A novel system with positive selection for the chromosomal integration of replicative plasmid DNA in Corynebacterium glutamicum

Masato Ikeda¹ and Ryoichi Katsumata²

A simple system has been developed for generating Corynebacterium glutamicum strains containing stable replicative plasmids integrated into the chromosome via homologous recombination. The system is based upon extremely strong incompatibility between two plasmids, which cannot be co-maintained even under antibiotic selective pressure. Integration of the resident plasmid that contained the trpD gene of C. glutamicum was achieved by introduction of a second plasmid and subsequent selection for the maintenance of both plasmids. Plasmid integrates positive for both plasmid markers were obtained at a frequency about 10⁻³ of the normal transformation frequency with selection for the maintenance of only the second plasmid. Southern analysis revealed that the integration had occurred through a single-crossover homologous recombination between the trpD regions of the host genome and the plasmid. On the basis of the Campbell-type integration, chromosome walking was attempted by using Escherichia coli replication origins that were also present in the integrated plasmid. The chromosomal DNA was digested, ligated, and used to transform E. coli, which enabled recovery of the expected adjacent genomic DNA regions. The plasmid integrate was stably maintained for 30 generations under non-selective culture conditions, suggesting that the integrated sequences carrying a replicon active in the host were maintained as a stable chromosomal insert in C. glutamicum.

Keywords: Corynebacterium glutamicum, integration, plasmid incompatibility

INTRODUCTION

Non-pathogenic Gram-positive Corynebacterium glutamicum and related bacteria are industrially important micro-organisms, used in the production of various amino acids (Kinoshita & Nakayama, 1978). Amino-acid-producing strains of these bacteria have been constructed by random mutagenesis and screening procedures (Kinoshita & Nakayama, 1978). The development of gene cloning systems for these bacteria (Katsumata et al., 1984; Ozaki et al., 1984; Santamaria et al., 1984; Yoshihama et al., 1985) has enabled us to carry out further strain improvement. Several examples have already been reported of successful metabolic engineering of amino acid producers (Ikeda & Katsumata, 1992; Ikeda et al., 1993, 1994).

Plasmid-mediated amplification of the gene for the rate-limiting enzyme is a common strategy in metabolic engineering. Plasmids generally carry antibiotic-resistance genes, which allow for selection, but these genes are not always maintained in the absence of the selective pressure (Katsumata & Ikeda, 1993). In addition, overexpression of a plasmid-born gene can markedly inhibit the growth of the host cells owing to metabolic imbalance or accumulation of intermediates (Aiba et al., 1982; Archer et al., 1991; Sugiura et al., 1987). Thus the integration of genes into the chromosome rather than via recombinant plasmids may be more useful for generating production strains for industrial-scale fermentation processes.

In bacteria, the most straightforward approach to integration relies on a non-replicating plasmid which carries a homologous chromosomal DNA segment and an antibiotic-resistance marker (Gutterson & Koshland, 1983; Leenhouts et al., 1989). Introduction of such a plasmid yields transformants when a single crossover between the plasmid and the recipient chromosome occurs at the region of homology, which results in
integration of the entire plasmid flanked by copies of the homologous DNA. This Campbell-type integration event was first demonstrated in the corynebacteria by Schwarzer & Pühler (1991). They succeeded in selecting for plasmid integrates that resulted from highly efficient conjugal transfer from *Escherichia coli* to *C. glutamicum* of mobilizable *E. coli* vector plasmids carrying *C. glutamicum* DNA fragments. One limitation of this approach is the presence of strong restriction barriers in corynebacteria (Katsumata *et al.*, 1984; Smith *et al.*, 1986), although Reyes *et al.* (1991) and Vertès *et al.* (1993) described methods suitable for chromosomal integration in the highly restrictive corynebacteria.

