

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
24 January 2008 (24.01.2008)

PCT

(10) International Publication Number
WO 2008/010228 A2

(51) International Patent Classification:
A01H 5/00 (2006.01)

(21) International Application Number:
PCT/IL2007/000920

(22) International Filing Date: 22 July 2007 (22.07.2007)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
60/831,937 20 July 2006 (20.07.2006) US

(71) Applicant (for all designated States except US): **YEDA RESEARCH AND DEVELOPMENT CO. LTD.** [IL/IL]; at the Weizmann Institute of Science, P.O. Box 95, 76100 Rehovot (IL).

(72) Inventors; and

(75) Inventors/Applicants (for US only): **SCHERZ, Avigdor** [IL/IL]; 4 Tor HaAviv Street, 76329 Rehovot (IL). **SHLYK-KERNER, Oksana** [IL/IL]; 6 Shkolnik Street, 76209 Rehovot (IL). **SAMISH, Ilan** [IL/IL]; 7 HaPartizanim Street, 74037 Nes Ziona (IL). **KAFTAN, David** [CZ/CZ]; Komenskeho 395, CZ-37333 Nove Hradky (CZ). **DINAMARCA, Jorge** [CL/CL]; Saavedra 680, Victoria (CL).

(74) Agents: **G.E. EHRLICH (1995) LTD.** et al.; 11 Menachem Begin Street, 52521 Ramat Gan (IL).

(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

- without international search report and to be republished upon receipt of that report
- with sequence listing part of description published separately in electronic form and available upon request from the International Bureau
- the filing date of the international application is within two months from the date of expiration of the priority period

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: PHOTOSYNTHETIC ORGANISMS AND COMPOSITIONS AND METHODS OF GENERATING SAME

(57) Abstract: An isolated polynucleotide is provided. The isolated polynucleotide comprising a nucleic acid sequence encoding a polypeptide of a Type II reaction center of a photosynthetic organism, the nucleic acid sequence being capable of imparting the type II reaction center with an activity under a temperature range different than that of the type II reaction center endogenous to the photosynthetic organism. Also provided are methods of using the sequences for generating photosynthetic organisms or tailor-made thermotolerance.



WO 2008/010228 A2

PHOTOSYNTHETIC ORGANISMS AND COMPOSITIONS AND METHODS OF
GENERATING SAME

FIELD AND BACKGROUND OF THE INVENTION

5 The present invention relates to photosynthetic organisms and, compositions and methods of generating same.

 The energy crises of the 1970's combined with world-wide climate changes linked to the accumulation of excess carbon dioxide in the atmosphere, has initiated a significant return to the development and use of biomass as a resource for fuel and
10 chemical products. Presently, society's dependence is shifting away from petroleum to renewable biomass and energy resources, in order to aid in the development of a sustainable industrial society and to manage the green house effect. The US department of energy has set goals to replace 30 % of the liquid petroleum transportation fuel with biofuels and to replace 25 % of the industrial organic
15 chemicals with biomass-derived chemicals by 2025 [Ragauskas et al, *Science*, 311, 484-489 (2006)]. The EU has set targets at 5.75 % for all petrol and diesel transport fuels to be biomass derived by the end of 2010 [Ragauskas et al, *supra*].

 Biomass production methods presently enable different chemical and biofuel (e.g. biodiesel and ethanol) production from plant resources. Current production of
20 biomass in general and biofuel in particular mainly relies on higher plants and trees, as for example corn and sugar cane plants. However, biofuel production from these plants is limited in yield and enhanced consumption of such plants as a biofuel source would lead to a severe shortage in food supply worldwide. Furthermore, maximum productivities of higher plants and trees are restricted to areas with prime soil, water,
25 and climate (primarily the tropics). Plant leaves exist in an aerial environment and are subject to large evaporative moisture losses, which directly inhibit the process of photosynthesis.

 Biomass production by cyanobacteria and microalgae, which are the most productive carbon dioxide users of all photosynthetic organisms and can fix greater
30 amounts of carbon dioxide per land area than higher plants, is restricted to a relatively narrow range of temperatures defined by their native habitat. Mesophilic organisms show maximal rate constant at 20-30 °C while thermophilic strains achieve similar rates at 60-70 °C. This temperature range mainly reflects constraints of the

photosynthetic energy conversion machinery (photosystems) and of the carbon fixation one (the Rubisco complex).

Biomass is mainly generated in the course of photosynthesis which photocatalyses carbon dioxide fixation via the Rubisco complex. Plants, microalga and cyanobacteria use photosystems I and II (PSI and PSII), to convert light energy into chemical energy. The central unit of a PSII protein complex is the reaction center (RC). The functional core of PSII RC consists of a heterodimer made of the two homologous protein subunits D1 and D2 along with one unit of cytochrome b559. The D1 and D2 protein subunits each have five (*A*, *B*, *C*, *D*, and *E*) transmembranal (TM) α helices. The cofactors that carry out electron transfer (chlorophyllus type molecules and quinones) in response to illumination and thereby perform the primary energy conversion, are mainly bound (non-covalently) to helices *D* and *E* of the D1/D2 subunits.

Numerous studies showed that the photosynthetic energy conversion by PSII RC is highly sensitive to irradiation and temperature variations [Takahashi et al., *Plant Cell Physiol* 45, 251-5 (2004); Murata et al *Biochim Biophys Acta* 1767, 414-21 (2007)]. Also, recent studies have suggested that the PSII RC is a key player in regulating the rate of photosynthetic energy conversion in response to the prevailing temperature [Yamasaki et al., *Plant Physiol* 128, 1087-97 (2002)].

Taking into consideration the global warming effect, annual and even daily changes in temperature in aquatic areas (e.g., oceans, small lakes and ponds), dramatically narrow the efficiency of biomass production of thermophilic and mesophilic microalgal strains as well as of different strains of cyanobacteria and thus limits their growth to the tropical arena. Even there, current global heating is expected to exceed the thermotolerance and production efficacy of these organisms [Wraight, *Front Biosci* 9, 309-37 (2004); Behrenfeld et al. *Nature* 444, 752-5 (2006)]. Although the expected changes in global temperatures are only in the order of several degrees, it is predicted that biomass production may be dramatically effected.

The increased need for biofuel and the concomitant shortage of food across the world underscores the urgent need for methods of increasing resistance of plants, alga and microalgae to ambient temperature changes.

SUMMARY OF THE INVENTION

The present invention successfully addresses the shortcomings of the presently known configurations by providing photosynthetic organisms and, compositions and methods of generating same.

5 Thus, according to one aspect of the present invention there is provided an isolated polynucleotide comprising a nucleic acid sequence encoding a polypeptide of a Type II reaction center of a photosynthetic organism, the nucleic acid sequence being capable of imparting the photosystem II reaction center with an activity under a temperature range different than that of the type II reaction center endogenous to the
10 photosynthetic organism.

According to further features in preferred embodiments of the invention described below, the polypeptide of the photosystem II reaction center is selected from the group consisting of D1, D2, L and M.

15 According to still further features in the described preferred embodiments, the polypeptide is D1.

According to still further features in the described preferred embodiments, the activity comprises solar energy conversion activity.

20 According to still further features in the described preferred embodiments, the D1 polypeptide comprises an amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 and/or at a position corresponding to 212 of the SEQ ID NO: 36.

According to still further features in the described preferred embodiments, the amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 comprises a Serine to Alanine mutation.

25 According to still further features in the described preferred embodiments, the amino acid sequence alteration at a position corresponding to 212 of SEQ ID NO: 36 comprises a Serine to Cysteine mutation.

30 According to still further features in the described preferred embodiments, the D1 polypeptide comprises an amino acid sequence alteration at any amino acid position corresponding to 208-212 of SEQ ID NO: 36.

According to still further features in the described preferred embodiments, the photosystem II reaction center is rendered more thermotolerant as compared to the photosystem II reaction center endogenous to the photosynthetic organism.

According to still further features in the described preferred embodiments, the photosystem II reaction center is rendered more thermoplastic as compared to the photosystem II reaction center endogenous to the photosynthetic organism.

5 According to still further features in the described preferred embodiments, the photosynthetic organism is selected from the group consisting of a higher plant, a photosynthetic bacteria and an algae.

According to still further features in the described preferred embodiments, the photosynthetic organism is selected from the group consisting of a mesophile, a thermophile and a psychrophile.

10 According to still further features in the described preferred embodiments, the temperature range comprises 10 °C – 43 °C.

According to still further features in the described preferred embodiments, the temperature range comprises an upwards shift.

15 According to still further features in the described preferred embodiments, the upwards shift is by at least 6 °C.

According to still further features in the described preferred embodiments, the temperature range comprises a downwards shift.

According to still further features in the described preferred embodiments, the downwards shift is by at least 2.5 °C.

20 According to still further features in the described preferred embodiments, the temperature range comprise a wider temperature range.

