Cloning Vectors Derived from Plasmids and Phage of Bacillus

JÜRGEN KREFT* AND COLIN HUGHES*

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1 Introduction

Bacillus subtilis is a well-characterized, gram-positive, non-pathogenic, spore-forming soil bacterium which produces a wide array of extracellular enzymes (for reviews see Young 1980; Priest 1977; Henner and Hoch 1980). The development of molecular cloning systems within this organism will not only greatly assist biochemical and genetic studies but should play a fundamental role in the further development of biotechnological processes based on the Bacilli.

As described by *Ehrlich* (this volume), following the realization that antibiotic resistance plasmids from *Staphylococcus aureus* could be transformed into *B. subtilis* (*Ehrlich* 1977), much effort has been devoted to developing *S. aureus* plasmid cloning vectors for the *Bacilli*. In this review we will describe the current status of vectors constructed from plasmids and phage indigenous to the *Bacilli*.

In the wake of the development of recombinant DNA techniques in *E. coli*, studies of *Bacillus* plasmids gave way to searches aimed at isolating potential cloning vectors. Extrachromosomal DNA in *Bacillus* was first demonstrated in *B. megaterium* and since then many reports have been made of plasmids in this species (*Carlton* and *Smith* 1974; *Rostas* et al. 1980), *B. subtilis* (*Lovett* and *Bramucci* 1975; *Tanaka* and *Koshikawa* 1977; *Bernhard* et al. 1978; *Uozumi* et al. 1980), and *B. pumilus* (*Lovett* et al. 1976).

^{*} Institut für Genetik und Mikrobiologie der Universität, Röntgenring 11, D-8700 Würzburg, West Germany

Table 1. Bacillus plasmids developed as cloning vehicles

Plasmid	Source	Mol. wt. (x 10 ⁻⁶)		Restriction sites	Marker	Reference
pLS 28	B. subtilis (natto)	4.1	5	EcoRI(2), BamHI(1), HindIII(5)	_	Tanaka and Koshikawa 1977
pBS 1	B. subtilis	5.5	6	EcoRI(1), BamHI(1), SaII(1), PsII(3), HindIII(6)	-	Bernhard et al. 1978
<i>pBC</i> 16	B. cereus	3.0	20	EcoRI(2), BamHI(1)	Tc^{R}	Bernhard et al. 1978
<i>pAB</i> 124	B. stearo- thermophilus	2.9	?	EcoRI(3), HindIII(2), single sites for BstEII, CauI, HpaI, XbaI	Tc ^R	Bingham et al. 1979

Numbers in parentheses indicate the number of restriction sites for this enzyme

However, the majority of these plasmids, in most cases isolated from culture collection strains, lack readily identifiable markers and are thus not of immediate use as cloning vehicles. Nevertheless two such cryptic plasmids, pBSI and pLS28, have been developed further. Examination of environmental isolates has proved more successful. Bernhard et al. (1978) isolated the tetracycline resistance (Tc^R) plasmid pBCl6 from a strain of B. cereus found in soil, and the antibiotic-resistant thermophiles discovered in river sludge and silage yielded the two Tc^R plasmids pAB124 and pAB128 (Bingham et al. 1979).

The following section describes the development of some of these plasmids (Table 1) as cloning vehicles.

2 Development and Use of Vector Plasmids

2.1 Vectors Capable of Replication Only in B. subtilis

The tetracycline resistance plasmid pBCl6 isolated from B. cereus can be transformed into B. subtilis (Bernhard et al. 1978) in which it replicates quite stably with no detectable segregation after more than 100 generations without selective pressure. It contains two EcoRI sites and in order to determine if these sites were within the tetracycline resistance gene(s) pBC16 was partially and completely digested with EcoRI and ligated with EcoRI linearized pBSI, a cryptic plasmid isolated from B. subtilis.

None of the Tc^R colonies obtained after transformation of competent cells of B. subtilis 168 with the ligated mixture contained a complete hybrid of the two parental plasmids but several derivatives were isolated (Fig. 1), pBC16-1, pBS161, and pBS161-1 being of particular interest (Kreft et al. 1978). The plasmid pBC16-1 is the circularized large Eco-RI fragment (mol. wt. 1.8×10^6) of pBCl6 which obviously carries both the replication functions and the tetracycline resistance determinant.

pBS161 and pBS162 have been found together in a large number of tetracycline-resistant colonies, the former plasmid alone carrying a TcR determinant.