Single-crossover integration of a plasmid after transformation generally occurs at a low frequency in the corynebacteria when the plasmid carries chromosomal sequences. To select for the plasmid integrates, we have devised a simple and efficient method based on extremely strong incompatibility between two plasmids. It is well known that two or more co-resident plasmids that have identical nucleotide sequences for all regions involved in replication and maintenance, which are referred to as isologous plasmids, are not stably inherited without external selection (Novick, 1987). However, preliminary studies described herein revealed that two vector plasmids which have the same pCG1 replication origin but differ in their antibiotic resistance markers could not be co-maintained in a *C. glutamicum* strain even under antibiotic selective pressure. This phenomenon prompted us to employ the two strictly incompatible plasmids to generate a *C. glutamicum* strain containing a replicative plasmid integrated into the chromosome via homologous recombination.

We report here the use of extremely strong incompatibility between the two plasmids to generate Campbell-type integration of a replicative plasmid with the trpD gene into the *C. glutamicum* genome. The stability of the recombinants and the facile recovery of adjacent genomic DNA regions containing the other tryptophan-biosynthetic genes is also reported.

**METHODS**

**Bacterial strains, plasmids and media.** The strains and plasmids used in this study are listed in Table 1. *C. glutamicum* TA2 was isolated by selection for tryptophan auxotrophy after mutagenesis of strain L22 with NTG. The genetic lesion was identified as trpD by both enzyme assay and complementation analysis with the intact trpD gene of the organism. Plasmids pCG1 and pCG2 are endogenous cryptic plasmids isolated from *C. glutamicum* ATCC 31808 and ATCC 31832, respectively, in our laboratories (Ozaki *et al.*, 1984). Vector plasmid pCG17 was constructed by ligation of the cryptic plasmid pCG2 and the Sp' and Sm' determinants from another endogenous plasmid, pCG4 (Katsumata *et al.*, 1984). For growth of *C. glutamicum*, complete medium BY (Katsumata *et al.*, 1984), minimal medium MM (Ozaki *et al.*, 1985) and enriched minimal medium MMYE (Katsumata *et al.*, 1984) were used. Solid plates were made by the addition of 1.6% (w/v) Bacto-Agar (Difco). RCGA medium (Katsumata *et al.*, 1984) containing tetracyphon (50 µg ml⁻¹) was used for regeneration of *C. glutamicum* protoplasts. When required, supplements or antibiotics were added at the following final concentrations (µg ml⁻¹): spectinomycin (Sp), 400 for RCGA plates or 100 for BY plates; kanamycin (Km), 200 for RCGA plates or 20 for BY plates; chloramphenicol (Cm), 5 for BY plates; tetracycline (Tc), 2.5 for BY plates; streptomycin (Sm), 20 for BY plates; tetracyphon, 50 for MM plates and MMYE medium. For growth of *E. coli*, LB medium (Sambrook *et al.*, 1989) was used. For selecting transformants and examining their phenotypes, Km, Cm or Tc was added at a final concentration of 20 µg ml⁻¹ for LB plates.

**Preparation and manipulation of DNA.** Plasmid DNA of *C. glutamicum* was isolated by the alkaline lysis method (Sambrook *et al.*, 1989) and, if necessary, purified by CsCl/ethidium bromide equilibrium density-gradient centrifugation (Katsumata *et al.*, 1984). Chromosomal DNA was extracted from protoplasts of *C. glutamicum* strains by the method of Saito & Miura (1963). The protoplasts were prepared as described previously (Katsumata *et al.*, 1984). Because of their lysozyme sensitivity, cells could be easily converted to protoplasts by lysozyme digestion without prior exposure to penicillin during growth. Preparation of plasmid DNA from *E. coli*, DNA digestion and ligation were carried out by standard procedures (Sambrook *et al.*, 1989). Restriction enzymes and T4 DNA ligase were obtained from Takara Shuzo Co.

**Transformations.** Transformation procedures used were the protoplast method (Katsumata *et al.*, 1984) for *C. glutamicum* and the CaCl₂ method (Sambrook *et al.*, 1989) for *E. coli*.

**Southern hybridization.** Digested chromosomal DNA was electrophoresed and transferred onto nitrocellulose membranes (Amersham), as described by Sambrook *et al.* (1989). Plasmid pCA45 was used as a hybridization probe; it was labelled to high specific activity with [α-³²P]dCTP by the nick-translation method of Smith *et al.* (1977). The membranes were prehybridized for at least 2 h at 65 °C and then hybridized overnight with the labelled probe at the same temperature. After being washed, the membranes were autoradiographed.