According to still further features in the described preferred embodiments, an amino acid at position 209 or 212 is selected from the group consisting of glycine, cysteine, alanine, threonin, asparagine, aspartanine, proline, valine, isoleucine, 25 leucine, glutamine and glutamic acid.

According to another aspect of the present invention there is provided a nucleic acid construct comprising the polynucleotide.

According to still further features in the described preferred embodiments, the nucleic acid construct further comprises a cis-regulatory element.

30 According to still further features in the described preferred embodiments, the cis-regulatory element is a promoter.

According to yet another aspect of the present invention there is provided a cell of a photosynthetic organism comprising the nucleic acid sequence.

According to still another aspect of the present invention there is provided an isolated polypeptide comprising an amino acid sequence encoding a polypeptide of a photosystem II reaction center of a photosynthetic organism, the nucleic acid sequence being capable of imparting the photosystem II reaction center with an activity under a temperature range different than that of the type II reaction center endogenous to the photosynthetic organism.

According to still further features in the described preferred embodiments, the polypeptide is selected from the group consisting of D1, D2, L and M.

According to still further features in the described preferred embodiments, the polypeptide is D1.

According to still further features in the described preferred embodiments, the activity comprise solar energy conversion activity.

According to still further features in the described preferred embodiments, the D1 comprises an amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 and/or at a position corresponding to 212 of the SEQ ID NO: 36.

According to still further features in the described preferred embodiments, the amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 comprises a Serine to Alanine mutation.

According to still further features in the described preferred embodiments, the amino acid sequence alteration at a position corresponding to 212 of SEQ ID NO: 36 comprises a Serine to Cysteine mutation.

According to still further features in the described preferred embodiments, the D1 comprises an amino acid sequence alteration at any amino acid position corresponding to 208-212 of SEQ ID NO: 36.

According to still further features in the described preferred embodiments, the photosystem II reaction center is rendered more thermotolerant as compared to the photosystem II reaction center endogenous to the photosynthetic organism.

According to still further features in the described preferred embodiments, the photosystem II reaction center is rendered more thermoplastic as compared to the photosystem II reaction center endogenous to the photosynthetic organism.

According to still further features in the described preferred embodiments, the photosynthetic organism is selected from the group consisting of a higher plant, a cyanobacteria and an algae.

According to still further features in the described preferred embodiments, the photosynthetic organism is selected from the group consisting of a mesophile, a thermophile and a psychrophile.

5 According to still further features in the described preferred embodiments, the temperature range comprises 10 °C – 43 °C.

According to still further features in the described preferred embodiments, the temperature range comprises an upwards shift.

According to still further features in the described preferred embodiments, the upwards shift is by at least 6 °C.

10 According to still further features in the described preferred embodiments, the temperature range comprises a downwards shift.

According to still further features in the described preferred embodiments, the downwards shift is by at least 2.5 °C.

15 According to still further features in the described preferred embodiments, the temperature range comprise a wider temperature range.

According to still further features in the described preferred embodiments, an amino acid at position 209 or 212 is selected from the group consisting of glycine, cysteine, alanine, threonin, asparagine, aspartanine, proline, valine, isoleucine, leucine, glutamine and glutamic acid.

20 According to still further features in the described preferred embodiments, the amino acid sequence is selected from the group consisting of SEQ ID NO: 37, 38, 50, 52 and 56.

25 According to an additional aspect of the present invention there is provided a photosynthetic organism comprising an exogenous nucleic acid sequence encoding a polypeptide of a photosystem II reaction center of the photosynthetic organism, the nucleic acid sequence imparting the photosystem II reaction center with an activity under a temperature range different than that of the photosystem II reaction center endogenous to the photosynthetic organism.

30 According to still further features in the described preferred embodiments, the photosynthetic organism is selected from the group consisting of a higher plant, a cyanobacteria and an algae.

According to still further features in the described preferred embodiments, the photosynthetic organism is selected from the group consisting of a mesophile, a thermophile and a psychrophile.

5 According to still further features in the described preferred embodiments, the exogenous nucleic acid sequence further renders the organism more thermoplastic than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence.

10 According to still further features in the described preferred embodiments, the exogenous nucleic acid sequence further renders the organism capable of expressing more Rubisco than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence.

15 According to still further features in the described preferred embodiments, the exogenous nucleic acid sequence further renders the organism capable of expressing more psuC than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence.

According to still further features in the described preferred embodiments, the exogenous nucleic acid sequence further renders the organism capable of growing faster than a photosynthetic organism not comprising the exogenous nucleic acid sequence.

20 According to still further features in the described preferred embodiments, the exogenous nucleic acid sequence further renders the organism capable of accumulating more chlorophyll than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence.

25 According to yet an additional aspect of the present invention there is provided a method of improving biomass/vigor/yield of a photosynthetic organism comprising introducing into the photosynthetic organism the isolated polynucleotide, thereby increasing biomass, vigor and/or yield of the photosynthetic organism.

30 According to still further features in the described preferred embodiments, the method further comprising growing the photosynthetic organism under abiotic stress conditions.

According to still further features in the described preferred embodiments, the abiotic stress conditions comprise heat, cold and alternating temperatures.

According to still an additional aspect of the present invention there is provided a method of increasing activity of a photosystem II reaction center of a photosynthetic organism under non-physiological temperature, the method comprising introducing into the photosynthetic organism the isolated polynucleotide, thereby increasing the activity of a photosystem II reaction center under a non-physiological temperature.

According to a further aspect of the present invention there is provided a method of identifying mutations which impart a photosynthetic organism with a photosynthetic activity under a non-physiological temperature, the method comprising: subjecting the photosynthetic organisms to a mutagen, so as to obtain mutated photosynthetic organisms; identifying an organism of the mutated photosynthetic organisms exhibiting at least one parameter associated with enhanced photosynthetic activity under non-physiological temperature as compared to corresponding wild type photosynthetic organisms; and identifying mutations in a polypeptide of a photosystem II reaction center of the organism, the mutations being correlated with photosynthetic activity under a non-physiological temperature.

According to still further features in the described preferred embodiments, the at least one parameter associated with enhanced photosynthetic activity is selected from the group consisting of biomass, growth rate, rubisco level, chlorophyll content, psaC level and thermoplasticity.

According to still further features in the described preferred embodiments, the polypeptide of a photosystem II reaction center is selected from the group consisting of D1, D2, L and M.

According to yet a further aspect of the present invention there is provided a method of increasing hydrogen production in a photosynthetic organism, the method comprising introducing into the photosynthetic organism capable of generating hydrogen the isolated polynucleotide, thereby increasing hydrogen production in the photosynthetic organism.

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, suitable methods and materials are described below. In case of conflict, the patent

specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

As used herein, the terms “comprising” and “including” or grammatical variants thereof are to be taken as specifying the stated features, integers, steps or components but do not preclude the addition of one or more additional features, integers, steps, components or groups thereof. This term encompasses the terms "consisting of" and "consisting essentially of".

The phrase "consisting essentially of" or grammatical variants thereof when used herein are to be taken as specifying the stated features, integers, steps or components but do not preclude the addition of one or more additional features, integers, steps, components or groups thereof but only if the additional features, integers, steps, components or groups thereof do not materially alter the basic and novel characteristics of the claimed composition, device or method.

The term "method" refers to manners, means, techniques and procedures for accomplishing a given task including, but not limited to, those manners, means, techniques and procedures either known to, or readily developed from known manners, means, techniques and procedures by practitioners of the chemical, biological and biophysical arts.

BRIEF DESCRIPTION OF THE DRAWINGS

The invention is herein described, by way of example only, with reference to the accompanying drawings. With specific reference now to the drawings in detail, it is stressed that the particulars shown are by way of example and for purposes of illustrative discussion of the preferred embodiments of the present invention only, and are presented in the cause of providing what is believed to be the most useful and readily understood description of the principles and conceptual aspects of the invention. In this regard, no attempt is made to show structural details of the invention in more detail than is necessary for a fundamental understanding of the invention, the description taken with the drawings making apparent to those skilled in the art how the several forms of the invention may be embodied in practice.

In the drawings:

FIGs. 1A-B are graphs depicting conserved structural and functional elements in photosystem II reaction centers (PTII RCs). Figure 1A depicts (by the broken arrows) light induces electron transfer (ET) from a special pair of chlorophylls (P_{D1} P_{D2} , green) through an accessory chlorophyll (Chl_{D1} , yellow) and pheophytin ($Pheo_{D1}$, purple) to quinones (Q_A and Q_B , red) cofactors held by the D1 (magenta) and D2 (blue) subunits. The non-heme iron (pink) and the atoms lining the three main cavities (orange) are depicted; Figure 1B depicts potential intersubunit hydrogen bonding (ISHB, dashed lines) between helices *D* and *E* of the two subunits. Residues involved in these bonds, e.g. D1-C212 (sulfur marked with a star) are depicted. For clarity, only the two largest cavities are drawn. The arrangement of the protein subunits L and M in purple bacteria well overlap with that of the D1 and D2 subunits, respectively, and the same holds for the chlorophyll and quinone cofactors [Kerner-Shlik et, *Nature*, 2006].