Recircularization in vitro of the largest *HindIII* fragment of pBS161 yielded pBS161-1,

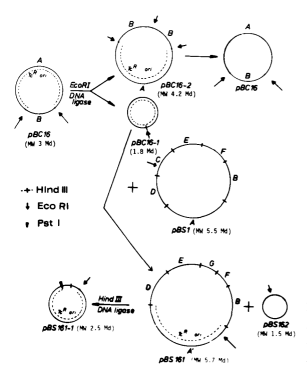


Fig. 1. Genealogy of Bacillus plasmids derived from pBC16 and pBS1. With the exception of pBS161-1 only the restriction sites for EcoRI and HindIII are shown

a small Tc^R plasmid (mol. wt. 2.5×10^6) with a high copy number (ca. 20), and single sites for HindIII, EcoRI, and PstI, none of which lies within the replication region or the tetracycline resistance determinant.

The recombinant plasmid RSF2124-B. leu (Nagahari and Sakaguchi 1978), contains the leucine A, B, and C genes of B. subtilis and can transform not only leu-E. coli but also B. subtilis to prototrophy. This plasmid is able to replicate only in E. coli. After digestion with EcoRI, ligation to EcoRI-cleaved B. subtilis (natto) plasmid pLS28 (Table 1), and transformation of B. subtilis RM125 (leu-, recE4), leu+ transformants yielded two recombinant plasmids pLS101 (mol. wt. 6.5×10^6) and pLS102 (mol. wt. 10.7×10^6). After subcultivation of these clones slowly growing colonies have been observed containing only the plasmid pLS103, which is indistinguishable from pLS101 (Fig. 2).

Insertion of foreign DNA into the single BamHI site inactivates leuA but not leuC, which can thus be used as a marker (Tanaka and Sakaguchi 1978). A derivative of pLS103 termed pLL10, has only one EcoRI site and complements leuA and leuB but not, in contrast to pLS103, leuC. In order to see whether DNA insertion into the remaining EcoRI site inactivated the leu function and also to introduce another marker, an EcoRI fragment carrying a B. subtilis 168 trimethoprim resistance determinant was recloned from pBR322-Tmp^R into pLL10 and transformed into B. subtilis ML112. leu+ Tmp^R clones yielded pTL10 (Fig. 3), a plasmid of mol. wt. 9.4×10^6 giving two fragments of 5.7×10^6 and 3.7×10^6 after EcoRI digestion. The latter fragment could convert B. subtilis to Tmp^R when inserted in both orientations, indicating its retention of the promoter (Tanaka and Kawano 1980).

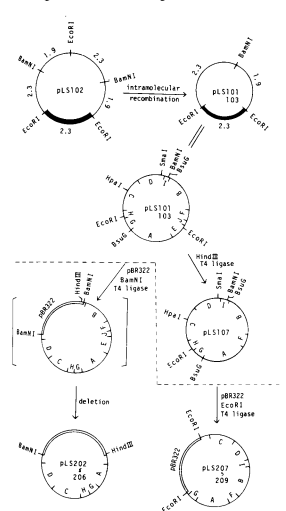


Fig. 2. Structure of constructed plasmids pLS101 etc. Numbers denote the molecular weights (\times 10⁻⁶) of the DNA fragments. The *thick* and *thin lines* indicate the vector and the DNA segments containing the leucine gene respectively. Cleavage sites of EcoRI, BamNI, SmaI, and BsuG are shown inside the circles and those of HindIII outside. Courtesy of T. Tanaka and K. Sakaguchi

In order to reduce both the size and number of EcoRI and Bg/II sites on pTL10, the derivative pTL12 has been constructed (Fig. 3); a leu^+ Tmp^R plasmid of mol. wt. 6.4×10^6 carrying single sites for EcoRI, Bg/II, BamHI, and XmaI. leu inactivation occurs following insertion at the BamHI and XmaI sites, and BamHI cleavage leaves a cohesive end (GATC), making possible the use of Bg/II, Bc/II, and MboI, which also leave this sequence. The presence of Tmp^R as a marker allows direct selection of transformants.

The Tc^R plasmid pAB124 isolated from B. stearothermophilus has three EcoRI sites (Table 1). The circularized EcoRI-A fragment (pAB224) (Bingham et al. 1980) is capable of autonomous replication and carries the tetracycline resistance determinant. It contains single sites for seven restriction enzymes, three of which produce cohesive termini. pAB524 has only one EcoRI fragment of pAB124 deleted (Fig. 4). Table 2 summarizes the properties of the plasmids described in this section. With the exception of pTL10 and pTL12, all these plasmids carry only one easily detectable genetic marker and do not allow

