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Identification of additional homologues of subunits VII and VIII of the ubiquinol-cytochrome c oxidoreductase enables definition of consensus sequences

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Abstract The Candida utilis QCR7 gene encoding subunit VII of the ubiquinol-cytochrome c oxidoreductase was isolated by functional complementation of the Saccharomyces cerevisiae subunit VII-null mutant. Several other subunit VII homologues as well as homologues for subunit VIII were identified by screening the GenBank database. Some of these homologues for subunit VII could only be identified as such using a consensus sequence that was derived from the multiple sequence alignment. Definition of the consensus should facilitate further analysis of structure/function relationships in this protein.

Key words: Ubiquinol-cytochrome c oxidoreductase; Multiple sequence alignment; Consensus sequence; Candida utilis

1. Introduction

The ubiquinol-cytochrome c oxidoreductase (bc1 complex) plays a central role in the respiratory chain, transferring electrons from ubiquinol to cytochrome c [1]. Coupled to this reaction is the translocation of protons across the inner membrane of mitochondria in eukaryotic organisms and across the cytoplasmic membrane in many bacteria. The bc1 complex of these prokaryotes consists of 3 or 4 subunits [1] while those from eukaryotes may consist of up to 11 subunits [2]. Three subunits, cytochrome b, cytochrome c1, and the Rieske Fe-S protein, contain prosthetic groups and are present in all known bc1 complexes. Since both prokaryotic and eukaryotic bc1 complexes function in a basically similar fashion, the function of the so-called supernumerary subunits is a matter for speculation.

For the bc1 complex from Saccharomyces cerevisiae 10 subunits have been described until now [3,4]. Both subunits VII and VIII (14 kDa and 11 kDa, respectively) are thought to be in close association with cytochrome b [5], the core subunit of the complex, and a role in the binding of ubiquinol/ubiquinone has been suggested [6-8]. Disruption of the genes encoding these subunits showed that both are essential for correct assembly of the complex [9,10].

Here we report on the isolation of the subunit VII homologue of Candida utilis by functional complementation and the identification of a number of other homologues of both subunits VII and VIII by database searches (Table 1). Multiple sequence alignment permits definition of a consensus sequence for both proteins which should prove to be of value in the identification of new members of the two families and in the design of mutant subunits for study of structure/function relationships.

2. Materials and methods

Escherichia coli strain JF 1754 (lac, gal, metB, leuB, hisB, hsdR) was used for DNA manipulations throughout. E. coli transformants were grown in YT medium (1.6% (w/v) bactotryptone, 1% (w/v) yeast extract and 0.5% NaCl) containing 200 µg/ml ampicillin. S. cerevisiae strain DLL70 (a, his3, ura3, leu2; qcr7::LEU2) [9] was used for the transformation of plasmids. Transformation of yeast was performed according to [11]. Transformants were selected on minimal media containing 0.67% (w/v) yeast nitrogen base (Difco), 2% (w/v) glucose, 2% (w/v) ethanol, isolation of plasmid DNA from yeast was performed according to [12].

Genomic DNA was isolated from C. utilis CBS 621 and partially digested with SauIIA. Fragments were separated by electrophoresis. Fragments ranging from 3.5 to 15 kb were isolated from agarose gel and ligated into the BamHI site of YepLac195 [13]. Transformation into E. coli resulted in about 75,000 transformants. Over 95% of these contained the YepLac195 vector with an insert. Transformants were pooled and plasmid DNA was isolated.

An GCG sequence analysis package was made available by the CAOS/CAMM center in Nijmegen, The Netherlands.

