to the speculation that an oligomeric complex of the RepT proteins is coiled by the DNA duplex with the direct repeat sequences. Plasmid maintenance in PFW. Both the remarkably low efficiency of transformation of PFW by pUC27H12 (Table 3) and the lack of confirmation of the plasmid prepared from the resultant transformant culture had made us presume that the pTT27-borne RNR genes were indispensable for megaplasmid replication. However, this speculation was ruled out by the addition of AdoCbl, which caused the transformation efficiency to increase dramatically and encouraged stable maintenance of pUC27H12 in...
PFW (Table 3). Since RNR generally catalyzes the formation of deoxyribonucleotides from ribonucleotides, it is indispensable for DNA replication and repair and is essential for cell growth (42). The megaplasmid-borne RNR would be a class I enzyme using only NDP under aerobic conditions, while in *T. thermophilus* HB27, a probable class II RNR that uses both NDP and NTP as a substrate under aerobic and anaerobic conditions is encoded by TT_C1930 on the chromosome. Owing to the presence of another RNR, the absence of the megaplasmid-encoded enzyme in the PFW strain would not eliminate growth in TR medium. However, the decrease in viable cells in TR medium without AdoCbl (Fig. 5B) suggests the failure of some cellular mechanisms in PFW. The decrease in viable cells could be improved by AdoCbl supplementation or by the introduction of the class I RNR genes into the chromosome, as shown in Fig. 5B. These results suggest that AdoCbl would work on the class II RNR, which requires AdoCbl as a cofactor involved in radical generation (42). The PFW cells probably would be under dNTP-deficient conditions in the normal TR medium, in which the amount of AdoCbl might be insufficient to stimulate the class II RNR completely, because the PFW strain loses both the class I RNR and AdoCbl biosynthesis ability as a result of pTT27 curing. The dNTP shortage or the imbalanced dNTP pool in PFW would lead to noncompletion or infidelity of chromosome and plasmid replication. Chromosome replication especially is crucial to the survival of the cells. In addition, the polyploidy of *T. thermophilus* and the high G+C content of its genome might make the situation serious. Introduction of the class I RNR or stimulation of the class II enzyme by the addition of AdoCbl resolves the dNTP shortage or the imbalanced dNTP pool in a cell, and then the completion or improved fidelity of chromosomal replication would result in an increase of viable PFW cells (Fig. 5B). For the same reason, the addition of AdoCbl would enable the stable maintenance of pUC27H12 as well as of other exogenous plasmids in PFW, and consequently, transformation efficiency would be improved also (Table 3). Replication failure would occur equally in the chromosome and plasmid, but failure of chromosome replication must be directly linked to death. As a result, the population of cells lacking the plasmid would increase with growth. It is probable that in place of the AdoCbl supplementation, the presence of cobalamin biosynthesis-related genes would enable stable maintenance of the plasmid in PFW. Whereas our stepwise deletions in pTT27 showed that the combination of the repT and class I RNR genes was important in megaplasmid maintenance
combination of repT and the cobalamin biosynthesis-related genes can be also assumed.

Necessity of megaplasmid pTT27 for growth. First, it should be mentioned that the megaplasmid pTT27 must be highly beneficial for maintenance of genome stability in T. thermophilus HB27 because it carries the class I RNR genes and the biosynthesis genes for AdoCbl stimulating the chromosomal class II RNR. However, the megaplasmid also carries some toxin-antitoxin system genes, implying its stable maintenance in a cell. Because the miniaturized pTT27 derivatives such as pUC27H11 and pUC27K11 lack all the toxin-antitoxin systems, curing pTT27 from HB27 might be achieved rather easily in the way described in Materials and Methods. The PFW strain can grow in nutrient-rich TR medium (Fig. 5A), suggesting that all megaplasmid-borne genes would be nonessential for cell growth under this condition.

A growth defect in minimal medium was observed also when the pTAV3 megaplasmid was lost in P. versutus (25). It has been discussed that the megaplasmid required for growth in minimal medium might carry housekeeping genes and thus not fit the strict definition of a plasmid (25). It remains to be judged whether this kind of DNA replicon, which is required only for growth in minimal medium, is a megaplasmid or a minichromosome.

DNA engineering in T. thermophilus. In this study, an ~110-kb region on the megaplasmid was deleted by replacement by the drug resistance gene, as shown in Fig. 1B. This means that two regions with sufficient length for homologous recombination enable the replacement of an intervening part as long as 100 kb without a nonessential gene. The longer the prepared homologous regions for recombination, the broader the part in the genome might be substituted or deleted. Such a dynamic would introduce the possibility of genomic engineering in T. thermophilus. Incompatibility between the substituted and parental plasmids is implied to have priority over homologous recombination between them, according to a comparison of the results in Fig. 1B for...
the deletion from locus 4 to locus 2 and for the other three trials. In *T. thermophilus*, a polyploid bacterium, the nonessential genes on all genomic copies were replaced easily, whereas at least one copy had to remain intact in the case of the essential gene (35). The easy establishment of the deletion from locus 4 to locus 2 might have been caused by the same undefined mechanism that is involved in the replacement of the nonessential gene.

The PFW strain obtained in this study lost its pTT27-derived yellow pigment. This plasmid-free white strain with natural competence might have great potential as a host strain for DNA engineering in *T. thermophilus*. The absence of a plasmid in the host cell is attractive for plasmid transformation, and white colonies are suitable for color-based selection, such as yellow caused by carotenoid production (13) or blue caused by *X-Gal* (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) and β-galactosidase (15). In this aspect, the PFW I-RNR strain, in which the megaplasmid-originated class I RNR genes were carried in the chromosome, would be better for use as the host cells, because it improves growth and stably maintains exogenous plasmids (Fig. 5A). Understandably, the class I RNR genes can also be cloned in the plasmid vector, as in the case of pUC27H11 (Fig. 2B). Alternatively, it is possible to clone the pTT27-derived cobalamin biosynthesis-related genes into the chromosome of PFW, but AdoCbl supplementation to the medium will make it easy. Another advantage of PFW is its higher transformation efficiency than WT, as shown in Table 3. pTT27 encodes Argonaute protein, which acts as a barrier for the uptake and propagation of foreign DNA. Deletion of the Argonaute gene has been reported to increase the transformation efficiency, and the transformation efficiency of the deletion mutant is 10-fold higher than that of the WT (46). In addition, the PFW host enables the plasmid-borne genes to be transiently kept, as with using a conditionally replicable vector. When it is not needed, cancellation of the AdoCbl supplementation would lead to lack of the genes with the plasmid in PFW. This can make a significant contribution to experiments involving verification of an essential gene in PFW.

When PFW is the host strain, the repT-dependent plasmid can be used as a cloning vector. The pUC27H13 plasmid, in which a 2.3-kb region with only repT is contained as the minimum replicon of pTT27, is a compact *E. coli-Thermus* shuttle vector. The repT vector can be expected to be suitable for large-scale cloning, because it was the origin of the 233-kb megaplasmid. This pTT27 replicon study, as well as the generation of the plasmid-free strain and analysis of exogenous plasmid maintenance in the resultant PFW strain, will serve as a foundation for the next wave of large-scale *Thermus* genetic/genomic engineering research, in addition to ensuring the availability of the HB27 (PFW) strain for thermo-philic cell factories.

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