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ANALYSIS OF EUGLENA GRACILIS CHLOROPLAST DNA

Mapping of a DNA sequence complementary to 16 S rRNA outside of the three rRNA gene sets

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Received 9 April 1979

1. Introduction

It is accepted now that Euglena gracilis chloroplast DNA contains 3 rRNA gene sets (16 S, 23 S, 5 S RNA) which are located in 3 tandemly repeated DNA segments of 5600 base pairs each [1–3]. Both the repeats and the rRNA genes were mapped on the circular DNA molecule in relation to numerous restriction enzyme cleavage sites and in particular it was shown that the DNA fragments EcoF, EcoL and EcoP (for DNA fragment nomenclature see legend to fig.1 and [3]) hybridized to rRNA. It was also shown that the 16 S rRNA hybridized exclusively to fragment EcoP [2] and that the three sets of genes were arranged from 5' to 3' end in the order of 16 S–23 S–5 S [4].

However, according to [3,5–8] and contrary to others [1,2,4] total rRNA hybridized also to the fragment *EcoB* which according to present mapping data [1,2,4] should not carry any structural rRNA genes. We have investigated this problem and give in this report unequivocal experimental evidence that the fragment *EcoB* does have a short DNA segment complementary to 16 S but not to 23 S rRNA, suggesting that *Euglena gracilis* chloroplast DNA contains in addition to three complete rRNA gene sets a supplementary 16 S rRNA cistron.

2. Methods

2.1. Preparation of chloroplast DNA and rRNA
Chloroplast DNA was isolated from purified chloroplasts of Euglena gracilis (Z. strain, culture collection

of Algae, Indiana University, no. 753), as in [3]. Chloroplast 16 S and 23 S rRNA were obtained from purified chloroplast ribosomes [9]. The 23 S and 16 S rRNA were purified by twice centrifuging in a 5–20% linear, 5 ml sucrose gradient in a SW50.1 Beckman rotor, 50 000 rev./min, 3 h, 4°C.

2.2. Preparation of DNA fragments BG 16, BG 17 and BG 18

The DNA fragments BG 16 and 17 were prepared from cloned DNA fragments BamD and/or BamE [10]. The respective recombinant DNA was digested first with endo R. BglII and the resulting BG 17 fragment (equivalent to BglQ ~2000 base pairs) was separated on a 5-20% linear 5 ml sucrose gradient (10 mM Tris-HCl (pH 7.9), 1 mM EDTA) in a SW50.1 Beckman rotor, 50 000 rev./min, 3 h, 15°C from the rest of the recombinant DNA consisting of the DNA from the plasmid pBR322 (4362 base pairs [11]) plus the sequences remaining from either BamE (3600 base pairs) or BamD (5000 base pairs). The fragment BG 16 was obtained by a second digestion of this remaining DNA with endo R. BamHI and by centrifugation in an identical sucrose gradient for 5 h. Fragment BG 18 was obtained by double limit digestion of total chloroplast DNA with endo R. BgIII and endo R, BamHI, separation by electrophoresis [3] and electrophoretic elution from a 1% agarose gel.

2.3. Labelling of DNA fragments and rRNA
DNA fragments were labelled by nick-translation
[12] using d- $[\alpha^{-32}P]$ ATP. Ribosomal RNA was
labelled by terminal phosphorylation using $[\gamma^{-32}P]$ ATP

essentially as reported for DNA labelling [13] Complementary labelled RNA was obtained by in vitro transcription of DNA templates using $[\alpha^{-32}P]$ ATP [14]

2 4 DNA RNA and DNA DNA hybridization
DNA fragments were transferred from gels into
millipore HAWP 304 FO filter strips according to
[15]. The filters were wetted with the labelled RNA
in 50% formamide, 5X NaCl/Cit, (1X NaCl/Cit,
0 15 M NaCl and 0 015 M sodium citrate) then
wrapped in Saran foil and incubated for 20 h, at
42°C The filters were washed 4–6 times in 2X NaCl/Cit, 65°C, treated with RNase A (20 μg/ml), 1 h,
20°C and rewashed in 2X NaCl/Cit at 65°C For
DNA DNA hybridization the conditions were the
same as for DNA RNA hybridization with the
exception that the filters were treated before and
after hybridization according to [16]