**Determination of antibiotic resistance.** *C. glutamicum* strains INT9 and TA2 carrying pCA45 were grown in 3 ml BY medium supplemented with Km. After appropriate dilutions of the culture broth, an aliquot (10⁴ cells) was spread on BY plates containing various concentrations of the indicated antibiotics. Growth was assessed after 42 h incubation at 30 °C.

**Stability of transformants.** Seed cultures of *C. glutamicum* strains INT9 and TA2 carrying pCA45 grown in BYG medium (BY medium containing 10% glucose) were diluted in fresh BYG medium and grown for 30 generations aerobically at 30 °C. No differences in growth rate between the prototrophic strains and the plasmid-less tryptophan auxotroph were observed under the conditions used, confirming that there was no selection for the trpD gene in its mutant. After every ten generations, an aliquot was withdrawn from the culture and appropriate dilutions were spread on BY plates. After incubation at 30 °C for 2 d, 100 colonies from each sample were tested for their antibiotic-resistance phenotype.
Table 1. Bacterial strains and plasmids

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Relevant characteristics</th>
<th>Source or reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C. glutamicum L22</td>
<td>Lysozyme-sensitive mutant of wild-type ATCC 31833; no indigenous plasmids</td>
<td>Katsumata et al. (1987)</td>
</tr>
<tr>
<td>C. glutamicum TA2</td>
<td>trpD mutant of L22</td>
<td>This study</td>
</tr>
<tr>
<td>C. glutamicum INT9</td>
<td>TA2 containing pCA45 integrated into the chromosome supE44 hsdR endA1 pro thi</td>
<td>This study</td>
</tr>
<tr>
<td>E. coli MM294</td>
<td></td>
<td>Sambrook et al. (1989)</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pCE53</td>
<td>pCG1-ori, pACYC177-ori, Km' Cm' Tc'</td>
<td>Ozaki et al. (1984)</td>
</tr>
<tr>
<td>pCG11</td>
<td>pCG1-ori, Sp' Sm'</td>
<td>Katsumata et al. (1987)</td>
</tr>
<tr>
<td>pCG17</td>
<td>pCG2-ori, Sp' Sm'</td>
<td>This study</td>
</tr>
<tr>
<td>pCA45</td>
<td>pCE53 derivative containing a 2-1 kb trpD fragment from C. glutamicum KY10894, Km' Cm' Tc'</td>
<td>Katsumata &amp; Ikeda (1993); see Fig. 1</td>
</tr>
</tbody>
</table>

RESULTS AND DISCUSSION

Incompatibility between C. glutamicum plasmid vectors

Plasmid incompatibility was examined in C. glutamicum strain L22 using three multicopy vectors, pCE53 (Km' Cm' Tc'), pCG11 (Sp' Sm') and pCG17 (Sp' Sm'). The former two have an identical pCG1 replication origin and the last has a pCG2 origin. Protoplasts of strain L22 carrying pCE53 were transformed with pCG11 or pCG17 and selected for Sp'. Regenerated Sp' colonies were tested for the presence of Km'. None of the pCG11-carrying colonies checked had the Km' phenotype, whereas about 80% of the pCG17-carrying colonies were Km', indicating that pCE53 could be stably co-maintained with pCG17, but not with pCG11. Hence, the isologous plasmids pCE53 and pCG11 were confirmed to be incompatible as predicted. Interestingly, double selection by Km' and Sp' after transformation of the pCE53 carrier with pCG11 gave no transformants, whereas with pCG17 the frequency was 2 x 10^6 per 10^9 cells. This inability of pCE53 and pCG11 to co-exist even under antibiotic selective pressure indicated strong incompatibility between the two plasmids.