FIG. 1C is a table depicting sequence alignment of the *D* helices in reaction centers from photosystem II (D1 and D2) and from purple non sulfur bacteria and thermophilic green non sulfur bacteria (M and L). The conserved GxxxG-like sequence motif (were the G stands for the small amino acid residues Gly, Ala, Cys, Ser, Thr, gray shading) is found in the center of the *D* helix. Sequences were taken from the Swissprot database (or PDB and NCBI when available). Inter subunit hydrogen bond (ISHB) donor and acceptor residues are highlighted in bold. The alignment of the protein subunits is based on their structure and sequence homologies

as previously described in the literature [Deisenhofer et al., *JMB* 246, 429-457 (1995)]. Of note, the GxxxG-like motifs in the protein subunits of the purple bacteria is shifted by one site compared with their position in photosystem II.

FIGs. 2A-B are graphs depicting the $Q_A^- \rightarrow Q_B$ ET rate in mesophilic and thermophilic cyanobacteria. Figure 2A are Eyring plots for ET in a mesophile (*Synechocystis* 6803, closed circles) and a thermophile (*Thermosynechococcus elongatus*, open circles); Figure 2B are Eyring plots for ET in D1-212 class I mutants Cys (triangles), and Ala (diamonds) compared to the *Synechocystis* 6803 Ser wild-type (closed circles).

FIGs. 3A-B are graphs depicting the effect of D1-212 residues on the photosystem II RC cavity in *Thermosynechococcus elongatus* structure. Figure 3A depicts class I residue (Ser, space-fill representation of side-chain) maintains the cavity open; Figure 3B depicts class II residue (Gln) abolishes one of the cavities and lines the largest cavity that is otherwise lined by flexible side-chains.

FIGs. 4A-B are graphs depicting temperature dependence and activation parameters of the $Q_A^- \rightarrow Q_B$ ET in D1-212 mutants of *Synechocystis* 6803. Figure 4A are Eyring plots for D1-S212 (closed circles), D1-V212 (open squares) and D1-Q212 (closed squares); Figure 4B depicts correlation between the activation entropy and enthalpy for the $Q_A^- \rightarrow Q_B$ ET and the packing values of residues at the D1-212 site. Class I residues (closed circles) are distinct from the class II residues (closed squares).

FIGs. 5A-E are graphs depicting structural Motifs within the photosystem II RCs. Figure 5A depicts ISHBs between the D and E helices of the D1 and D2 subunits of the photosystem II RC. This figure zooms into Figure 1B (detailed hereinabove) excluding for clarity the cartoon diagram of the D1 and D2 protein subunits as well as the cavities. Residues participating in transmembrane domain ISHBs (see Figure 1C) are marked; Figure 5B depicts ISHBs involving D1-212 and immediate vicinity. Viewing from the D2 subunit viewpoint it is shown that the D1-C212 SG atom (marked with an ' * ') can act as both an ISHB acceptor and a donor. As a donor, the hydrogen must "choose" between two alternative H-bond options. Atoms participating in ISHBs are marked; Figures C-E depict intra-protein cavities found in the transmembrane region between the D1 and D2 subunits of photosystem II RC. The largest three cavities (see Table 2) form a continuous area of high flexibility from just below the histidines ligating the non-heme iron and virtually till the center of

mass of this protein complex. The atoms lining these cavities (Figure 5C) are equally distributed between the D1 and D2 subunits (space-fill, color by chain color) and consist solely of helices D and E. The largest cavity is located at the center of the four-helix-bundle (Figure 5D) as shown from a top view. A focus on the cavities and the residues participating in ISHBs (Figure 5E) demonstrates that these two structural features are intimately entwined.

FIG. 6 is a graph depicting the amplitude of the shortest life-time component of the photosystem II RC fluorescence obtained at different temperatures as explained in the Methods section hereinbelow. Wild type fluorescence (D1- S212, full circles), class I mutants Val (open triangles) and Gln (open circles).

FIG. 7 is a graph depicting the Eyring plots for $Q_A^- \rightarrow Q_B$ ET in wild type (Δ KS, open circles), single mutants D1-212 Cys 212 (filled diamonds) and Pro 212 (triangles) and double mutants D1-209/212 SS209/212AA (open squares) and SS209/212AC (closed squares). The T_o values are depicted for individual strains.

FIGS. 8A-L are graphs depicting the temperature effect on wild type (Δ KS) and the different mutants. FIG. 8A shows the colors of the different liquid cultures after incubation at 30 °C for six days; FIG. 8B shows the colors of the different liquid cultures after incubation at 43 °C for six days; FIG. 8C is a graph showing growth at 30 °C measured as turbidity at 730 nm; FIG. 8D is a graph showing growth at 43°C measured as turbidity at 730 nm; FIG. 8E is a graph showing chlorophyll *a* accumulation rates in cell cultures grown at 30 °C; FIG. 8F is a graph showing chlorophyll *a* accumulation rates in cell cultures grown at 43 °C; FIG. 8G is an immunoblot showing the levels of D1 protein in cell cultures grown at 30 °C; FIG. 8H is an immunoblot showing the levels of D1 protein in cell cultures grown at 43 °C; FIG. 8I is an immunoblot showing the levels of the Rubisco large protein subunit in cell cultures grown at 30 °C. FIG. 8J is an immunoblot showing the levels of the Rubisco large protein subunit in cell cultures grown at 43 °C; FIG. 8K is an immunoblot showing the levels of psaC protein in cell cultures grown at 30 °C; and FIG. 8L is an immunoblot showing the levels of psaC protein in cell cultures grown at 43 °C.

FIGS. 9A-L are graphs depicting the thermoplasticity of the wild type (Δ KS) and the different mutants under different temperature cycles (43 °C – 30 °C -43 °C or 43 °C – 10 °C – 43 °C). FIG. 9A shows the colors of the different liquid cultures after

incubation at 43 °C – 30 °C -43 °C; FIG. 9B shows the colors of the different liquid cultures after incubation at 43 °C – 10 °C – 43 °C; FIG. 9C is a graph showing growth at 43 °C – 30 °C -43 °C measured as turbidity at 730 nm; FIG. 9D is a graph showing growth at 43 °C – 10 °C – 43 °C measured as turbidity at 730 nm; FIG. 9E is a graph showing chlorophyll *a* accumulation rates in cell cultures grown at 43 °C – 30 °C -43 °C; FIG. 9F is a graph showing chlorophyll *a* accumulation rates in cell cultures grown at 43 °C – 10 °C – 43 °C; FIG. 9G is an immunoblot showing the levels of D1 protein in cell cultures grown at 43 °C – 30 °C -43 °C; FIG. 9H is an immunoblot showing the levels of D1 protein in cell cultures grown at 43 °C – 10 °C – 43 °C; FIG. 9I is an immunoblot showing the levels of the Rubisco large protein subunit in cell cultures grown at 43 °C – 30 °C -43 °C; FIG. 9J is an immunoblot showing the levels of the Rubisco large protein subunit in cell cultures grown at 43 °C – 10 °C – 43 °C; FIG. 9K is an immunoblot showing the levels of *psaC* protein in cell cultures grown at 43 °C – 30 °C -43 °C; and FIG. 9L is an immunoblot showing the levels of *psaC* protein in cell cultures grown at 43 °C – 10 °C – 43 °C.

FIGs. 10A-F are graphs depicting the thermoplasticity of the wild type (Δ KS) and the different *Synechocystis* 6803 mutants and wild type cultures under 10 °C - 43 °C -10 °C cycle. FIG. 10A shows the colors of the different liquid cultures after incubation; FIG. 10B is a graph showing growth measured as turbidity at 730 nm; FIG. 10C is a graph showing chlorophyll *a* accumulation rates in cell cultures; FIG. 10D is an immunoblot showing the levels of D1 protein in cell cultures; FIG. 10E is an immunoblot showing the levels of the Rubisco large protein subunit in cell cultures; and FIG. 10F is an immunoblot showing the levels of *psaC* protein in cell cultures.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention is of photosynthetic organisms and, compositions and methods of generating same. Specifically, the present invention can be used to render a mesophile photosynthetic organism more thermotolerant enabling high yield of photosynthetic energy conversion at higher temperatures than the physiological temperature environment of the mesophilic organism.

The principles and operation of the present invention may be better understood with reference to the drawings and accompanying descriptions.

