# DNA rearrangements generating artificial promoters

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The promoter-cloning plasmid pBRH4 (a derivative of pBR322 with a partially deleted promoter of the *tet* gene) is shown to contain a sequence which is located near the *Eco*RI site and can operate as an effective Pribnow box, but is not the remainder of the deletion-inactivated *tet* promoter of pBR322. If there is a sequence homologous to the '-35' promoter region at the border of the DNA fragment inserted at the *Eco*RI site, then a compound promoter arises and activates the *tet* gene. Point mutations in the nonfunctional -35 region of pBRH4 also activate the cryptic Pribnow box. Several compound promoters were obtained through deleting small portions of DNA around the *Hin*dIII site of pBR322; the deletions moved various sequences that could operate as Pribnow boxes towards the -35 region of the *tet* promoter.

Pribnow box '-35' region S1-mapping

## 1. INTRODUCTION

Several plasmids with a partially deleted promoter of the tetracycline resistance gene have been used for promoter cloning [1-3]. It has already been noted that the activation of the *tet* gene does not always indicate the presence of the natural promoter in the inserted fragment but may reflect an incidental fit of the end of the fragment and the remainder of the *tet* gene promoter giving rise to a compound promoter [3]. This property of the pBRH4 plasmid, which makes it an imperfect vector for promoter cloning, has been used here to generate artificial promoters.

# 2. MATERIALS AND METHODS

Plasmids were isolated as in [4].

DNA was sequenced as in [5] as modified in [6]. Deletions in pBR322 were obtained through treatment of a *Hin*dIII-linearized plasmid  $(2 \mu g)$ with 5 units S1 nuclease in 100  $\mu$ l of 100 mM NaCl, 3 mM ZnSO<sub>4</sub>, 30 mM sodium acetate (pH 4.8) for 10-30 min at 20°C. The resulting DNA after blunt-end ligation was used to transform *E. coli*  HB101 cells, then ampicillin resistant clones were selected and tested for tetracycline resistance.

Mutations that activate the *tet* gene were obtained either by selecting spontaneously arising Tet<sup>R</sup> clones of *E. coli* C600 (pBRH4) or through treatment of these cells with  $10 \,\mu\text{g/ml}$  *N*-nitro-*N'*-nitrosoguanidine.

The *Msp*I fragment of pBR322, which contains the *tet* gene promoter, was used for the S1-mapping of transcription starting points. The fragment was labelled at the 5'-end with T4 polynucleotide kinase [5], cleaved with *Bsp*RI, then the large subfragment was isolated. RNA extraction and S1 mapping were carried out as in [7] with the following modifications: during S1 treatment NaCl was added to 500 mM, the duration of treatment was reduced to 5 min and the temperature was 20°C.

## 3. RESULTS AND DISCUSSION

The plasmid pEG3503, obtained earlier through inserting the *Eco*RI-G fragment of the *rpo*BC operon into pBRH4 [8], makes cells resistant to  $20 \ \mu g/ml$  tetracycline despite the fact that the pro-

moter contained in the EcoRI-G fragment was reported to be weak [9,10]. To localize the promoter that caused tetracycline resistance, the transcription initiation point was determined by the S1 mapping experiments. The transcription of the *tet* gene in pEG3503 containing cells was shown to be initiated near the border of pBRH4 and the inserted fragment (see fig.1,2). One can see that the Pribnow box, which does not coincide with the Pribnow box of the pBR322 *tet* promoter, belongs to pBRH4, and the -35 region belongs to the fragment of the *rpo*BC operon which normally encodes

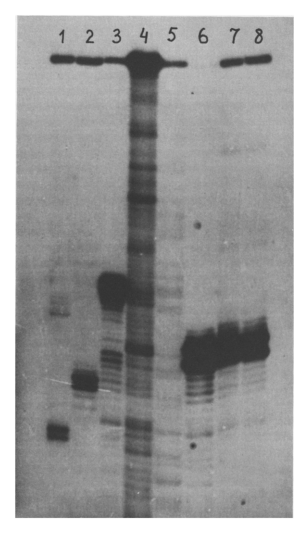


Fig.1. S1-mapping of 5'-ends of plasmid transcripts: 1, pBR $\Delta$ 22; 2, pBR $\Delta$ 15; 3, pBR322; 6, pBR $\Delta$ 10; 7, pEG3503; 8, pBRH4-25; 4,5, DNA fragments obtained as in [5] and used as  $M_r$  markers. the amino acids of the  $\beta$ -subunit of RNA polymerase.