Isolation of transformants possessing all selectable markers of two strictly incompatible plasmids

When strain TA2, a trpD mutant of L22, carrying pCE53 was transformed with pCG11 (100 ng DNA), 5 x 10^5 transformants were obtained upon selection for only Sp'. However, as with the experiments with strain L22, no transformants appeared under selection for Km' and Sp'. In contrast, transformation of strain TA2 carrying pCA45, a pCE53-derived plasmid that contained a 2-1 kb fragment with the intact trpD gene of C. glutamicum, with pCG11 gave about 10^6 Km' Sp' transformants, corresponding to about 10^-8 of the frequency for Sp' transformants. Similar results were obtained from two repeat experiments, confirming the reproducibility of the transformation. Ten randomly chosen clones from these Km' Sp' transformants were purified by single-colony isolations and their phenotypes were examined. All clones tested showed a phenotype of Km' Cm' Tc', and prototrophy as well as Sp' Sm', which indicated that they possessed all selectable markers of both plasmids. However, restriction cleavage analysis of the plasmids isolated from these clones allowed the detection of only pCG11 as a free plasmid, suggesting that the other plasmid, pCA45, had integrated into the host chromosome. Three such clones, chosen randomly, designated INT strains, were used for further analysis.

Phenotypic and molecular analyses of INT strains

If plasmid pCA45 was being integrated into the genome of INT strains in a Campbell-type manner, the resistance of cells to the plasmid-encoded antibiotics would be predicted to be lower in the putative single-copy plasmid integrates than in the multi-copy plasmid carrier. To examine this possibility, the MICs of Km, Cm and Tc were determined as described in Methods.}

<table>
<thead>
<tr>
<th>Strain</th>
<th>MIC (µg ml^-1)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Km</td>
<td>Cm</td>
<td>Tc</td>
<td></td>
</tr>
<tr>
<td>TA2(pCA45)</td>
<td>&gt; 800</td>
<td>50</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>INT9</td>
<td>200</td>
<td>10</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>TA2</td>
<td>0.4</td>
<td>1.6</td>
<td>0.8</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Antibiotic resistances of strains INT9, TA2 and TA2(pCA45)
Fig. 1. Schematic representation of the relevant regions of the chromosomes of strains TA2 and INT9 after Campbell-type integration of plasmid pCA45. □, Cloned 2.1 kb trpD fragment from C. glutamicum; ◇, pCG1-ori; ▲, pACYC177-ori; △, pACYC184-ori. Restriction sites: B, BamHI; E, EcoRI; P, PstI.

Fig. 2. Southern hybridization analysis of chromosomal DNAs from strains TA2 and INT9. (a) DNA samples on a 0.8% agarose gel. Lanes: λ, lambda HindIII size markers; A and B, genomic DNA from strain TA2 digested with BamHI and EcoRI, respectively; C and D, genomic DNA from strain INT9 digested with BamHI and EcoRI, respectively. (b) Autoradiogram of a Southern blot of (a) (lanes A to D), hybridized with 32P-labelled pCA45. Numbers (1–7) in (b) indicate hybridizing bands. Sizes of bands 3–7 are indicated on the right.

To verify the occurrence of a single crossover event and confirm the chromosomal structure of the plasmid integrate, genomic digests were analysed by Southern hybridization using 32P-labelled pCA45 as a probe. Probing genomic digests from the non-transformed host strain TA2 should give signals with sizes corresponding to those expected from the restriction cleavage map of the C. glutamicum tryptophan-biosynthetic gene cluster (Katsumata & Ikeda, 1993) depicted schematically in Fig. 1. After Campbell-type integration, these hybridizing signals should be absent and followed by a shift to new signals in genomic digests from the plasmid integrate INT9 (Fig. 1). The results are presented in Fig. 2. As expected, the DNA from TA2 gave a unique band of 11.0 kb when digested with BamHI (Fig. 2b, lane A, band 1) and a 5.0 kb band when digested with EcoRI (Fig. 2b, lane B, band 2). When genomic digests from INT9 were probed, band 1 disappeared in the BamHI digests and two hybridizing bands, of 15.3 kb and 8.7 kb, were detected instead (Fig. 2b, lane C, bands 3 and 4). In addition, in the EcoRI digests, band 2 was replaced by three hybridizing bands, of 7.3, 6.5 and 4.2 kb (Fig. 2b, lane D, bands 5, 6 and 7). The sizes of these five new bands (3–7) were in agreement with those expected after Campbell-type integration of pCA45 through a single-crossover homologous recombination between the trpD regions of the host genome and the plasmid. A similar analysis of two other INT strains revealed that in these strains also the plasmids were integrated via a Campbell-type mechanism (data not shown).