Before explaining at least one embodiment of the invention in detail, it is to be understood that the invention is not limited in its application to the details set forth in the following description or exemplified by the Examples. The invention is capable of other embodiments or of being practiced or carried out in various ways. Also, it is to be understood that the phraseology and terminology employed herein is for the purpose of description and should not be regarded as limiting.

Whilst conceiving the present invention, the present inventors uncovered a role of a specific amino acid sequence motif located at the D1/D2 interface of photosystem II reaction center, which contributes to functional flexibility and temperature adaptation of photosystem II reaction center (RC) in all photosynthetic organisms. This finding allowed the generation of photosynthetic organisms with "tailor-made" thermotolerance.

Thus, based on these novel findings it is now possible to increase the thermostability and thermoplasticity of photosystem II reaction center, thereby shifting maximal photosynthetic rates of the genetically modified organisms to higher temperatures, consequently enabling higher growth and/or biomass-generating rates compared with the wild type in non-optimal temperatures, e.g. by achieving a closer-to-optimum growth rates under nonphysiological conditions where such rates are suppressed. In addition, the present teachings enable photosynthetic light conversion in PSII and biomass production at wider range of temperatures, above and below that presented by wild type mesophilic strain. Furthermore, the present teachings allow maintenance of high productivity and activity of the Rubisco complex at such wider range of temperatures.

As is illustrated hereinbelow and in the Examples section which follows, whilst searching for motifs that account for the temperature adaptation of PSII RC, the present inventors examined the sequences of the two major protein subunits found in all purple bacteria and photosystem II RCs and found two sites, D1-209 and D1-212, that undergo consistent changes between mesophilic, thermotolerant and thermophilic organisms including cyanobacteria, algae and green plants (Example 1). The sites are positioned in a GxxxG-like sequence motif (where 'G' denotes small residues such as Gly, Ala, Ser, Cys and Thr) found at the closest contact of the two major protein subunits (D1 and D2 and corresponding L and M in purple bacteria for example). This motif and the structurally homologous motif in purple bacteria RC (Example 1)

participates in an intersubunit hydrogen bonding (ISHB) network probably providing local flexibility while maintaining a protein overall stability. Thus, the present invention contemplates amino acid alterations within the identified GxxxG-like motif for modifying the RC localized flexibility and consequently, changing the electron transfer rate constant. Two sizable cavities augmented by several small ones complement the ISHB cluster (Example 1). Such cavities in the inner core of a protein may facilitate the conformational rearrangement required for enhancing the protein's local flexibility during discrete functional steps. Relying on the above understanding the present inventors have modified D1 of *Synechocystis* sp. 6803 in the identified domain to thereby shift the maximum PSII RC energy conversion rate of the mesophilic cyanobacteria to higher temperatures as observed when comparing mutated mesophiles with thermotolerant and thermophilic bacteria (Examples 2 and 3). Importantly, this shift is independent of the proteins and whole cell thermostability. The present inventors have further shown that some single or double mutation of the identified GxxxG motif grant thermostability over the requested temperature range (45 °C - 10 °C - 45 °C) while maintaining high photosynthetic productivity to mesophilic cyanobacteria (Examples 6, and 7). Since the observed GxxxG-like motif is conserved in all photosynthetic organisms (Example 1) the present teachings can be easily applied to algae, microalgae and plants to thereby establish new energy, biomass and food resources. The present inventors have further uncovered that the modified thermotolerant organisms maintain Rubisco content at higher temperature than the physiological temperature of the wild type, thus assuring carbone-dioxide fixation and biomass generation (Example 4, 5 and 7). The increased stability and functionality of PSII at high temperatures should also enable provision of thermoplasticity and enable crop growth and production in hot or warming climates.

Thus, according to one aspect of the present invention there is provided an isolated polynucleotide comprising a nucleic acid sequence encoding a polypeptide of a photosystem II reaction center of a photosynthetic organism, said nucleic acid sequence being capable of imparting said photosystem II reaction center with an activity under a temperature range (e.g., 10-50 °C, 10-43 °C or 30- 43 °C) different than that of said type II reaction center endogenous to said photosynthetic organism.

As used herein the phrase "photosynthetic organism" refers to an organism (prokaryote or eukaryote) capable of converting light energy and a carbon source (e.g., carbon dioxide) to triose phosphates.

5 The photosynthetic organism of this aspect of the present invention can be a mesophile, a thermophile and a psychrophile.

As used herein the phrase "mesophile organism" refers to an organism with a physiological growth temperature at a range of about 15-35 °C.

As used herein the phrase "thermotolerant organism" refers to an organism with a physiological growth temperature at a range of about 35-45 °C.

10 As used herein the phrase "thermophile organism" refers to an organism with a physiological growth temperature at a range of about 45-70 °C.

As used herein the phrase " psychrophile organism" refers to an organism with a physiological growth temperature at a range of about 0-15 °C.

15 Thus, the photosynthetic organism of this aspect of the present invention refers to a bacteria (aerobic and anaerobic), an algae (prokaryotic and eukaryotic) and a higher plant.

Examples of photosynthetic bacteria include without limitation cyanobacteria e.g. *Synechocystis* sp. PCC 6803, *Prochlorothrix hollandica*, *Thermosynechococcus elongatus*, purple non sulfur e.g., *Rhodospirillum rubrum* *Rhodobacter sphaeroides*,
20 *Rhodopseudomonas viridis* and green non sulfur photosynthetic bacteria e.g., *Chloroflexus aurantiacus*.

Examples of algae include without limitation green algae e.g., *Chlamydomonas reinhardtii*, *Chlorella vulgaris* and red algae e.g. *Cyanidium caldarium*, *Porphyridium purpureum*.

25 Examples of plants [The term "plant" as used herein encompasses whole plants, ancestors and progeny of the plants and plant parts, including seeds, shoots, stems, roots (including tubers), and plant cells, tissues and organs. The term "plant" also therefore encompasses suspension cultures, embryos, meristematic regions, callus tissue, leaves, etc.]. include in particular monocotyledonous and dicotyledonous plants
30 which are of commercial value, including a fodder or forage legume, ornamental plant, food crop, tree, or shrub selected from the following non-limiting list comprising maize, sweet potato, tubers such as cassarva, sugar beet, wheat, barely, rye, oat, rice, soybean, peanut, pea, cowpea, lentil and alfalfa, cotton, rapeseed, canola, pepper,

sunflower, potato, tobacco, tomato, eggplant, trees such as eucalyptus and poplars, an ornamental plant, a perennial grass and a forage crop.

As used herein the phrase "photosystem II reaction center" or a "type II reaction center" refers to eukaryotic and prokaryotic type II reaction centers. The photochemical reaction center (RC) of photosystem II (PSII, the D1-D2 cyt.*b559* complex) is the smallest unit in PSII that shows photochemical activity. The RC contains 8 chlorophyll (Chl) *a* and two pheophytin (Pheo) *a* molecules that all have their lowest electronic transition around 675 nm, as well as two β -carotenes. The D1 and D2 polypeptides are homologous to the L and M subunits of bacterial RCs, suggesting an arrangement of the core pigments in the RC of PSII similar to that in the bacterial RC. The two additional Chl molecules are probably located near the periphery of the D1-D2 complex.

As used herein the phrase "activity of a photosystem II reaction center" refers to a solar energy conversion activity which is essential for the photosynthetic organism. A number of proxies can be used to determine the activity of the photosystem II reaction center. Examples include, but are not limited to, growth rate, viability, chlorophyll concentration, Q_A - Q_B electron transfer rate and Rubisco levels. Assays for qualification/quantification are described at length in the Examples section which follows.

As used herein the phrase "a polypeptide of a photosystem II reaction center" refers to D1, D2, L and M (see SEQ ID NOs. 30-34 and 36). Exemplary GenBank Accession Numbers are provided in Figure 1C.

As mentioned herein above, the present inventors have identified a nucleic acid sequence motif GXXXG (corresponding to positions 208-212 of SEQ ID NO: 36), on D1 (located at the D1/D2 interface) which contributes to the functional flexibility and temperature adaptation of photosystem II reaction center.

Thus, according to an exemplary embodiment of this aspect of the present invention the nucleic acid sequence encodes a polypeptide having an amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 and at a position corresponding to 212 of SEQ ID NO: 36 (e.g., see SEQ ID NOs. 37-61).

According to other exemplary embodiments the isolated polypeptide sequences of D2, L and M polypeptides SEQ ID NO: 30 – SEQ ID NO: 34 with

suggested sites for mutations for D2 polypeptide at a positions corresponding to 207 and 211 in *Synechocystis sp* 6803 (SEQ ID NO: 30); for M polypeptide at positions corresponding to 211 and 215 in *Rhodopseudomonas viridis* (SEQ ID NO: 31) and 213 and 217 in *Rhodobacter sphaeroides* (SEQ ID NO: 32); for L polypeptide at positions corresponding to 184 and 188 in both *Rh.viridis* and *Rh.sphaeroides* (SEQ ID NOs. 33 and 34).