3. Results

3.1. Isolation and sequencing of C. utilis QCR7

The C. utilis genomic library was transformed into the S. cerevisiae QCR7-null mutant DLL70 [9]. Of the approximately 10,000 transformants that were replica-plated on ethanol/glycerol (EG) medium, four complemented the respiratory deficiency of the DLL70 strain. Plasmid DNA was isolated and amplified in E. coli. Restriction analysis showed that three clones were identical while the fourth, larger clone contained an overlapping DNA insert. The size of the smallest clone was approximately 7000 bp and a restriction map for enzymes present in the multiple cloning site of YepLac195 is shown in Fig. 1. Three subclones were constructed using the SstI restriction sites and subsequently transformed into DLL70. Subclone 1, containing the first 1750 bp, was able to suppress the EG-negative phenotype of DLL70.

Suitable fragments were subcloned into vector pBluescript SK (Stratagene) and the DNA sequence was determined on one strand only [14]. Only one open reading frame (ORF) longer than 200 bp was found. The DNA sequence of the ORF plus some 400 bp of flanking sequence was determined on the other
The QCR7 gene of *C. utilis* is 378 bp long and codes for a protein of 126 amino acids with a calculated molecular weight of 14.1 kDa. Alignment with the *S. cerevisiae* subunit VII shows 54% sequence identity, which is more or less equally distributed over the entire protein. Secondary structure predictions and hydrophilicity plots also indicate a high level of conservation between the two proteins (data not shown).

The capacity of Dll70 complemented with the *C. utilis* QCR7 gene to grow on a non-fermentable carbon source indicates that the heterologous *bc* complex in which subunit VII is replaced by its homologue from *C. utilis* is (at least partially) functional. However, whether lower efficiency is due to sequence differences or lower expression of the *C. utilis* gene in *S. cerevisiae* still has to be resolved.

### 3.2. Database search for subunit VII homologues

We have performed a database search for proteins or ORFs showing sequence identity with subunits VII and VIII. A TBLASTn search was performed using yeast subunit VII as the query sequence. Of the sequences retrieved only those with a Smallest Sum Probability *P(N)* smaller than 10^-32 were considered to be significantly similar. In addition to the known homologues of this protein a *Drosophila melanogaster* clone was found with sequence similarity of statistical significance (*P(N)* = 8.2 e^-16). This clone contains the Mst98Ca and Mst98Cb genes. Downstream of these genes is an ORF encoding part of subunit VII. According to the multiple sequence alignment in Fig. 3 about 75 bp at the 3’-end of the gene are missing.

In another TBLASTn search we used the *Solana tuberosum* (potato) subunit VII as the query sequence. This resulted in the retrieval of 3 clones containing rice (*Oryza sativa*) cDNA with significant similarity. These clones were expressed sequence tags (EST) and each contained only part of the QCR7 gene. However, when aligned these 3 sequences permit reconstruction of a complete QCR7 gene. The overlapping regions showed complete identity, suggesting that these cDNAs were translated from the same gene. In addition, one maize (*Zea mays*) cDNA was retrieved containing part of the QCR7 gene. Several ambiguous nucleotides were present and an additional ambiguity had to be introduced at position 99 in order to align the translated sequence with the potato protein. Two clones containing *Arabi-...
S. cerevisiae protein) with $P = 8.7 \times 10^{-7}$. Two clones were found for *P. falciparum* which were almost identical. Since several ambiguous nucleotides are included by the authors it is likely that the differences between the two clones are due to sequence errors. The homologues from these two organisms clearly match the consensus sequence (see Fig. 3). However, the statistical value $P(N)$ given by TBLASTn was 0.40 for *P. falciparum* and 0.72 for *C. elegans.*

3.3. Database search for subunit VIII homologues

Similar search strategies were used to identify new members of the subunit VIII family. Using the potato subunit VIII as a query sequence resulted in the identification of the complete *QCR8* gene of *A. thaliana.* The sequence contains several ambiguous nucleotides and is 13 amino acids longer than the potato protein. The proteins differ extensively at the C-terminus, in clear contrast to the rest of the protein. However, dele-