**Recovery of chromosomal regions flanking the integrated pCA45**

As shown in Fig. 1, the integrated vector flanked by a duplication of the trpD region contains two kinds of *E. coli* replication origins. Therefore, it should be possible to clone other tryptophan-biosynthetic genes into *E. coli* if the adjacent genomic DNA regions are rescued as circular molecules so that they contain either origin as well as one of the selectable markers through digestion with an appropriate restriction enzyme and subsequent self-ligation. To examine this possibility, *BamHI* was chosen as the restriction enzyme. Chromosomal DNA of strain INT9 was completely digested with *BamHI* and treated with T4 DNA ligase at high dilution to maximize intramolecular ligation and therefore circularization of each fragment. The ligation mixture was then used to transform competent cells of *E. coli* MM294. Transformants were selected for Cmr or KmS and their phenotypes determined. As expected from the chromosomal structure depicted in Fig. 3(a), the former and the latter transformants showed the Cmr KmS TcS phenotype and the KmS CmS TcS phenotype, respectively. Restriction cleavage analysis (Fig. 3b) of each plasmid isolated from the former and the latter transformants and designated pEX1 and pEX2, respectively, yielded the same restriction cleavage map as that depicted in Fig. 3(a), indicating that pEX1 and pEX2 contained the trpCBA genes and the trpE gene, respectively, in addition to the trpD gene. These results suggest that chromosomal sequences flanking any cloned genes of *C. glutamicum* could be obtained as plasmids by using the same procedure as described here. These plasmids can be physiologically analysed to give restriction maps of the flanking regions and then be used as a source of DNA fragments for subsequent cycles of cloning. Thus, this method should provide alternative tools not only for integrating replicative plasmids but for 'chromosome walking', which is useful for genomic studies in *C. glutamicum*.

**Stability of the chromosomal insert**

The segregational stability of the antibiotic-resistance phenotype was examined under non-selective culture conditions in both INT9 and TA2 carrying pCA45. The
Plasmid integration in the *C. glutamicum* genome

**Fig. 3.** Restriction cleavage analysis of excision plasmids pEX1 and pEX2. (a) Predicted structure of genomic DNA regions yielding pEX1 and pEX2 after digestion of chromosomal DNA of strain INT9 with BamHI. Symbols are as described in the legend to Fig. 1. Restriction sites: B, BamHI; Bg, BgIII; E, EcoRI; S, Sall. (b) Restriction cleavage patterns of pEX1 (lanes 1-4) and pEX2 (lanes 5-8) on a 0.8% agarose gel. Each plasmid was digested with BamHI (lanes 1 and 5), EcoRI (lanes 2 and 6), Sall (lanes 3 and 7) or BgIII (lanes 4 and 8). The numbers on the right indicate the sizes of λ DNA digested with HindIII. Faint bands visible were due to incomplete digestion of DNA. In lane 2, the smallest (approx. 0.7 kb) fragment is not visible, although it was present.

Concluding remarks

Some research groups have used plasmid incompatibility to facilitate plasmid integration or gene replacement in bacteria, including *E. coli* (Simon *et al.*, 1983) and *Pasteurella haemolytica* (Fedorova & Highlander, 1997). In these studies, a rare recombination event was promoted through prolonged cultivation of cells where two incompatible plasmids were forced to co-exist. However, the last step was based on negative selection, because transformants that had been selected for both plasmid markers still consisted mostly of cells with two incompatible plasmids co-existing. In contrast, the system described here is based on a positive selection. To our knowledge, this is the first report of strong incompatibility being used to positively select transformants containing replicative plasmid DNA integrated into prokaryotic genomes. This system is efficient and thus should be a useful addition to the tools used in genetic engineering of the industrially important corynebacteria.

ACKNOWLEDGEMENTS

We are grateful to Yoshie Kaneko for her excellent technical assistance.

REFERENCES


Received 12 December 1997; revised 24 February 1998; accepted 20 March 1998.