Thus, the present inventors have shown that *Synechocystis sp.* 6803 modified with a polynucleotide of the present invention encoding a D1 polypeptide with a serine to alanine substitution at a position corresponding to 209 of SEQ ID NO: 36 and/or a serine to cysteine substitution at a position corresponding to 212 of SEQ ID NO: 36 imparts the modified photosystem II RC with an activity under non-physiological temperature [e.g., upwards shift in temperature conditions of (e.g., 30>43 °C)].

As used herein the phrase "physiological conditions" refers to a temperature range which mediates optimal growth (e.g., +/- 50 %) of the naïve (non-genetically modified) organism.

Thus, according to one embodiment of this aspect of the present invention the isolated polynucleotide (or an expression product of same i.e., the isolated polypeptide) of this aspect of the present invention is capable of rendering the photosystem II reaction center which comprises same more thermotolerant as compared to a non-modified reaction center.

According to another embodiment of this aspect of the present invention the isolated polynucleotide (or an expression product of same i.e., the isolated polypeptide) of this aspect of the present invention is capable of rendering the photosystem II reaction center which comprises same more thermoplastic as compared to a corresponding non-modified reaction center.

According to yet another embodiment of this aspect of the present invention, the polynucleotide is capable of imparting the photosystem II reaction center with an activity at an upwards shifted range (e.g., at least 4 °C, 6 °C, 7 °C or 10 °C higher than that of a corresponding non-modified type II reaction center).

According to a still embodiment of this aspect of the present invention, the polynucleotide is capable of imparting the photosystem II reaction center with an activity at an downwards shifted range (e.g., at least 2.5 °C or 4 °C lower than that of a corresponding non-modified type II reaction center).

According to a further still embodiment of this aspect of the present invention, the polynucleotide is capable of imparting the photosystem II reaction center with an activity at an downwards shifted range (e.g., at least 2.5 °C or 4 °C lower than that of a corresponding non-modified type II reaction center).

5 According to a further still embodiment of this aspect of the present invention, the polynucleotide is capable of imparting the photosystem II reaction center with biomass productivity at a wider temperature range (e.g., at least 10 °C wider than that of a corresponding non-modified type II reaction center).

The nucleic acid sequence (also termed herein as "isolated polynucleotide") of
10 the present invention refers to a single or double stranded nucleic acid sequence which is isolated and provided in the form of an RNA sequence, a complementary polynucleotide sequence (cDNA), a genomic polynucleotide sequence and/or a composite polynucleotide sequences (e.g., a combination of the above).

As used herein the phrase "complementary polynucleotide sequence" refers to
15 a sequence, which results from reverse transcription of messenger RNA using a reverse transcriptase or any other RNA dependent DNA polymerase. Such a sequence can be subsequently amplified *in vivo* or *in vitro* using a DNA dependent DNA polymerase.

As used herein the phrase "genomic polynucleotide sequence" refers to a
20 sequence derived (isolated) from a chromosome and thus it represents a contiguous portion of a chromosome.

As used herein the phrase "composite polynucleotide sequence" refers to a
25 sequence, which is at least partially complementary and at least partially genomic. A composite sequence can include some exonal sequences required to encode the polypeptide of the present invention, as well as some intronic sequences interposing therebetween. The intronic sequences can be of any source, including of other genes, and typically will include conserved splicing signal sequences. Such intronic sequences may further include cis acting expression regulatory elements.

Nucleic acid sequences of the polypeptides of the present invention may be
30 optimized for plant expression. Examples of such sequence modifications include, but are not limited to, an altered G/C content to more closely approach that typically found in the plant species of interest, and the removal of codons atypically found in the plant species commonly referred to as codon optimization.

The phrase "codon optimization" refers to the selection of appropriate DNA nucleotides for use within a structural gene or fragment thereof that approaches codon usage within the plant of interest. Therefore, an optimized gene or nucleic acid sequence refers to a gene in which the nucleotide sequence of a native or naturally occurring gene has been modified in order to utilize statistically-preferred or statistically-favored codons within the plant. The nucleotide sequence typically is examined at the DNA level and the coding region optimized for expression in the plant species determined using any suitable procedure, for example as described in Sardana *et al.* (1996, Plant Cell Reports 15:677-681). In this method, the standard deviation of codon usage, a measure of codon usage bias, may be calculated by first finding the squared proportional deviation of usage of each codon of the native gene relative to that of highly expressed plant genes, followed by a calculation of the average squared deviation. The formula used is: $1 \text{ SDCU} = n = 1/N \sum [(X_n - Y_n) / Y_n]^2 / N$, where X_n refers to the frequency of usage of codon n in highly expressed plant genes, where Y_n to the frequency of usage of codon n in the gene of interest and N refers to the total number of codons in the gene of interest. A table of codon usage from highly expressed genes of dicotyledonous plants is compiled using the data of Murray *et al.* (1989, Nuc Acids Res. 17:477-498).

One method of optimizing the nucleic acid sequence in accordance with the preferred codon usage for a particular plant cell type is based on the direct use, without performing any extra statistical calculations, of codon optimization tables such as those provided on-line at the Codon Usage Database through the NIAS (National Institute of Agrobiological Sciences) DNA bank in Japan (<http://www.kazusa.or.jp/codon/>). The Codon Usage Database contains codon usage tables for a number of different species, with each codon usage table having been statistically determined based on the data present in Genbank.

By using the above tables to determine the most preferred or most favored codons for each amino acid in a particular species (for example, rice), a naturally-occurring nucleotide sequence encoding a protein of interest can be codon optimized for that particular plant species. This is effected by replacing codons that may have a low statistical incidence in the particular species genome with corresponding codons, in regard to an amino acid, that are statistically more favored. However, one or more less-favored codons may be selected to delete existing restriction sites, to create new

ones at potentially useful junctions (5' and 3' ends to add signal peptide or termination cassettes, internal sites that might be used to cut and splice segments together to produce a correct full-length sequence), or to eliminate nucleotide sequences that may negatively effect mRNA stability or expression.

5 The naturally-occurring encoding nucleotide sequence may already, in advance of any modification, contain a number of codons that correspond to a statistically-favored codon in a particular plant species. Therefore, codon optimization of the native nucleotide sequence may comprise determining which codons, within the native nucleotide sequence, are not statistically-favored with
10 regards to a particular plant, and modifying these codons in accordance with a codon usage table of the particular plant to produce a codon optimized derivative. A modified nucleotide sequence may be fully or partially optimized for plant codon usage provided that the protein encoded by the modified nucleotide sequence is produced at a level higher than the protein encoded by the corresponding naturally
15 occurring or native gene. Construction of synthetic genes by altering the codon usage is described in for example PCT Patent Application 93/07278.

Thus, the present invention encompasses nucleic acid sequences described hereinabove; fragments thereof, sequences hybridizable therewith, sequences homologous thereto, sequences encoding similar polypeptides with different codon
20 usage, altered sequences characterized by mutations, such as deletion, insertion or substitution of one or more nucleotides, either naturally occurring or man induced, either randomly or in a targeted fashion.

Since the present invention provides previously unknown polypeptides, the present invention also encompasses polypeptide sequences, fragments of the above
25 described polypeptides and polypeptides having mutations, such as deletions, insertions or substitutions of one or more amino acids, either naturally occurring or man induced, either randomly or in a targeted fashion.

Methods of identifying mutations which impart a photosynthetic organism with a photosynthetic activity under a non-physiological temperature are outlined infra. Basically, subjecting the photosynthetic organisms to a mutagen, so as to obtain mutated photosynthetic organisms. Identifying an organism of said mutated photosynthetic organisms exhibiting at least one parameter associated with enhanced photosynthetic activity under non-physiological temperature as compared to

corresponding wild type photosynthetic organisms. Identifying mutations in a polypeptide of a photosystem II reaction center of said organism, said mutations being correlated with photosynthetic activity under a non-physiological temperature.

Exemplary parameters which are associated with photosynthetic activity include, but are not limited to, biomass, growth rate, rubisco level, chlorophyll content, psaC level and thermoplasticity.

Polynucleotides and polypeptides of the present invention are used for expression in a cell of a photosynthetic organism.

The expression system will naturally depend on the organism.

5 Methods of expressing the nucleic acid sequences of this aspect of the present invention in photosynthetic bacteria are well known in the art and described at length in the Examples section which follows. Generally these include site-directed mutagenesis and methods of transgenesis.

10 Methods of expression of exogenous polynucleotides in microalgae are well known in the art. See for example WO2006/013572 incorporated herein by reference in its entirety.

Expressing the exogenous polynucleotide of the present invention within higher plants can be effected by transforming one or more cells of the plant with the exogenous polynucleotide, followed by generating a mature plant from the transformed cells and cultivating the mature plant under conditions suitable for
15 expressing the exogenous polynucleotide within the mature plant.