Table 2. Vectors derived from plasmic	ls listed in Table 1
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Vector	Source	Mol. wt. (x 10 ⁻⁶)	Single restriction sites	Markers
pBC 16-1	pBC 16	1.8	<i>Eco</i> RI	TcR
pBS 161-1	pBC 16/pBS 1	2.5	EcoRI, HindIII, PstI	$\mathrm{Tc}^{\mathbf{R}}$
pLS 103	<i>pLS</i> 28/ RSF 2124-B. leu	6.5	BamHI, SmaI, HpaI, XmaI	leu
pLL 10	pLS 103	5.7	EcoRI, BamHI, Bg/I, XmaI	leu
pTL 10	pLL 10	9.4	BamHI, BglI, XmaI	leu, Tmp ^R
pTL 12	pTL 10	6.4	BamHI, EcoRI, Bg/II, XmaI	leu, Tmp ^R Tc ^R
pAB 224	pAB 124	1.95	EcoRI, HpaI, HpaII, HhaI, ThaI, CauII, BstEII	TeŘ
pAB 524	pAB 124	2.3	HpaI, HpaII, CauII, BstEII	Tc^{R}

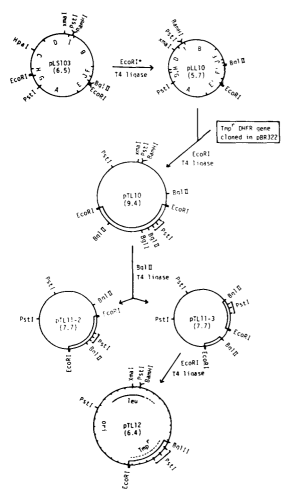


Fig. 3. Structure of plasmids pLS103 to pTL12. Numbers in parentheses are molecular weights. HindIII sites of pLS103/pLL10 are shown inside the circles and these were preserved in pTL10. Courtesy of T. Tanaka and N. Kawano

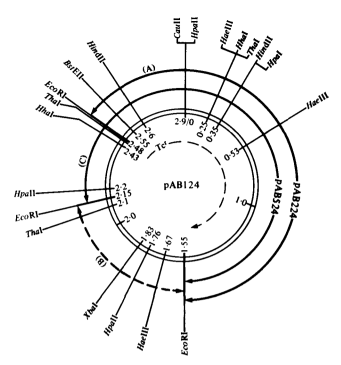


Fig. 4. Restriction endonuclease cleavage maps of pAB124, pAB224, and pAB524. Courtesy of A.H.A. Bingham, C.J. Bruton, and T. Atkinson

identification of recombinant molecules by insertional inactivation. They are, therefore, of limited value as vectors for molecular cloning.

2.2 Hybrid Vectors Capable of Replication in E. coli and B. subtilis

The construction of hybrid vectors attempts to combine the advantages of the well-defined *E. coli* cloning systems with those of the *Bacillus* host.

Hybrid replicons comprising *E. coli* vector plasmids and antibiotic resistance plasmids from *Staphylococcus aureus* have been described by *Ehrlich* (this volume). We have also constructed several *S. aureus/E. coli* hybrid plasmids of this type (pJK310, pJK312, pJK321, pJK521, and pJK523) (*Goebel* et al. 1979; *Kreft* and *Goebel*, manuscript in preparation). Two of them, pJK310 (pUB110 + pBR325) (*Gryczan* et al. 1978; *Bolivar* 1978) and pJK523 (pC221 + pBR322) (*Novick* 1976; *Bolivar* et al. 1977), express resistance to two antibiotics in *B. subtilis* and carry single restriction sites in these markers.

In addition we have developed hybrid replicons consisting of the *E. coli* vectors *pBR*322 (*Bolivar* et al. 1977), *pACYC*184 (*Chang* and *Cohen* 1978) and the *Bacillus* plasmids *pBS*161-1 and *pBS*1 (*Kreft* et al. 1978; *Goebel* et al. 1979).

pJK3 and pJK3-1 have been constructed by ligation of HindIII-cleaved pBR322 and pBS161-1. From the resulting complete hybrid pJK3 several duplex restriction sites have been removed by religation of PstI cleaved pJK3, thus yielding pJK3-1 (Fig. 5). This plas-

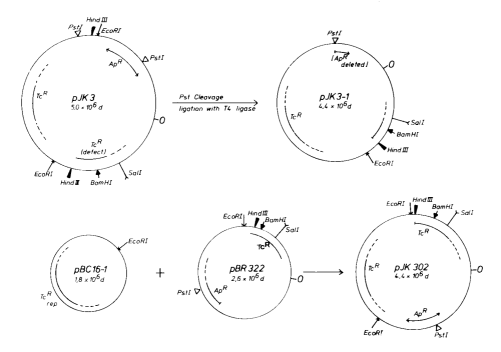


Fig. 5. Construction and restriction endonuclease cleavage maps of pJK3-1 and pJK302. Construction of pJK3 is described in the text

mid has retained only the tetracycline resistance determinant, but has single sites for five restriction enzymes and is capable of replication in both E. coli and B. subtilis. After transformation of E. coli with a ligation mixture from HindIII-cleaved pACYC184 and pBS161-1, the largely deleted hybrid plasmid pJK201, which carries Cm^R and Tc^R determinants, has been obtained (Goebel et al. 1979).

pJK302 is a hybrid consisting of EcoRI cleaved pBR322 and pBC16-1. It has single sites for four restriction enzymes, the PstI site being situated in the Ap^R gene. Cleavage with EcoRI of both parental plasmids does not inactivate the Tc^R determinants on these; the hybrid pJK302 expresses a high level (more than 100 µg/ml) in both E. coli and B. subtilis (Fig. 5).