2.5 Electrophoresis and autoradiography

Gel electrophoresis and autoradiography were carried out as in [3] and as specified in the respective legends

26 Enzymes and radioisotopes

Endo R *BgI*II was a kind gift from Dr T Bickle, University of Basle The following enzymes were purchased endo R *Eco*RI and DNA polymerase (*E coli*), from Boehringer Biochem, Mannheim, RNA polymerase (*E coli* K-12) and polynucleotide kinase (T₄-infected *E coli* B) from Miles Labs, Elkart IN 46514, endo R *Hae*III, from Biolabs, Beverly, MA 01915, endo R *Bam*HI, from Bethesda Res Labs, Rockville, MD 20850 Radioisotopes were from Radiochemical Center, Amersham

3. Results and discussion

In fig.1 a short segment of the Euglena graculus chloroplast genome map is presented. It shows part of the rDNA region with one complete gene set for 16 S + 23 S + 5 S rRNA located in fragment BamD, the beginning of another gene set in BamE, left hand side, and a small part of the fragment BamB with the crucial segment BG 18 at the right-hand side which contains, according to this report, the supplementary 16 S rRNA cistron. Details of the mapping procedure

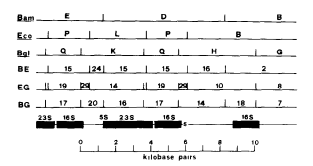


Fig 1 Segment of the rDNA region of the chloroplast genome The 7 horizontal lines symbolize a segment of the double stranded chloroplast DNA The vertical lines indicate the position of the cleavage sites of restriction enzymes. The abbreviations Bam, Eco and Bgl stand for endo R bamHI, endo R EcoRI, and endo R BglII, respectively BF, EG and BG stand for endo R BamHI plus endo R EcoRI, endo R EcoRI plus endo R BglII, and endo R BamHI and endo R BglII The numbers and letters above the lines refer to restriction fragments described in [3] Restriction enzyme nomenclature is according to [17] On the bottom line the ribosomal RNA genes are positioned in scale with respect to the various cleavage sites The 5' end of the 16 S rRNA is 170 base pairs apart from the endo R BglII cleavage site BG 17/14 and BG 17/20 as determined by base sequence analysis (Schwarz, Kossel, Graf, E.S., in preparation) The sizes of the 16 S and 23 S structural genes are taken as 1500 and 2800 base pairs [18], respectively The 5 S rRNA is placed according to [4] The supplementary 16 S rRNA cistron in tragment BG 18 is positioned in accordance to the position of the 16 S rRNA gene in fragment BamD (E) taking as reference point the endo R BglII cleavage site BG 18/7 as discussed here

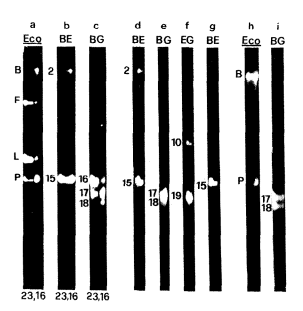
have been published [1-3] and the fragment nomenclature is given in the legend

Total chloroplast DNA was digested with endo R EcoRI (Eco fragments), endo R BamHI and endo R EcoRI (BE fragments), endo R BamHI and endo R BgIII (BG fragments) or endo R EcoRI and endo R BgIII (EG fragments) and the respective fragments were separated by gel electrophoresis. The fragments were hybridized to ³²P-labelled 16 S or 23 S rRNA according to [15]. The respective autoradiographs are shown in fig.2 (panels a—c). We can see that only the 16 S rRNA hybridized to EcoB (a), BE 2 (b) or BG 18 (c). Otherwise the hybridization data were as expected and published [2,3]. As shown in [3] and drawn up in fig 1 these 3 DNA fragments are located outside of a complete rRNA gene set.