Preferably, the transformation is effected by introducing to the plant cell a nucleic acid construct which includes the exogenous polynucleotide of the present invention and at least one promoter capable of directing transcription of the exogenous polynucleotide in the plant cell. Further details of suitable transformation approaches
20 are provided hereinbelow.

As used herein, the term "promoter" refers to a region of DNA which lies upstream of the transcriptional initiation site of a gene to which RNA polymerase binds to initiate transcription of RNA. The promoter controls where (e.g., which portion of a plant, which organ within an animal, etc.) and/or when (e.g., which stage
25 or condition in the lifetime of an organism) the gene is expressed.

Any suitable promoter sequence can be used by the nucleic acid construct of the present invention. Preferably the promoter is a constitutive promoter.

Suitable constitutive promoters include, for example, CaMV 35S promoter.

Suitable tissue-specific promoters include, but not limited to, leaf-specific promoters such as described, for example, by Yamamoto *et al.*, Plant J. 12:255-265, 1997; Kwon *et al.*, Plant Physiol. 105:357-67, 1994; Yamamoto *et al.*, Plant Cell Physiol. 35:773-778, 1994; Gotor *et al.*, Plant J. 3:509-18, 1993; Orozco *et al.*, Plant Mol. Biol. 23:1129-1138, 1993; and Matsuoka *et al.*, Proc. Natl. Acad. Sci. USA 90:9586-9590, 1993.

The nucleic acid construct of the present invention preferably further includes an appropriate selectable marker and/or an origin of replication. Preferably, the nucleic acid construct utilized is a shuttle vector, which can propagate both in *E. coli* (wherein the construct comprises an appropriate selectable marker and origin of replication) and be compatible for propagation in cells. The construct according to the present invention can be, for example, a plasmid, a bacmid, a phagemid, a cosmid, a phage, a virus or an artificial chromosome.

The nucleic acid construct of the present invention can be utilized to stably or transiently transform plant cells. In stable transformation, the exogenous polynucleotide of the present invention is integrated into the plant genome and as such it represents a stable and inherited trait. In transient transformation, the exogenous polynucleotide is expressed by the cell transformed but it is not integrated into the genome and as such it represents a transient trait.

There are various methods of introducing foreign genes into both monocotyledonous and dicotyledonous plants (Potrykus, I., Annu. Rev. Plant. Physiol., Plant. Mol. Biol. (1991) 42:205-225; Shimamoto *et al.*, Nature (1989) 338:274-276).

The principle methods of causing stable integration of exogenous DNA into plant genomic DNA include two main approaches:

(i) *Agrobacterium*-mediated gene transfer: Klee *et al.* (1987) Annu. Rev. Plant Physiol. 38:467-486; Klee and Rogers in Cell Culture and Somatic Cell Genetics of Plants, Vol. 6, Molecular Biology of Plant Nuclear Genes, eds. Schell, J., and Vasil, L. K., Academic Publishers, San Diego, Calif. (1989) p. 2-25; Gatenby, in Plant Biotechnology, eds. Kung, S. and Arntzen, C. J., Butterworth Publishers, Boston, Mass. (1989) p. 93-112.

(ii) Direct DNA uptake: Paszkowski *et al.*, in Cell Culture and Somatic Cell Genetics of Plants, Vol. 6, Molecular Biology of Plant Nuclear Genes eds. Schell, J., and Vasil, L. K., Academic Publishers, San Diego, Calif. (1989) p. 52-68; including methods for direct uptake of DNA into protoplasts, Toriyama, K. *et al.* (1988) 5 Bio/Technology 6:1072-1074. DNA uptake induced by brief electric shock of plant cells: Zhang *et al.* Plant Cell Rep. (1988) 7:379-384. Fromm *et al.* Nature (1986) 319:791-793. DNA injection into plant cells or tissues by particle bombardment, Klein *et al.* Bio/Technology (1988) 6:559-563; McCabe *et al.* Bio/Technology (1988) 6:923-926; Sanford, Physiol. Plant. (1990) 79:206-209; by the use of micropipette 10 systems: Neuhaus *et al.*, Theor. Appl. Genet. (1987) 75:30-36; Neuhaus and Spangenberg, Physiol. Plant. (1990) 79:213-217; glass fibers or silicon carbide whisker transformation of cell cultures, embryos or callus tissue, U.S. Pat. No. 5,464,765 or by the direct incubation of DNA with germinating pollen, DeWet *et al.* in Experimental Manipulation of Ovule Tissue, eds. Chapman, G. P. and Mantell, S. 15 H. and Daniels, W. Longman, London, (1985) p. 197-209; and Ohta, Proc. Natl. Acad. Sci. USA (1986) 83:715-719.

The *Agrobacterium* system includes the use of plasmid vectors that contain defined DNA segments that integrate into the plant genomic DNA. Methods of inoculation of the plant tissue vary depending upon the plant species and the 20 *Agrobacterium* delivery system. A widely used approach is the leaf disc procedure which can be performed with any tissue explant that provides a good source for initiation of whole plant differentiation. Horsch *et al.* in Plant Molecular Biology Manual A5, Kluwer Academic Publishers, Dordrecht (1988) p. 1-9. A supplementary approach employs the *Agrobacterium* delivery system in combination with vacuum 25 infiltration. The *Agrobacterium* system is especially viable in the creation of transgenic dicotyledonous plants.

There are various methods of direct DNA transfer into plant cells. In electroporation, the protoplasts are briefly exposed to a strong electric field. In microinjection, the DNA is mechanically injected directly into the cells using very 30 small micropipettes. In microparticle bombardment, the DNA is adsorbed on microprojectiles such as magnesium sulfate crystals or tungsten particles, and the microprojectiles are physically accelerated into cells or plant tissues.

Following stable transformation plant propagation is exercised. The most common method of plant propagation is by seed. Regeneration by seed propagation, however, has the deficiency that due to heterozygosity there is a lack of uniformity in the crop, since seeds are produced by plants according to the genetic variances governed by Mendelian rules. Basically, each seed is genetically different and each will grow with its own specific traits. Therefore, it is preferred that the transformed plant be produced such that the regenerated plant has the identical traits and characteristics of the parent transgenic plant. Therefore, it is preferred that the transformed plant be regenerated by micropropagation which provides a rapid, consistent reproduction of the transformed plants.

Micropropagation is a process of growing new generation plants from a single piece of tissue that has been excised from a selected parent plant or cultivar. This process permits the mass reproduction of plants having the preferred tissue expressing the fusion protein. The new generation plants which are produced are genetically identical to, and have all of the characteristics of, the original plant. Micropropagation allows mass production of quality plant material in a short period of time and offers a rapid multiplication of selected cultivars in the preservation of the characteristics of the original transgenic or transformed plant. The advantages of cloning plants are the speed of plant multiplication and the quality and uniformity of plants produced.

Micropropagation is a multi-stage procedure that requires alteration of culture medium or growth conditions between stages. Thus, the micropropagation process involves four basic stages: Stage one, initial tissue culturing; stage two, tissue culture multiplication; stage three, differentiation and plant formation; and stage four, greenhouse culturing and hardening. During stage one, initial tissue culturing, the tissue culture is established and certified contaminant-free. During stage two, the initial tissue culture is multiplied until a sufficient number of tissue samples are produced to meet production goals. During stage three, the tissue samples grown in stage two are divided and grown into individual plantlets. At stage four, the transformed plantlets are transferred to a greenhouse for hardening where the plants' tolerance to light is gradually increased so that it can be grown in the natural environment.

Preferably, mature transformed plants generated as described above are further selected for increase biomass, alcohol production, vigor and/or yield.

Although stable transformation is presently preferred, transient transformation of leaf cells, meristematic cells or the whole plant is also envisaged by the present invention.

Transient transformation can be effected by any of the direct DNA transfer methods described above or by viral infection using modified plant viruses.

Viruses that have been shown to be useful for the transformation of plant hosts include CaMV, TMV and BV. Transformation of plants using plant viruses is described in U.S. Pat. No. 4,855,237 (BGV), EP-A 67,553 (TMV), Japanese Published Application No. 63-14693 (TMV), EPA 194,809 (BV), EPA 278,667 (BV); and Gluzman, Y. *et al.*, Communications in Molecular Biology: Viral Vectors, Cold Spring Harbor Laboratory, New York, pp. 172-189 (1988). Pseudovirus particles for use in expressing foreign DNA in many hosts, including plants, is described in WO 87/06261.