Ligation of the EcoRI-cleaved plasmids pBR322 and pBSI yields in E. coli the expected complete hybrid pJK501. After transformation of competent cells or protoplasts of B. subtilis tetracycline-resistant colonies yield numerous derivatives of pJK501 which have deleted different parts of the original plasmid (Kreft and Parrisius, unpublished observations). One of those derivatives which do not undergo further rearrangements, pJK502, has single restriction sites for HindIII, BamHI, SalI (in the Tc^R determinant), and PstI (in the ApR gene).

In order to convert such a bifunctional plasmid into a cosmid system, we have introduced the cos site from pHC79 (Hohn and Collins 1980) into pJK3, yielding pLK103 (Fig. 6). But for unknown reasons all attempts to package this plasmid in vitro into lambda heads have failed so far (G. Luibrand, unpublished observations).

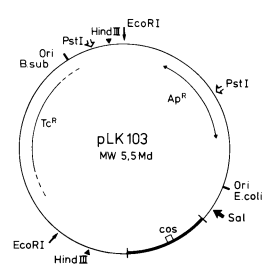


Fig. 6. Restriction endonuclease cleavage map of *pLK*103. The *thick line* indicates the *Bg/*II fragment containing the *cos* site which has been inserted into the *Bam*HI site of *pJK*3

A third type of bifunctional plasmid has been obtained by ligation of EcoRI-cleaved pBSI61-1 and M13mp2 phage (Gronenborn and Messing 1978) RF double-stranded DNA. The resulting hybrid molecule pKJB200 (mol. wt. 7.4×10^6) replicates and expresses tetracycline resistance in both $E.\ coli$ and $B.\ subtilis$, and gives rise to single-stranded DNA and phage in $E.\ coli\ (K.J.\ Burger,\ unpublished\ observations)$. This plasmid has single restriction sites for HindIII and PstI and should facilitate DNA sequencing of cloned fragments.

The plasmids pJK3, pJK302, pKJ502 and pLK103 carry two antibiotic resistance markers (Ap^R and Tc^R), both spanning single restriction sites, thus allowing the detection of recombinant plasmids by insertional inactivation. However, due to the nonexpression of the Ap^R gene from pBR322 in B. subtilis (see Sect. 4) its inactivation can only be detected in E. coli. Nevertheless recombinant plasmids can subsequently be transformed into B. subtilis, using Tc^R as a selective marker.

As will be discussed later, nonexpression and instability of cloned DNA fragments is an important problem in *B. subtilis*. It is, therefore, worthwhile to clone foreign DNA first in *E. coli* and to use the "bridge" character of the described hybrid replicons to introduce the cloned genes into *B. subtilis*.

2.3 Transformation

Bacillus subtilis cells can be transformed by DNA either at the stage of natural competence (*Spizizen* 1958; *Bott* and *Wilson* 1976) or after creation of protoplasts by lysozyme treatment (*Chang* and *Cohen* 1979).

In competent cells the transformability follows the same time course for chromosomal and plasmid DNA (*Contente* and *Dubnau* 1979). Competent cells are efficiently transformed only by oligomers of plasmid DNA (*Mottes* et al. 1979), whereas protoplasts can be transformed with equal efficiency by monomeric or oligomeric plasmid DNA (*Kreft*, unpublished observations).

It has been reported that a restriction-deficient mutant strain of B. subtilis can be more efficiently transformed by plasmid DNA than the restriction-proficient parental strain (Tanaka 1979). It has also been claimed that strains carrying the recE mutation transform poorly (Dubnau et al. 1980). However, in our hands the protoplast system shows no difference between the transformation rates of B. subtilis BR151 and BR151 recE4 or between B. subtilis MT120 r_M m_M recE4 and MT128 r_M m_M recE4 (Tanaka 1979). In all cases plasmids pBS161-1 (a Bacillus plasmid) and pJK3 (a Bacillus/E. coli hybrid plasmid) have been tested and, interestingly, no difference in transformation rate was seen regardless of whether the pJK3 plasmid DNA used for transformation had been isolated from E. coli or from B. subtilis (Kreft, unpublished observations). Clearly the influence of restriction and/or recombination systems on the transformation rate depends upon the particular plasmid used. The transformation efficiency with competent cells or protoplasts is highest with small plasmids, but nevertheless we have been able to transform B. subtilis protoplasts with a plasmid of mol. wt. 17×10^{6} .