Although we had carefully purified the rRNA

sample we could not totally exclude the possibility that, e.g., some contaminating precursor rRNA or mRNA hybridized with the fragments EcoB, BE 2 and BG 18, respectively. In order to eliminate this possibility we prepared complementary ³²P-labelled RNA from purified fragments BG 16 and BG 17 (equivalent to BglQ). The autoradiographs in fig.2 (panels d-f) show the results when complementary RNA of fragment BG 17 was hybridized with the BE, BG and EG fragments of total chloroplast DNA and panel g represents the autoradiograph obtained when complementary RNA of BG 16 was hybridized with the BE fragments. An essential result is that complementary RNA from the fragment BG 17 but not from fragment BG 16 hybridized with fragment BE 2. Furthermore we see that the fragments BG 18 (panel e) and EG 10 (panel f) also hybridized with complementary RNA of BG 17 proving that the fragment BG 17 from BamD (or BamE) contains DNA sequences complementary to sequences within fragment BG 18. Since we definitely know from base sequencing studies (Schwarz, Kössel, Graf, E. S., in preparation) that the fragment BG 17 carries the structural gene for 16 S rRNA, we can deduce that the fragment BG 18 also contains DNA sequences complementary to 16 S rRNA.

In a somewhat reciprocal experiment we hybridized nick-translated BG 18 DNA to *Eco* fragments (panel h)



and to BG fragments (panel i) from total chloroplast DNA. It can be seen that the nick-translated BG 18 DNA hybridized to fragments Eco B and EcoP (panel g) and to BG 17 and BG 18 (panel i). This is in full agreement with all the results shown in fig.2.

In order to estimate the extent of sequence homologies between the fragments BG 17 and BG 18 we digested both kinds of fragments with endo R. HaeIII, an enzyme known to cleave Euglena gracilis chloroplast DNA frequently [9]. The respective results are shown in fig.3. The HaeIII fragment patterns as obtained after gel electrophoresis look almost identical for digests of fragments BG 17 (panel b) and

Fig.2. Autoradiographs of DNA: RNA and DNA: DNA hybridization experiments. Panels a,b,c show, respectively, the hybridization of labelled 23 S and 16 S rRNA to Eco fragments, BE fragments and EG fragments from total chloroplast DNA. Solutions of 32P-labelled 23 S or 16 S rRNA $(4.5 \times 10^{5} \text{ cpm. } \mu\text{g RNA}^{-1}, 10 \,\mu\text{g rRNA/ml})$ with a 5-fold concentration of cold competitor rRNA were used to separately incubate the halved DNA filter strips. Electrophoresis conditions were: 1% agarose, 18 mA, 20°C, 18 h; buffer, 0.04 M Tris, 0.02 M Na-acetate (pH 7.8). The filters were autoradiographed at room temperature for three days (Typon-X-ray film). Panels d,e,f show, respectively, the hybridization of labelled complementary RNA of BG 17 with BE fragments, BG fragments and EG fragments from total chloroplast DNA. Panel g shows the hybridization of labelled complementary RNA of BG 16 fragment with the BE fragments from total chloroplast DNA. Purified fragment BG 17 (BglQ) (~5 µg) was used as template to prepare complementary 32P-labelled RNA according to [14]. Incubation volume for labelling was 80 µl in buffer Hepes-KOH (pH 7.9), 20 mM, Mg-acetate 10 mM, spermine 200 μ M, KCl 200 mM, ATP, GTP, CTP, UTP, 0.5 mM each, $[\alpha^{-32}P]$ ATP, 10 μ Ci (10 Ci/mmol), dithiothreitol 5 mM, 5 units E. coli RNA polymerase, 37°C, 20 min. The reaction was stopped by adding formamide (final conc. 50%) and NaCl/Cit (final conc. 5 X). Panels h,i show, respectively, the hybridization of nick-translated fragment BG 18 with Eco and BG fragments from total chloroplast DNA. Fragment BG 18 (~10 ng) was nick-translated [12] in buffer Tris-HCl, 50 mM (pH 7.8), MgCl 5 mM, 2-mercaptoethanol 10 mM, bovine serum albumin 50 μ g/ml, dCTP, dTTP, dGTP, 10 μ M each, d[α -32P]ATP, $10 \,\mu\text{Ci}$ (200 Ci/mmol), 5 units DNA polymerase, 20°C , 1 h. After passing through Sephadex G-75 the DNA containing fraction was treated with ethanol. The precipitated DNA was resuspended in Tris-HCl 10 mM, EDTA 1 mM (pH 7.8), heat denatured at 97°C, 10 min, chilled in ice, and the solution adjusted to 50% formamide and 5X NaCl/Cit. The hybridization conditions were as given in section 2.