The tet gene does not work in pBRH4 without the insertion, even though there is a Pribnow box, i.e., the sequence occupying the -35 position is not functional. However, pBRH4 gives rise to mutants which sustain cell growth on  $20 \,\mu g/ml$ tetracycline with a spontaneous frequency of  $10^{-8}$ and a nitrosoguanidine-induced frequency of  $10^{-5}-10^{-4}$ . The S1 mapping of 1 spontaneous and 6 nitrosoguanidine-induced mutant variants of pBRH4 (fig.1 shows the data for one of them) revealed one starting point in all cases, the same as in pEG3503, i.e., the same Pribnow box as in the above described compound promoter operates here. One of the nitrosoguanidine-induced pBRH4 mutants (pBRH4-25) has been sequenced. As shown in fig.2, the mutation substitutes T for C at position -34 from the starting point, i.e., creates a functional -35 region.

These results show that when there is a readymade Pribnow box it is easy enough to select a sequence that will function as a -35 region. That is what happened in the case of pEG3503 and, presumably, in other cases when artifacts were observed during cloning in pBRH4. A similar situation may be expected to arise when there is a ready-made -35 region and various sequences are fitted to it. To test this supposition, we obtained several deletion derivatives of pBR322 with an 'ideal' sequence in the -35 region of the *tet* promoter. Deletions were obtained in the vicinity of the *Hin*dIII site (see section 2). After transforma-

	pEG3503
TGGCTGGACTTCGAATTCA	==TTAATGCGGTAGTTTA
	pBRH4-25
TTTCGTTTTCAAGAATTCA	===TTAATGCGGTAGTTTA
ł	pBRH4
TTTCGTCTTCAAGAATTCA	TTAATGCGGTAGTTTA
	pBR∆22.
TTTCGTCTTCAAGAATTCTCATGTTTGACAGCTTATCATCG=====	A
	pBR <b>4</b> 15
TTTCGTCTTCAAGAATTCTCATGTTIGACAGCTTA1CATCG=====	======GTAGTTTA
	pBR <b>41</b> 0
TTTCGTCTTCAAGAATTCTCATGTTTGACAGCTTATCATCG=====	ATGCGGTAGTTTA
	pBR322 *
TTTCGTCTTCAAGAATTCTCATGTTTGACAGCTTATCATCGATAAGCTTTAATGCGGTAGTTTA	
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Fig.2. Mutation changes in the sequence preceding the *tet* gene. =, deleted nucleotides. The point mutation in pBRH4-25 is indicated by an arrow. The bacterial insertion in pEG3503 and the *Eco*RI and *Hind*III restriction sites are underlined.

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tion we selected 3 clones containing deleted plasmids and having different levels of tetracycline resistance. Plasmid sequences are shown in fig.2. In accordance with the deletion length, they are denoted pBR $\Delta 10$ , pBR $\Delta 15$  and pBR $\Delta 22$ . The S1 mapping of these plasmids (fig.1) has shown, as expected, that one and the same -35 region operates in all 3 cases and the promoters differ in their Pribnow boxes. The pBR $\Delta 10$  plasmid, which has a stronger tet promoter than pBR322, proved to have the same transcription starting point, hence the same Pribnow box, as pEG3503 and the pBRH4 mutants with an activated tet gene. In pBR $\Delta$ 15 and pBR $\Delta$ 22 the *tet* promoter is not very effective, but at least in pBR $\Delta 22$  nitrosoguanidine induces mutations (approximate frequency  $10^{-7}$ ) which increase the promoter efficiency several times.

Our results show that a promoter arises with comparative ease if either of its 'halves' is already present. By fitting to it various DNA sequences to recreate a whole promoter one can, in principle, construct many artificial promoters. The process may also occur naturally. For example, it was in this way that a promoter appeared in the gal operon on the insertion of the IS2 element [11].

Our data allow the efficiency and the sequence to be compared for 7 variants of the tet promoter: 4 promoters have the same -35 region but different Pribnow boxes (pBR322, pBR $\Delta 10$ , pBR $\Delta 15$ and pBR $\Delta 22$ ), and 4 variants have the same Pribnow box but different -35 sequences (pEG3503, pBRA10, pBRH4 and pBRH4-25). Fig.3 shows the promoter sequences and summarizes the efficiency data. Promoter efficiency was evaluated by the level of tetracycline resistance of cells carrying the plasmid concerned and by the amount of nucleaseresistant DNA fragments in S1 mapping experiments (the two methods correlate quite well). The relation between the efficiency of these promoters and their sequence is discussed at length elsewhere.

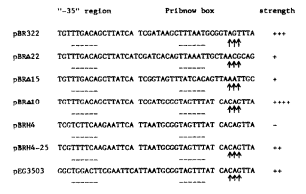


Fig.3. Mutant *tet* promoter sequences. The sequences are aligned with respect to their functionally important parts. Positions of the S1 nuclease cleavage sites are shown by arrows. +, the qualitative level of promoter efficiency.

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