2.4 Stability of Vector Plasmids

One prerequisite for the application of a host-vector system to the molecular cloning of DNA is the stable replication of vector plasmids and recombinant molecules in the host. It seems, however, that recombinant plasmids show a remarkable tendency to undergo alterations (mainly deletions) in *B. subtilis*.

Several different mechanisms may be responsible for the phenomena observed. Plasmids can of course recombine with each other or with the chromosome if they contain homologous segments (Keggins et al. 1978; Tanaka and Sakaguchi 1978), the recE4 mutation (Dubnau and Cirigliano 1974) preventing this recombination process. Intramolecular recombination, on the other hand, can occur without involvement of the recE4 function (Tanaka 1979b). In one case it has been shown that such an event can occur in a site-specific way, giving rise to two daughter plasmids containing the entire DNA sequence of the parental plasmid (Fujii and Sakaguchi 1980). The plasmids pBS161 and pBS162 (see. Sect. 2.1) may have been generated by the same process, as suggested by the distribution of restriction sites on these and the parental plasmids and the fact that the sum of the molecular weights of pBS161 and pBS162 is roughly the same as the sum of pBS1 and the large *Eco*RI fragment of *pBC*16.

All the hybrid plasmids described in Sect. 2.2 replicate stably in E. coli without detectable segregation. In B. subtilis the segregation rate after ten generations without selective pressure is 4%-65%, depending on the plasmid examined. They show no deletions or rearrangements while replicating in E. coli, but frequently display extensive deletions when isolated from B. subtilis transformants.

For example, from B. subtilis transformed by pJK501 a large variety of deleted plasmids may be obtained, the deletions affecting both parts (pBSI and pBR322) of the hybrid plasmid and ranging from 3.6-6.1 \times 10⁶ in size. In some cases if plasmid DNA from B. subtilis cells transformed with this plasmid is isolated immediately after transformation, plasmids indistinguishable in size from pJK501 are found, while after subcultivation large deletions are again observed. This clearly demonstrates that in this case the deletion event occurs after the uptake of the plasmid into the cell, as has also been proposed by others (Gryczan and Dubnau 1978).

The occurrence of deletions, at least in the case of pJK501, is independent of the recE function and the restriction/modification system of the recipient. Also the type of E. coli modification of pJK501 plasmid DNA isolated from E. coli and used for transformation of E. subtilis had no influence on the deletion phenomenon (E and E arrisius, unpublished observations).

Insertion of different foreign DNA segments into the same vector plasmid showed that not the vector itself but rather the particular combination of vector with another segment of DNA determines whether this new structure is stable in *B. subtilis* (see. Sect. 4).

2.5 Clone Analysis

The screening for recombinant plasmids is difficult in cases where no marker inactivation and no primary selection for a cloned fragment is possible. To screen for plasmids with inserts (or deletions) the use of a rapid lysis procedure is of great advantage, the method of *Birnboim* and *Doly* (1979) giving in our hands satisfactory results for *B. subtilis*.

It should be kept in mind, however, that due to the remarkable tendency for deletions to occur in *B. subtilis* the mere size of a plasmid isolated from transformants is not a reliable criterion in assessing its structure.

Immunologic (*Broome* and *Gilbert* 1978) and colony hybridization methods (*Grunstein* and *Hogness* 1975) can help to identify particular recombinant plasmids, but have yet to be adapted to *B. subtilis*.

For studies on the expression of cloned DNA fragments a minicell system is available for *B. subtilis* (*Reeve* et al. 1973).