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**Fig. 3.** Multiple sequence alignment of the subunit VII homologues. Sequences were aligned using the GCG PileUp program (gap weight, 3.00; gap length weight, 0.10) [21]. Amino acids conserved in all sequences are indicated by a black box; amino acids appearing in at least 75% of the sequences are indicated by a grey box. Gaps are indicated (—). 1 = *S. cerevisiae* [15]; 2 = *K. lactis* [16]; 3 = *C. utilis* (this study); 4 = human [17]; 5 = rat [17]; 6 = bovine heart [18]; 7 = *D. melanogaster* (this study); 8 = *P. falciparum* (this study); 9 = *O. sativa* (this study); 10 = *S. tuberosum* [19]; 11 = *T. aestivum* [20]; 12 = *C. elegans* (this study); 13 = *Z. mays* (this study); 14 = *A. thaliana* (this study). Note that sequences from 5, 7, 8, 11, 12 and 13 are only partial.
Here we show that the use of a consensus sequence as a tool for identifying homologous proteins. Not all of the sequences found in this study could functionally complement the subunits VII and VIII homologues. However, we generated and used to search the SwissProt database similar to the approach for subunit VII. Also in this search no proteins lacking a clear consensus sequence were found with significant similarity.

The second part of this study involves the identification of new homologues of both subunits VII and VIII. Three clones were found containing rice subunit VIII homologues. These clones are expressed sequence tags and all contain the complete QCR8 gene. Similar to *A. thaliana*, two of these clones contained two ambiguous nucleotides and a frame shift was found in one of them (A34 was deleted in clone C0237-1A). Alignment of the three genes showed that two of them were identical while the third showed six differences on the nucleotide level resulting in four amino acid changes. This third sequence is displayed in the sequence alignment in Fig. 4 as it did not contain any ambiguous nucleotides.

Although the multiple sequence alignment of the subunit VII homologues lacks a clear consensus sequence, a profile was generated and used to search the SwissProt database similar to the approach for subunit VII. Also in this search no proteins were found with significant similarity.

4. Discussion

We have isolated a gene from *Candida utilis* and identified it as the QCR7 gene encoding subunit VII of the bc1 complex based on sequence identity with the *S. cerevisiae* subunit VII and on the fact that a plasmid carrying this gene is able to functionally complement the *S. cerevisiae* QCR7-null mutant.

The second part of this study involves the identification of new homologues of both subunits VII and VIII. These homologues were identified by screening DNA and EST databases using protein sequences of several known subunit VII and VIII homologues. Not all of the sequences found in this study could be identified based on sequence identity with known proteins. Here we show that the use of a consensus sequence as a tool in screening sequence databases increases the chance of identifying homologous proteins.

The conserved amino acids, indicated by a black box in Fig. 3, are not distributed over the entire protein, but are predominantly found in the central domain (residues 31–89 in *S. cerevisiae*). The N-terminus (residues 1–30 in *S. cerevisiae*) is highly variable, both in amino acid composition and in length. The N-terminus of *S. cerevisiae* subunit VII (residues 1–18) has been proposed to contain a mitochondrial targeting sequence [24] as it may form an amphipathic helix with a high hydrophilic moment. Apart from the N-terminal methionine, this targeting sequence is not cleaved [1].

The C-terminus (residues 90–127 in *S. cerevisiae*) contains only a few conserved amino acids. However, the sequences do show additional similarity in that they all contain a large number of both acidic and basic residues, making this part of the protein highly hydrophilic. Another interesting feature of this region is that five positions in the alignment contain more than 50% glutamate residues. A previous study on this C-terminus in *S. cerevisiae* showed the importance of this part of the protein for correct assembly of the bc1 complex, but not for electron transport [25]. The study also implied that the overall features of this region (hydrophilicity and charges) rather than specific residues are important, a feature which is confirmed by the sequence alignment in Fig. 3.

Three clones were found containing rice *QCR8*. Two were identical while the third contained changes both in nucleotide and amino acid sequence. Although sequence errors could be responsible for these differences, it should be noted that gene duplicates are relatively common in plants. In the case of the Rieske Fe-S protein it was shown that in tobacco at least 5 different genes are present and transcribed [26]. Most of these proteins differ only by a few amino acids. For subunit VII also five different genes are present and transcribed [26]. Most of these proteins differ only by a few amino acids. For subunit VII also five different genes are present and transcribed [26].

