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Abstract
Gene cloning in Neurospora crassa is often achieved by mutant complementation. However, the cloning strategy sometimes requires the isolation of a specific genomic region (by chromosome walking) before transformation of N. crassa. This is the case, for example, if the gene to be isolated has a non-selectable phenotype. Here we specifically describe the construction of the cosmid vector, pAC3, which is designed for direct transformation of N. crassa, its utilization for the construction of a genomic library, and chromosome walking in the region of un-10 on linkage group VII.

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CBM1, a *Neurospora crassa* genomic cosmid library in pAC3 and its use for walking on the right arm of linkage group VII

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Gene cloning in *Neurospora crassa* is often achieved by mutant complementation. However, the cloning strategy sometimes requires the isolation of a specific genomic region (by chromosome walking) before transformation of *N. crassa*. This is the case, for example, if the gene to be isolated has a non-selectable phenotype. Here we specifically describe the construction of the cosmid vector, pAC3, which is designed for direct transformation of *N. crassa*, its utilization for the construction of a genomic library, and chromosome walking in the region of un-10 on linkage group VII.

EXPERIMENTAL PROCEDURES

The pAC3 cosmid vector was constructed as follows:

a) The prokaryotic origin of replication, the antibiotic marker and cos sequences were derived from pJB8 (Ish-Horowicz et al. 1981. Nucl. Acids Res. 9:2989).

b) The pJB8 EcoRI fragment containing the BamHI cloning site was replaced by the EcoRI-EcoRI segment from pWE15 (Catalogue no. 851201 from Stratagene). This 500 bp segment contains two NotI sites and the T3 and T7 phage promoters flanking the BamHI cloning site.

c) A BamHI-BglII fragment from the plasmid MSK338, containing the Neurospora qa-2 gene (coding for catabolic dehydroquinase) was treated with Klenow DNA polymerase 1 to fill in the protruding ends. Such a fragment was blunt ligated in the filled-in SalI site of the construct obtained in b). The final product is the cosmid vector pAC3 (Fig. 1).
We used the pAC3 vector to construct the CBM1 \textit{N. crassa} genomic library. We isolated partially digested chromosomal DNA directly from 74A \textit{N. crassa} agarose plugs prepared as for pulse field electrophoresis (Ballario et al. 1989. Fungal Genetics Newsletter 36:38). The chromosomal DNA was dephosphorylated and ligated at a pAC3 arms-insert ratio of 3:1 (see below for the definition of arms). To avoid the loss of methylated sequences (Grant et al. 1990. Proc. Natl. Acad. Sci. 87:4645) and the recombination events between repeated sequences, a Gigapack Gold packaging extract (Catalogue no. 200214 from Stratagene) was used for packaging and \textit{E. coli} PLK-F' cells (\textit{recA lac mcrA mcrB hsdR gal supE} [\textit{F'} \textit{proAB lacqZ M15 Tn10 (tetr)}]) were infected. An efficiency of about 1.2 x 10(5) clones per ug of \textit{N. crassa} chromosomal DNA was obtained.

**RESULTS AND DISCUSSION**

The characteristics of pAC3 are summarized as follows:
1) The cosmid pAC3 allows the use of the strategy suggested by Ish-Horowicz and Burke for cloning in pJB8. In fact, the presence in pAC3 of asymmetric sites (like HindIII and PstI) enables the preparation of dephosphorylated left and right cos fragments (defined arms). The chromosomal DNA, partially digested by Sau3A or MboI, is dephosphorylated and used without a previous step of size enrichment. Before ligation, the cos fragments are digested with \textit{BamHI} to create the phosphorylated cloning site. This cloning procedure allows one to avoid two potential
problems: a) the formation of cosmids containing inserts with non-contiguous genomic fragments, b) the formation of cosmids containing multiple copies of the vector.

2) The origin of replication of pJB8, present in pAC3, maintains the cosmid at a low copy number and may permit survival of clones containing sequences toxic for *E. coli*.

3) The presence of T3 and T7 phage promoters flanking the cloning site allows both the preparation of RNA probes representing the ends of the cosmids and restriction mapping using T3 and T7 DNA sequencing primers (Evans et al. 1989. Gene 79:9).

4) The sites for a rare cutter restriction enzyme (*Not*I) flanking the cloning site make it possible to remove the inserts and facilitate physical restriction mapping.

5) A selectable marker for direct selection of *N. crassa* transformants has been included.

We have used the CBM1 library for a chromosome walk from *un-10* toward white collar 1 (*wc-l*) on the right arm of linkage group VII. As a starting point for the walk, the cosmid clone 10:E12 from the Vollmer-Yanofsky (V.Y.) library complementing the *un-10* mutation was used. In the absence of indications about the genomic orientation of cosmid 10:E12, we were forced to walk in both directions. Fragments A and B (in Fig. 2) of cosmid 10:E12 were identified by conventional restriction mapping and Southern hybridization and used as starting points in the walk.

![Chromosome walking from un-10 toward wc-l and oli. Part of the linkage groups VII map is shown on the top, the genetic distances in the region between wc-l and oli are indicated. The inserts of the walking cosmids and lambda clone are shown as horizontal lines below the chromosome. The cosmids indicated with the symbol V.Y. are from the Vollmer-Yanofsky library, the others are from CBM1. Probes A and B, used to start the walks, are indicated by boxes. Cosmid 10:E12 was kindly provided by T. Schmidhauser. The clone Az was isolated from the lambda J1 library (M. Orbach) available from FGSC.](image)
Starting from probe A (Fig. 2) we made four steps of walking, covering about 120 kb, and finally found the cosmid 62:D7 from CBM1 overlapping the 2:A10 cosmid from the V.Y. library. This clone was previously mapped at the extreme left of oli by McClung et al. (1989. Nature 339:558) and revealed the walking orientation toward frq. The isolation of the last cosmid (62:D7) was preceded by a walking step in a Neurospora lambda library to isolate the clone Az. This step was necessary to cover about 4 kb apparently absent in both the V.Y. library and ours.

Starting from the probe B we have made four steps so far, representing about 80 kb of genomic sequences in the direction of wc-1. All the walking steps have been checked with the RFLP mapping technique (done by A. Folgori) always confirming the assignment of the selected cosmids to the correct linkage group VII region.

This chromosome walking in the CBM1 library, together with that done by McClung et al., results in the cloning of about 390 kb from the right arm of linkage group VII represented by a series of contiguous cosmids. Having this large region already cloned could allow the molecular investigation of the several known genetic markers mapped in the region.

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