3 Use of Bacteriophage Vectors

Several phage systems of B. subtilis have been well characterized (Graham et al. 1979; Cregg and Ito 1979; Mizukami et al. 1980). In particular, early work has involved the phages Φ3T and ρ11 which may be termed specialized transducing phages as they carry the thymidilate synthetase gene thyP3 (Dean et al. 1976). Kawamura et al. described in 1979 a method to construct specialized transducers of B. subtilis based on the phage pl1. This involved cleavage by EcoRI of DNA from pll and the defective phage PBSX induced from B. subtilis 168. The latter phage contains only host chromosomal DNA (Okamoto et al. 1968), thus limiting the method to the cloning of homologous DNA. It has been extended by Yoneda et al. (1979) to permit the cloning into B. subtilis of foreign DNA for which no primary selection exists. They chose to construct a specialized transducing phage containing the α -amylase gene(s) from B. amyloliquefaciens. Chromosomal DNA from this strain and Φ 3T DNA were digested with BgIII, mixed, and ligated. This ligated mixture was then added to a preparation of chromosomal DNA from B. subtilis RUB200, a strain prototrophic for threonine and defective in α -amylase synthesis. This mixture was in turn incubated with B. subtilis RUB201, a threonine auxotroph lysogenic for Φ 3T. thr⁺ transformants were selected and tested for α-amylase production. Competent cells may take up more than one fragment of DNA (congression) so that by selecting, in this case, thr+ transformants one effectively enriches for cells carrying foreign DNA (such selection gave a 10⁴-fold enrichment for amy⁺ clones). Seven of 10⁵ thr⁺ transformants of RUB201 acquired α -amylase activity and from five of these Φ 3T could be induced. Infection of the amy strain RUB200 with these phages showed a 100% correlation between establishment of lysogeny and the amy+ phenotype. Transformation and selection of amy+ clones showed cotransformation with the phage specific thyP3 gene.

The technique has been utilized for the cloning of spo 077 (quoted in Kawamura et al. 1980) and amyE (Nomura et al. 1979) into p11, but this phage, having a genome with mol. wt. 80 x 10⁶, generates a large number of fragments after routine digestions. *Iijima* et al. (1980) have, therefore, adapted the procedure to the temperate phage Φ 105, which has a genome size of 26 x 10⁶. Chromosomal DNA of B. subtilis 168 (trpC2) was prepared from phage PBSX and after EcoRI digestion ligated with EcoRI digested DNA from Φ105C. Ligated DNA was used to transform B. subtilis (trpC2 lys 3 met B10) lysogenic for Φ105C. Selection for auxotrophic markers, subsequent mitomycin C induction, and transduction of the resulting lysate into B. subtilis (trpC2 lys 3 met B10) allowed isolation of met B+ transducing particles. While Φ105C DNA is insensitive to BamHI, incorporation of the new met B fragment introduced a single site for this enzyme. This seems possible with other phages, e.g., the virulent Φ 1 (Kawamura et al. 1980), which also have no BamHI site. A deletion mutant of Φ 1, Φ 1E2 Δ 1, has been isolated with increased cloning capacity and this has been used to clone pl1 fragments which introduced BamHI and HaeIII restriction sites.

To summarize, initial bacteriophage systems have been shown to be very efficient especially in shotgun cloning of heterologous DNA. Due to the selection marker thymidilate synthetase the phage Φ 3T is particularly useful. Major limitations of the method are that by lysogenization normally only single gene copies can be introduced into recipient cells and that induction of lysogens leads to lysis, which might cause containment problems.

4 Molecular Cloning with Plasmid Vectors

In a strict sense the construction of vector plasmids like pJK3 (Kreft et al. 1978) or pTL12 (Tanaka and Kawano 1980) has already involved the cloning of either heterologous or homologous DNA. This section describes the cloning and expression of isolated genomic fragments and the expression of genetic markers on hybrid replicons.

It has been shown that, at least in certain cases, even DNA cloned in an E. coli vector plasmid can transform B. subtilis without replication of the recombinant plasmid in the Bacillus host (see also Sect. 2.1). The thymidilate synthetase gene from B. subtilis bacteriophage Φ 3T, which has extensive homology to the chromosomal gene, can transform thy B. subtilis to thy when cloned in pSC101 or pMB9 (Ehrlich et al. 1976). On the other hand, the nonhomologous thymidilate synthetase gene from phage \$22 can only transform thy B. subtilis when cloned into an E. coli plasmid carrying a small fragment of DNA homologous to the B. subtilis chromosome (Duncan et al. 1978; Young 1980). In this case the whole recombinant plasmid becomes integrated into the chromosome. Recently it has been shown that the thymidilate synthetase gene from E. coli can also transform B. subtilis when cloned into pBR322 or pMB9 (Rubin et al. 1980). It is not yet clear, however, if there exists sequence homology between the cloned gene and the B. subtilis chromosome.

Most of the hybrid plasmids described in Sect. 2.2 carry more than one antibiotic

Plasmid	Marker	Source	Expression in	
			E. coli	B. subtilis
<i>pJK</i> 3	Ap ^R Tc ^R	pBR 322	+	_
_	Te^{R}	pBS 161-1	+	+
<i>pJK</i> 201	Cm^R	pACYC 184	+	_
-	Cm ^R Tc ^R	pBS 161-1	+	+
pJK 502	Ap^{R} Tc^{R}	pBR 322	+	_
•	Tc^{R}	pBR 322	+	+

Table 3. Expression of antibiotic resistance markers on hybrid replicons in E. coli and B. subtilis

resistance marker. Transformation studies have revealed, however, that these markers are not always expressed in both host bacteria, as is shown in Table 3.