Preferably, the virus of the present invention is avirulent and thus is incapable of causing severe symptoms such as reduced growth rate, mosaic, ring spots, leaf roll, yellowing, streaking, pox formation, tumor formation and pitting. A suitable avirulent virus may be a naturally occurring avirulent virus or an artificially attenuated virus. Virus attenuation may be effected by using methods well known in the art including, but not limited to, sub-lethal heating, chemical treatment or by directed mutagenesis techniques such as described, for example, by Kurihara and Watanabe (Molecular Plant Pathology 4:259-269, 2003), Gal-on *et al.* (1992), Atreya *et al.* (1992) and Huet *et al.* (1994).

Suitable virus strains can be obtained from available sources such as, for example, the American Type culture Collection (ATCC) or by isolation from infected plants. Isolation of viruses from infected plant tissues can be effected by techniques well known in the art such as described, for example by Foster and Tatlor, Eds. "Plant Virology Protocols: From Virus Isolation to Transgenic Resistance (Methods in Molecular Biology (Humana Pr), Vol 81)", Humana Press, 1998. Briefly, tissues of an infected plant believed to contain a high concentration of a suitable virus, preferably young leaves and flower petals, are ground in a buffer solution (e.g., phosphate buffer solution) to produce a virus infected sap which can be used in subsequent inoculations.

Construction of plant RNA viruses for the introduction and expression of non-viral exogenous polynucleotide sequences in plants is demonstrated by the above references as well as by Dawson, W. O. *et al.*, *Virology* (1989) 172:285-292; Takamatsu *et al.* *EMBO J.* (1987) 6:307-311; French *et al.* *Science* (1986) 5 231:1294-1297; and Takamatsu *et al.* *FEBS Letters* (1990) 269:73-76.

When the virus is a DNA virus, suitable modifications can be made to the virus itself. Alternatively, the virus can first be cloned into a bacterial plasmid for ease of constructing the desired viral vector with the foreign DNA. The virus can then be excised from the plasmid. If the virus is a DNA virus, a bacterial origin of replication 10 can be attached to the viral DNA, which is then replicated by the bacteria. Transcription and translation of this DNA will produce the coat protein which will encapsidate the viral DNA. If the virus is an RNA virus, the virus is generally cloned as a cDNA and inserted into a plasmid. The plasmid is then used to make all of the constructions. The RNA virus is then produced by transcribing the viral sequence of 15 the plasmid and translation of the viral genes to produce the coat protein(s) which encapsidate the viral RNA.

Construction of plant RNA viruses for the introduction and expression in plants of non-viral exogenous polynucleotide sequences such as those included in the construct of the present invention is demonstrated by the above references as well as in 20 U.S. Pat. No. 5,316,931.

In one embodiment, a plant viral polynucleotide is provided in which the native coat protein coding sequence has been deleted from a viral polynucleotide, a non-native plant viral coat protein coding sequence and a non-native promoter, preferably the subgenomic promoter of the non-native coat protein coding sequence, capable of 25 expression in the plant host, packaging of the recombinant plant viral polynucleotide, and ensuring a systemic infection of the host by the recombinant plant viral polynucleotide, has been inserted. Alternatively, the coat protein gene may be inactivated by insertion of the non-native polynucleotide sequence within it, such that a protein is produced. The recombinant plant viral polynucleotide may contain one or 30 more additional non-native subgenomic promoters. Each non-native subgenomic promoter is capable of transcribing or expressing adjacent genes or polynucleotide sequences in the plant host and incapable of recombination with each other and with native subgenomic promoters. Non-native (foreign) polynucleotide sequences may be

inserted adjacent the native plant viral subgenomic promoter or the native and a non-native plant viral subgenomic promoters if more than one polynucleotide sequence is included. The non-native polynucleotide sequences are transcribed or expressed in the host plant under control of the subgenomic promoter to produce the desired products.

5 In a second embodiment, a recombinant plant viral polynucleotide is provided as in the first embodiment except that the native coat protein coding sequence is placed adjacent one of the non-native coat protein subgenomic promoters instead of a non-native coat protein coding sequence.

10 In a third embodiment, a recombinant plant viral polynucleotide is provided in which the native coat protein gene is adjacent its subgenomic promoter and one or more non-native subgenomic promoters have been inserted into the viral polynucleotide. The inserted non-native subgenomic promoters are capable of transcribing or expressing adjacent genes in a plant host and are incapable of recombination with each other and with native subgenomic promoters. Non-native
15 polynucleotide sequences may be inserted adjacent the non-native subgenomic plant viral promoters such that the sequences are transcribed or expressed in the host plant under control of the subgenomic promoters to produce the desired product.

In a fourth embodiment, a recombinant plant viral polynucleotide is provided as in the third embodiment except that the native coat protein coding sequence is
20 replaced by a non-native coat protein coding sequence.

The viral vectors are encapsidated by the coat proteins encoded by the recombinant plant viral polynucleotide to produce a recombinant plant virus. The recombinant plant viral polynucleotide or recombinant plant virus is used to infect appropriate host plants. The recombinant plant viral polynucleotide is capable of
25 replication in the host, systemic spread in the host, and transcription or expression of foreign gene(s) (exogenous polynucleotide) in the host to produce the desired protein.

Techniques for inoculation of viruses to plants may be found in Foster and Taylor, eds. "Plant Virology Protocols: From Virus Isolation to Transgenic Resistance (Methods in Molecular Biology (Humana Pr), Vol 81)", Humana Press, 1998;
30 Maramorosh and Koprowski, eds. "Methods in Virology" 7 vols, Academic Press, New York 1967-1984; Hill, S.A. "Methods in Plant Virology", Blackwell, Oxford, 1984; Walkey, D.G.A. "Applied Plant Virology", Wiley, New York, 1985; and Kado

and Agrawa, eds. "Principles and Techniques in Plant Virology", Van Nostrand-Reinhold, New York.

In addition to the above, the polynucleotide of the present invention can also be introduced into a chloroplast genome thereby enabling chloroplast expression.

5 A technique for introducing exogenous polynucleotide sequences to the genome of the chloroplasts is known. This technique involves the following procedures. First, plant cells are chemically treated so as to reduce the number of chloroplasts per cell to about one. Then, the exogenous polynucleotide is introduced via particle bombardment into the cells with the aim of introducing at least one
10 exogenous polynucleotide molecule into the chloroplasts. The exogenous polynucleotides selected such that it is integratable into the chloroplast's genome via homologous recombination which is readily effected by enzymes inherent to the chloroplast. To this end, the exogenous polynucleotide includes, in addition to a gene of interest, at least one polynucleotide stretch which is derived from the chloroplast's
15 genome. In addition, the exogenous polynucleotide includes a selectable marker, which serves by sequential selection procedures to ascertain that all or substantially all of the copies of the chloroplast genomes following such selection will include the exogenous polynucleotide. Further details relating to this technique are found in U.S. Pat. Nos. 4,945,050; and 5,693,507 which are incorporated herein by reference. A
20 polypeptide can thus be produced by the protein expression system of the chloroplast and become integrated into the chloroplast's inner membrane.

Photosynthetic organisms generated according to the teachings described hereinabove may be rendered more thermoplastic than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence of the present
25 invention.

Alternatively or additionally, the photosynthetic organism of the present invention may be capable of expressing more Rubisco than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence of the present invention.

30 Yet alternatively or additionally, the photosynthetic organism of the present invention may be capable of expressing more psaC than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence of the present invention.

Still alternatively or additionally, the photosynthetic organism of the present invention may grow faster than a photosynthetic organism not comprising the exogenous nucleic acid sequence of the present invention.

5 Still alternatively or additionally, the photosynthetic organism of the present invention may be accumulating more chlorophyll than an identical photosynthetic organism not comprising the exogenous nucleic acid sequence.

Thus, the present teachings may be used to increase activity of a photosystem II reaction center of a photosynthetic organism under non-physiological temperature.

10 As used herein the phrase "non-physiological temperature" refers to a temperature which does not mediate ultimate growth of the non-genetically modified photosynthetic organism (e.g., at least about 70 % reduction in photosynthetic activity). This may be achieved by introducing into the photosynthetic organism the polynucleotide of the present invention and growing it in the non-physiological temperature, thereby increasing the activity of the photosystem II reaction center
15 under a non-physiological temperature.

The present teachings allow for improving biomass/vigor/yield of a photosynthetic organism comprising the nucleic acid sequence of the present invention, preferably when grown under non-physiological temperature (abiotic stress condition xxxe.g., heat and/or alternating temperatures).

20 As used herein the phrase "biomass" refers to the amount or quantity of tissue (in particular cellulose comprising tissue) produced from the photosynthetic organism in a growing season, which could also determine or affect the photosynthetic organism yield or the yield per growing area.

25 As used herein the phrase "vigor" refers to the amount or quantity of tissue produced from the photosynthetic organism in a given time. Hence increase vigor could determine or affect the photosynthetic organism yield or the yield per growing time or growing area.

30 As used herein the phrase "yield" refers to the amount or quantity of tissue produced and harvested as the photosynthetic organism produced product. Hence increase yield could affect the economic benefit one can obtain from the photosynthetic organism in a certain growing time.

