PIGMENT-PROTEIN COMPLEXES OF THYLAKOID MEMBRANES OF Deschampsia antarctica Desv. Plants

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Abstract. The molecular analysis results of pigment-protein complexes of photosynthetic membranes of Deschampsia antarctica plants, one of the native representatives of the Antarctic region are presented. Bioinformatics searching of genome products of Deschampsia antarctica in available protein databases related to the structure and function of plastids was carried. The searching of Arabidopsis thaliana and Oryza sativa japonica homolog search submitted in free databases is also performed. The results of comparative analysis of primary sequences of the known and potential products are presented.

Keywords: Deschampsia, pigment-protein complexes, proteome.

Introduction

The investigations of protection responses of plants on abiotic stress conditions is actual today. Antarctic species Deschampsia antarctica Desv. (Poaceae) is unique plant object for studying of adaptive mechanisms of plants on abiotic stress action. This can be explained both the absence of breeding and biotechnological interferences in the development of this species and its unique habitat.
Phenotypical and genotypic variability are the basis of plant adaptation on abiotic stress. Functioning of them provides the balance between the requirements of maximal ontogenetic adaptation and maintenance of phylogenetic flexibility of populations (Жученко, 1994; Кир’яченко, 2005).

Some abiotic factors in Antarctic are excessive illumination and elevated levels of UV-B radiation. Therefore formation of active paths of adaptation for maintenance of photosynthetic activity take place both at the level of the ultrastructural organization of organels (Gielwanowska, 2005), that provides effective mechanism of dissipation of excessive energy, and at a biochemical level of photosynthetic reactions.

Primary processes of photosynthesis take place in thylakoid membranes of chloroplasts during redox transformations of components of electron transport chain. They involved in adaptive transformations of photochemical processes. Thylakoid membranes contain five types of protein complexes, which realize distribution of a charge and transfer of electrons. There are photosystem I (PS I), photosystem II (PS II), their light-harvesting complexes, cytochrome $b_{5}/f$ complexes of proton ATP-synthase ($CF_{0}$-$CF_{1}$).

The aims of our investigation were isolation and analysis pigment-protein complexes of thylakoid membranes of chloroplasts and searching in accessible databases of amino acid sequences of $D. antarctica$ which belong to pigment-protein complexes of thylakoid membranes and also carrying out of their comparative analysis with Arabidopsis thaliana (L.) and Oryza sativa L. ssp. japonica cv. Nipponbare sequences for finding-out of particularities of formation of adaptive reactions of $D. antarctica$ plants at a level of primary processes of photosynthesis.

Materials and methods

The analysis of pigment-protein complexes of thylakoid membranes carried out by method of nondenaturing polyacrylamide gel electrophoresis in polyacrylamide gel with using of sodium lauryl sulfate (SLS) for decomposition of membranes desintegration (Anderson, 1980). The relative pigment-protein content was estimated by chlorophyll concentration that was determined in 80 % acetone (Arnon, 1949). Electrophoregrams scanned (‘Mustek’, China) and then analyzed by TotalLab V1.10.

Searching of $D. antarctica$ sequences was spent in open databases Swiss-prot / TREMBL (http://www.expasy.org). Searching of potential homologues was carried out by comparison of $D. antarctica$ sequences with proteomes of A. thaliana and O. sativa, as vegetative organisms which are most full presented in databases. Searching of homologous sequences and determination of a homologous level carried out with BLASTp tool (http://www.ncbi.nlm.nih.gov/blast/Blast.cgi). The selection of homologous sequences for further analysis was spent by standard method according to which proteins is homologous under identity $\geq 25 \%$ with E-value $<10^{-10}$. Global equalization of protein sequences carried out with on-line resource Clustal with using of BLOSUM matrix. The analysis of domain architecture of amino acid sequences spent with on-line resource ART (http://smart.embl.de).

Results and discussion

We identified seven strips according to non-denaturing green SDS-electrophoresis. Each of them associated with pigment-protein complexes according to Anderson nomenclature (Anderson, 1980) (fig. 1).