Hybridization of in vivo radioactively labeled RNA from B. subtilis carrying pJK3 or pJK201 (Goebel et al. 1979; Kreft et al., manuscript in preparation) to restriction fragments of these plasmids showed that the nonexpression of the E, coli Ap^R (β-lactamase) and the Cm^R (chloramphenicol acetyl transferase) genes in B. subtilis is due to a transcriptional block. That no structural rearrangement in the DNA sequence of these genes is responsible for this lack of expression was shown following successful retransformation of E. coli with plasmid DNA isolated from B. subtilis (Goebel et al. 1979).

The tetracycline resistance determinant of pBR322 (combined with pBSI to give pJK501) can be expressed in B. subtilis. It is not, however, clear if this resistance determinant indeed originates from E. coli. In addition, since the Tc^R determinant in pJK501 is very close to the *Bacillus* part of the hybrid plasmid, it is possible that its transcription starts at a *Bacillus* promoter.

The tetracycline resistance specified by pBC16 is expressed in E. coli, although at a reduced level (Kreft et al. 1978). The differences observed in expression of E. coli and Bacillus genes in the nonhomologous host might be explained by the promoter specificity of the RNA polymerase. In vitro studies with RNA polymerase from B. subtilis have shown that this enzyme preferentially binds to and transcribes from Bacillus promoters (Williamson and Doi 1978; Lee et al. 1980) in contrast to E. coli RNA polymerase, which transcribes nonhomologous genes quite efficiently (*Davison* et al. 1979).

It has been shown by Ehrlich (1977) that plasmids from S. aureus can express Cm^R or Tc^{R} in B. subtilis. In order to see whether the β -lactamase (E.C. 3.5.2.6.) mediating Ap^{R} in S. aureus can also be expressed in B. subtilis we recloned an EcoRI fragment, containing the β -lactamase gene (from pSC122, Timmis et al. 1975), into pJK3-1. Ap^R/Tc^R transformants of E. coli contained the recombinant plasmid pJK401 and restriction analysis showed that deletions had occurred in both the vector and in the fragment originating from S. aureus. B. subtilis can be transformed to Tc^R with pJK401 and the plasmid replicates stably in this host. Ap^R is not expressed nor can β-lactamase activity be detected intracellularly (Kreft, unpublished observations). It remains to be seen whether this nonexpression in B. subtilis is due to the removal of regulatory DNA sequences by the observed deletion.

Shotgun cloning experiments in B. subtilis seem, from the experience of ourselves and several others, to be rather difficult to perform. The main problem, in addition to nonexpression of heterologous genes, seems to be the difficulty, seen particularly in shotgun experiments, in cloning large fragments of DNA. In one case it was shown that the mean size (mol. wt. 1×10^6) of DNA inserts found in recombinant plasmids is only one-third of the mean size of the fragments in the *Eco*RI or *Hin*dIII digested donor DNA (*Michel* et al. 1980). It is not yet clear if this phenomenon reflects a preferential transformation of recombinant plasmids with small inserts or is due to posttransformational deletions.

A similar experience has been made during attempts to clone the sporulation gene *spo*OF from *B. subtilis* 60015. *Eco*RI digested DNA from this strain, enriched 870-fold for the *spo*OF gene, was ligated to *Eco*RI linearized plasmid *pBS*161-1, and the asporogenous mutant *B. subtilis* strain JH756b was transformed with the ligated mixture. One of the tetracycline-resistant transformants contained *pBS*161-1 with a small insert of DNA, but this plasmid could not complement the sporulation deficiency (*Rhaese* et al. 1979).

Several methods have been proposed to circumvent these difficulties (*Dubnau* et al. 1980). In order to examine the usefulness of cloning first into *E. coli* as an intermediate host we have tried to study the stability and expression in *B. subtilis* of DNA fragments cloned in *E. coli* following their recloning into *B. subtilis/E. coli* hybrid vectors.

We have recloned the *E. coli phoA* gene encoding the periplasmic enzyme alkaline phosphatase (E.C. 3.1.3.1) which is located on a *HindIII/Bam*HI fragment (mol. wt. 3×10^6) of *pSB*53 (*W. Boidol* and *G. Siewert*, manuscript in preparation) into *pJK*3-1. The resulting plasmid *pJK*353 can transform *E. coli* SB44 (*phoA*) to *pho*⁺ but not *B. subtilis* GSY172 (*phoP8 argA11*) (*Le Hégarat* and *Anagnostopoulos* 1973). All the Tc^R *B. subtilis* transformants tested contain a derivative of *pJK*353 with a large deletion of mol. wt. 5.6×10^6 .