<table>
<thead>
<tr>
<th>Clone</th>
<th>Species</th>
<th>Sequence Alignment</th>
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<tbody>
<tr>
<td>1</td>
<td>FRRFK</td>
<td>S. cerevisiae</td>
</tr>
<tr>
<td>2</td>
<td>FRRVK</td>
<td>K. lactis</td>
</tr>
<tr>
<td>3</td>
<td>RRTR</td>
<td>O. sativa</td>
</tr>
<tr>
<td>4</td>
<td>HKVSENW</td>
<td>S. cerevisiae</td>
</tr>
<tr>
<td>5</td>
<td>HKVSEN</td>
<td>S. cerevisiae</td>
</tr>
<tr>
<td>6</td>
<td>HKVSEP</td>
<td>S. cerevisiae</td>
</tr>
<tr>
<td>7</td>
<td>HKVSDX</td>
<td>S. cerevisiae</td>
</tr>
</tbody>
</table>

Fig. 4. Multiple sequence alignment of the subunit VIII homologues. Sequences were aligned using the GCG PileUp program (gap weight, 3.00; gap length weight, 0.10) [21]. Amino acids conserved in all sequences are indicated by a black box. A previous study on this C-terminus in *S. cerevisiae* showed the importance of this part of the protein for correct assembly of the bc1 complex, but not for electron transport [25]. The study also implied that the overall features of this region (hydrophilicity and charges) rather than specific residues are important, a feature which is confirmed by the sequence alignment in Fig. 3.
The multiple sequence alignment in Fig. 3 shows that there is a clear consensus sequence for subunit VII. This subunit VII ‘family’ now contains members of the classes of yeasts, mammals, plants, insects, protozoa and eumetazoa. In marked contrast to this, is the low sequence identity amongst the subunit VIII homologues. In line with this observation is that the QCR8 gene from *S. pombe* could be cloned on the basis of complementation of an *S. cerevisiae* deletion strain [27], but not the QCR7 gene from this organism. The requirements for subunit VII to allow assembly are apparently more strict than those for subunit VIII. Some of the homologues of subunit VII (from *P. falciparum* and *C. elegans*) were identified using the consensus sequence. It is likely that there are still several unidentified homologues of subunit VIII present in the databanks. The lack of a clear consensus sequence as a tool in screening the database would thus be the reason for this.

The very low sequence conservation amongst the subunits VIII (see Fig. 4) indicates that proteins even less related will most likely lack sequence similarity. Subunit IV of the *bc* complex from *Rhodobacter sphaeroides* has been suggested to fulfill the same function as the 9.5 kDa subunit from bovine heart, the subunit VIII homologue [28], this despite the fact that no amino acid sequence similarity could be found. However, structural features, such as hydrophobicity, are similar. The same features are conserved among the known subunit VIII homologues [27].

The SwissProt database searches that were performed using a profile of the multiple sequence alignment of both subunits did not find any protein showing similarity to the respective consensus sequences. For both subunits a role in the binding of ubiquinol/ubiquinone has been suggested [6-8]. It would be very useful if a consensus sequence for Q-binding could be defined. However, studies on other quinone binding proteins, such as cytochrome *b* [29] and those in the photosynthetic reaction center [30] indicate that the Q-binding pocket is such as cytochrome *b* [29] and those in the photosynthetic reaction center [30] indicate that the Q-binding pocket is

The level of sequence conservation among the two protein families provides information about the function of these subunits within the *bc* complex. It thus seems that subunit VIII predominantly requires certain structural features while subunit VII contains several specific amino acids required for the function of this protein.

In general we propose that the identification of homologous proteins and the definition of a consensus sequence provides targets for the design of mutant proteins for study of structure/function relationships.

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References