In result of comparison of pigment-protein complexes of thylakoid membranes of $D. antarctica$ plants with Pisum sativum L. plants (Топчий, 2006) the essential qualitative differences between them were not exposed.
Quantitative differences in general content of LHCII (oligomeric (LHCP1) and monomeric (LHCP3) forms), chlorophyll content in CPα, (protein complexes of complexes of near antenna) are revealed. In our opinion, quantitative differences associated with features of growth conditions of D. antarctica plants. Thus, the influences of excessive light and increased level of UV-B radiation.

The relative content of pigment-protein complexes of thylakoid membranes of D. antarctica plants is presented lower. CP1a and CP1 is 8,97 ± 0,38 (%) and 12,34 ± 0,26 (%), oligomeric form (LHCP1) of LHCII - 27,02 ± 0,53 (%), whereas monomeric form LHCP3) - 20,16 ± 0,61 (%). The relation between forms (LHCP 1/LHCP3) is equaled 1,34 ± 0,02, general content of LHC II (LHCP1 + LHCP3) is 47,17 ± 1,11 (%).

Scanning of databases has shown, that 37 amino acid sequences which belong to D. antarctica proteome currently there is in free access. From them 20 sequences are potential proteins or fragments of proteins with specified function and 17 are potential proteins or fragments of proteins which function is not determined. Thus, annotation of D. antarctica genome and proteome is on the initial stage of determination of function of all known proteins, and functions practically all known potential proteins are not determined finally.

In our work special attention pays to plastid proteins. We selected from databases 15 proteins, which had chloroplast origin and related to families RsbG and PsbC (Nixon, 1989, Luciski, 2006).

The comparison of Psb sequences of D. antarctica received from databases (ABS30954; ABS30955; ABS30956; ABS30957; ABS30958) demonstrated, that they are identical, and they are the sequencing of one potential product ('Clustal' program). The analysis of domain architecture of this fragment was conducted by online SMART recourse. It confirmed the accession of this fragment to PsbC. The determination of fragmen of protein domain architecture of Psb was impossible for D. antarctica plants. It associated with small sizes of amino-acid residues.

Searching of homologous sequences in A.thaliana and O.sativa proteomes was lead (BLASTp instrument) for specification of function of D. antarctica hypothetical product. As result, have been allocated amino acid sequences NP051055 A.thaliana (identity of 97 %, E-value 0,00) and NP039367 O.sativa (identity of 98 %, E-value 0,00). In articles of databases they are presented proteins of PsbC.
family are a part and participate in binding of chlorophyll and β-carotene and transfer of excitation energy to reaction center (Barber, 2002).

Multiple global alignment of found sequences of *A. thaliana* and *O. sativa* with fragment has revealed the significant level of conservatism of this proteins (fig. 2).

Fig. 2. Comparison of amino acid sequences of PS II protein fractions of *D. antarctica*, *A. thaliana* and *O. sativa*.

The structured area of these proteins includes from 31 to 473 amino-acid residues. The investigated fragment of *D. antarctica* sequence practically completely coincides with the sequence of proteins from 45 to 473 amino acid residue. But positions 106, 170, 185, 245, 264, 437, 463 had strong replacements, on 207, 277, 470 positions was weak replacement, and on 242, 282, 468 coincidence was absent. Probably strong replacements lead to the certain modifications in structure of this protein which provides the effective mechanism of excessive energy dissipation for maintenance photosynthesis in normal conditions.

**Conclusions**

Quantitative differences in general content of LHC II (oligomeric and monomeric forms), chlorophyll content in CPα zone (pigment-protein complexes of nearest antenna) are revealed. Results of global alignment and domain architecture have shown, that analyzed fragment of amino acid sequence of PS II protein fraction of *D. antarctica* has high affinity with sequences of *A. thaliana* and *O. sativa* which belong to PsbC family of proteins, and it is a fragment of a product of a gene of psbC family. Also it is possible to note, that in the given fragment is absent C-final sequence, which have size as one or several exons.

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