This deletion affects the *pho*A gene thereby precluding until now study of its expression in *B. subtilis*. Nevertheless, further investigations should indicate how *Bacilli*, which transport extracellular enzymes to the external medium (for a review see *Priest* 1977), transport *E. coli* enzymes which are normally periplasmic, i.e., are only carried through the inner (cytoplasmic) membrane.

Also using pJK3-1 as a vector the pen gene of B. licheniformis, specifying β -lactamase, has been cloned in B. subtilis. Either the gene already cloned into a λ vector (Brammar et al. 1980) or cloned during a shotgun experiment into phage fd was used as a DNA source. The EcoRI fragment of mol. wt. 2.8×10^6 carrying the β -lactamase genes was inserted into pJK3-1 in both orientations and in both cases gave Tc^R/Ap^R transformants in E. coli and B. subtilis. The recombinant plasmid containing the fragment from λ pen was rather stably maintained in B. subtilis, whereas the plasmid with the fragment from the shotgun cloning became deleted after a few generations (Neugebauer 1980). The expression of the β -lactamase was dependent upon the particular recombinant plasmid and the recipient strain. In B. subtilis SB202 only about 1% of the activity found in the donor strain of B. licheniformis was expressed. In B. subtilis BD170 the expression of β -lactamase from the fragment obtained by shotgun cloning was very good. With the fragment from lambda pen only 10%-30% of this activity was seen in BD170 (Sprengel, personal communication).

A similar cloning experiment has recently been reported (*Gray* and *Chang* 1981) using a bifunctional replicon from *E. coli* and *S. aureus* as a vector. The *B. licheniformis* β -lactamase was efficiently expressed and processed in both *E. coli* and *B. subtilis* BD 224 and secreted into the medium by the *Bacillus* host.

Recombinant plasmids containing genes for exoproteins like β-lactamase may be

very useful as exportation vectors facilitating the secretion of other proteins, the genes for which being fused to the essential parts of the expensive gene.

It has been shown in several cases that genes from the yeast S. cerevisiae can be expressed after cloning and introduction into E. coli and can complement auxotrophic mutations (Struhl et al. 1976; Ratzkin and Carbon 1977). A small HindIII fragment of mol. wt. 2×10^6 containing the arg4 gene of S. cerevisiae (Clarke and Carbon 1978) has been recloned from pYe(arg4) 402-11 into pJK3-1 and the resulting recombinant plasmid pJK3-1 (arg4) transforms E. coli JA228 argH to arg^+ , but not B. subtilis GSY172 phoP8 argA11. In addition, no argininosuccinate lyase (E.C. 4.3.2.1.) activity can be detected in the cells (Kreft, unpublished observations). The recombinant plasmid is stable in B. subtilis and plasmid DNA isolated from this host can retransform E. coli argH to arg^+ . It has been demonstrated that the HindIII fragment containing the arg4 gene carries a promoter which functions in E. coli (Clarke and Carbon 1978) but which from preliminary studies seems to allow no transcription in B. subtilis.

5 Conclusions

Compared to the very sophisticated vectors and recombinant DNA techniques available in *E. coli*, the application of this methodology to *B. subtilis* is still in its infancy. However, the rapidly increasing amount of research in this field may soon allow exploitation of the particular advantages of *B. subtilis* as a host for cloned DNA.

These include (i) nonpathogenicity, (ii) lack of endotoxin, (iii) direct selection of cloned genes which are specific for this species e.g., genes encoding for sporulation and exoenzymes, the latter being of additional interest in the development of "export vectors", and (iv) the possibility of using the transformability of competent cells of *B. subtilis* by homologous chromosomal DNA for the "scaffolding technique" (*Young* 1980) and also for the enrichment of specific markers in DNA samples prior to cloning.

Until now the use of indigenous bacteriophage systems and of *E. coli* as an intermediate host for *E. coli/Bacillus* hybrid plasmid vectors has proved most promising in shotgun cloning, but large problems remain, particularly the nonexpression of heterologous genes and the instability of cloned DNA fragments, which clearly are much more important in this host than in *E. coli*.

Further developments which seem particularly necessary to achieve successful application of the recombinant DNA technique in *Bacillus* are (i) host mutants which allow more stable maintenance of cloned DNA and (ii) vectors which can express heterologous genes regardless of the presence of suitable transcription and translation signals on the cloned fragment.

An in vitro packaging system into bacteriophage heads, comparable to the cosmid system of *E. coli*, should increase the cloning capacity of plasmid vectors.

For the practical application of the recombinant DNA technique in *B. subtilis* to the microbial production of commercially and medically important compounds, as well as the study of regulation processes in this bacterium, vectors with variable copy number or inducible expression of cloned genes will be of great importance.

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