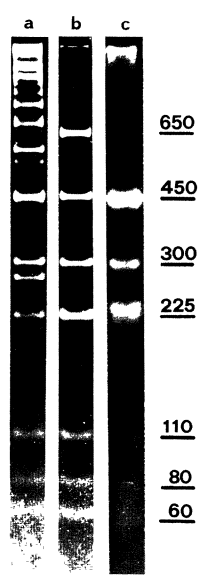


Fig 3 HaeIII fragment patterns of total chloroplast DNA and the DNA fragments BG 17 and BG 18 Panels a and b show, respectively, an ultraviolet photograph of ethidium bromide stained gels with HaeIII fragments of total chloroplast DNA and HaeIII digestion products of fragment BG 17 Panel c shows the autoradiograph of HaeIII fragments obtained by digestion of the fragment BG 18 labelled with 32 P by nicktranslation Gel electrophoresis conditions were as given in fig 2, except that the gel was 3% polyacrylamide–1% agarose Under these conditions only DNA fragments of $M_r < 1000$ base pairs were clearly resolved. The molecular weights as indicated on the right side of panel c are expressed in number of base pairs and were determined by calibrating the gels with HaeIII fragments from pBR322 [11]

BG 18 (panel c) In particular the fragments with sizes of 450, 300, 225, 110, 80 and 60 base pairs are present in both panels. The 650 base pair fragment seen in panel b, but not c, is a terminal piece, located adjacent to fragment BG 16 (B J, E S, unpublished) The corresponding terminal piece in BG 18 must be about 200 base pairs shorter, according to the fragment calibration data in [3] and most likely migrates along with the 450 base pair fragment, thereby generating an intensified band (panel c) The other terminal HaeIII fragment in fragment BG 17 is known to be 225 base pairs long and it comigrates with the internal DNA fragment of equal size. The respective band, therefore, has a stoichiometry of ~2 (B J, E S, unpublished) In panel c the equivalent band is also strongly intensified suggesting that the fragment BG 18 also gives two kinds of HaeIII fragments of 225 base pair length, 1 e, the BglIII cleavage site 18/7 is still part of the DNA segment homologous to the 16 S rRNA gene

In panel a the relevant *Hae*III digestion products (< 1000 base pairs) from total chloroplast DNA are displayed Most important is that those 6 bands which are apparent in both panel b and c are also present in panel a, but as expected, the 650 base pair fragment does not show up in panel a. This indicates that the fragment patterns b and c are not due to some artifacts introduced, e.g., through cloning or nick-translation procedures, but correlate with the *Hae*III fragment pattern of native chloroplast DNA

In summary we can say that the EcoB fragment contains a DNA segment of \sim 1500 base pairs which is complementary to 16 S rRNA. Since the corresponding 23 S rRNA sequence is lacking we postulate that the circular $Euglena\ graculis\ (Z\ strain)$ chloroplast DNA contains in addition to 3 complete rRNA gene sets a supplementary 16 S rRNA gene. It remains to be shown by base sequence analysis whether the 16 S rRNA cistron in fragment EcoB is fully identical to the 3 rRNA genes of the 3 repeat units

Acknowledgements

These investigations were supported by Fonds National Suisse de la Recherche Scientifique, grant 3 687 76 and Fondation Sandoz pour l'Avancement des Sciences Médicales et Biologiques We are grateful to Dr L. Graf and B. Schlunegger for providing the cloned DNA material and to Professor B. Allet, University of Geneva for helpful advice. This work is part of a doctoral program carried out by B.J.

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