Methods of determining biomass, yield and vigor are well known in the art and further described in Coleman et al, 2006, Plant Biotechnology Journal 4 (1), 87–101.

As used herein the term "improving" or "increasing" refers to improving or
5 increasing the biomass/yield/vigor of the photosynthetic organism of the present invention by at least about 2 % more, 5 % more, 10 % more, 20 % more, 30 % more, 40 % more, 50 % more, 60 % more, 70 % more, 80 % more, 90 % or more than that of the non-transgenic photosynthetic organism (e.g., mock transfected, or naïve).

As it has recently been suggested that temperature is the most important
10 parameter for hydrogen production (Tsygankov 1999 Biotechnol. Bioeng. 64:709-15), the present invention also contemplates use of the photosynthetic organisms provided herein for hydrogen production.

As used herein the term "about" refers to $\pm 10\%$.

15

Additional objects, advantages, and novel features of the present invention will become apparent to one ordinarily skilled in the art upon examination of the following examples, which are not intended to be limiting. Additionally, each of the various embodiments and aspects of the present invention as delineated hereinabove and as
20 claimed in the claims section below finds experimental support in the following examples.

EXAMPLES

Reference is now made to the following examples, which together with the
25 above descriptions, illustrate the invention in a non limiting fashion.

Generally, the nomenclature used herein and the laboratory procedures utilized in the present invention include molecular, biochemical, microbiological and recombinant DNA techniques. Such techniques are thoroughly explained in the literature. See, for example, "Molecular Cloning: A laboratory Manual" Sambrook et al., (1989); "Current Protocols in Molecular Biology" Volumes I-III Ausubel, R. M., ed. (1994); Ausubel et al., "Current Protocols in Molecular Biology", John Wiley and Sons, Baltimore, Maryland (1989); Perbal, "A Practical Guide to Molecular Cloning", John Wiley & Sons, New York (1988); Watson et al., "Recombinant DNA", Scientific

American Books, New York; Birren et al. (eds) "Genome Analysis: A Laboratory Manual Series", Vols. 1-4, Cold Spring Harbor Laboratory Press, New York (1998); methodologies as set forth in U.S. Pat. Nos. 4,666,828; 4,683,202; 4,801,531; 5,192,659 and 5,272,057; "Cell Biology: A Laboratory Handbook", Volumes I-III
5 Cellis, J. E., ed. (1994); "Current Protocols in Immunology" Volumes I-III Coligan J. E., ed. (1994); Stites et al. (eds), "Basic and Clinical Immunology" (8th Edition), Appleton & Lange, Norwalk, CT (1994); Mishell and Shiigi (eds), "Selected Methods in Cellular Immunology", W. H. Freeman and Co., New York (1980); available immunoassays are extensively described in the patent and scientific literature, see, for
10 example, U.S. Pat. Nos. 3,791,932; 3,839,153; 3,850,752; 3,850,578; 3,853,987; 3,867,517; 3,879,262; 3,901,654; 3,935,074; 3,984,533; 3,996,345; 4,034,074; 4,098,876; 4,879,219; 5,011,771 and 5,281,521; "Oligonucleotide Synthesis" Gait, M. J., ed. (1984); "Nucleic Acid Hybridization" Hames, B. D., and Higgins S. J., eds. (1985); "Transcription and Translation" Hames, B. D., and Higgins S. J., Eds. (1984);
15 "Animal Cell Culture" Freshney, R. I., ed. (1986); "Immobilized Cells and Enzymes" IRL Press, (1986); "A Practical Guide to Molecular Cloning" Perbal, B., (1984) and "Methods in Enzymology" Vol. 1-317, Academic Press; "PCR Protocols: A Guide To Methods And Applications", Academic Press, San Diego, CA (1990); Marshak et al., "Strategies for Protein Purification and Characterization - A Laboratory Course
20 Manual" CSHL Press (1996); all of which are incorporated by reference as if fully set forth herein. Other general references are provided throughout this document. The procedures therein are believed to be well known in the art and are provided for the convenience of the reader. All the information contained therein is incorporated herein by reference.

25

GENERAL MATERIALS AND EXPERIMENTAL PROCEDURES

PCR-based mutagenesis of the *Synechocystis* 6803

The *psbAII* gene mutation at positions 634-636, corresponding to D1-Ser212, was inserted using combinatorial DopA/DopB primers (Table 1 hereinbelow) coding
30 for 16 residues at this site. The oligonucleotides P5, P3, P209 and P212 were designed to match known sequences in the *psbAII* gene (Table 1). The *NsiI* restriction site modification was introduced in the third codon downstream to the 212 loci; facilitating screening of the transformant colonies. The final PCR product was used

to transform the recipient strain ΔSRS268-270 in the *psbAII* gene as previously described [Kless and Vermaas, *Biochemistry* 35, 16458-16464 (1996)]. Specific X212 oligonucleotides were designed to introduce codons in the D1-212 site corresponding to (Gly, Lys, His, Phe, Trp, Tyr, Arg, Met, Asp, Gln, and Glu) amino acids that were not obtained or introduced by the combinatorial mutagenesis approach. Additionally S209/212 nucleotide was prepared to insert degenerate sequence at 625-627, corresponding to D1-Ser209 and at 634-636 corresponding to D1-Ser212 sites to obtain double D1-209/212 mutants.

10

Table 1
Oligonucleotides used in mutagenesis

Oligonucleotides ^a	Nucleotide Sequences ^b	Gene Position ^c
DopA ^d SEQ ID NO: 1	CGGTGGTAGCTTGTTTCYDBGCCATGCA/TGGTTCC	618-651
DopB ^d SEQ ID NO: 2	CGGTGGTAGCTTGTTTCVHWGCCATGCA/TGGTTCC	618-651
P212 SEQ ID NO: 3	GAACAAGCTACCACCGAATACACCAGC	633-607
P3 SEQ ID NO: 4	GGATTAATTCTCTAGACTCTCTAATGG1	233-1179
P5 SEQ ID NO: 5	CCAAAACGCCCTCTGTTTACC	(-187)-(-166)
P260 SEQ ID NO: 6	GCCGTGGGCGGCAACGATGTTGTAGG	759-734
P4 SEQ ID NO: 7	CTTCCACATGTTAGGTGT	588-605
P209.....	ACCACCGAATACACCAGCCACACCTAA.....	624-598
SEQ ID NO: 8		
X212 ^e SEQ ID NO: 9	CGGTGGTAGCTTGTTTCxxxGCCATGCA/TGGTTCC	618-651
S209/212.....	GGTGTATTCGGTGGTNNSTTGTTCCNNSGCCATGCA/TGGTTCC.....	610-651
SEQ ID NO: 10		

^aThe P260 and P4 oligonucleotides were used for sequencing the PCR product. ^bOligonucleotides are listed in a 5' to 3' direction. Restriction sites of *NsiI* introduced as silent mutations are underlined, and the modifications are in italics. ^c*psbAII* numbering as described. ^dThe universal code for degenerate oligonucleotides is Y(C, T); D(g, A, T); B(g, T, C); V(g, A, C); H(A, T, C); W(A, T); N(A, T, C, g); S(C, g). ^eThe xxx represents specific codons for Trp (TGG), Tyr (TAC), Phe (TTT), Arg (CGG), Gln (CAA), Glu (GAA), His (CAC), Asp (GAT), Lys (AAA), Gly (GGT) and Met (ATG).

***Synechocystis* sp. PCC6803 growth conditions**

Synechocystis sp. PCC 6803 cells were grown in BG-11 mineral medium as previously described [Williams, *Methods in Enzymology* 167, 766-778 (1988)]. All wild type and mutant strains were maintained on 1.5 % agar (Difco) plates in the

20

presence of 5 µg/ml kanamycin (Monosulfate, Sigma). Five mM glucose was routinely added to plates to maintain PSII-independent growth. To get liquid cultures suitable for measurements, cells were scratched from the plates and transferred into a stirred 50ml BG-11 liquid medium in the absence of glucose for one day growth in order to get fresh starter culture. Liquid cultures were grown on gyratory shaker (200 rpm) at 30 °C and 40 µmol (photons) m⁻²s⁻¹ of white light. The growth chamber was aerated with fresh air. Cells for the photosynthetic measurements were transferred into stirred 250 ml BG-11 medium at the same conditions and harvested at the end of the logarithmic growth phase corresponding to a chlorophyll concentration of 4-7 µg/ml.

Temperature treatments

Cells grown for 3 days were diluted or concentrated to an OD₇₃₀ = 0.5 with fresh medium prior to treatments. Cultures of the similar cell density were transferred to 30 °C or 43 °C (same light conditions) for 7 days. For the plasticity test, cultures having OD₇₃₀ = 0.5 were incubated at a 43 °C - 30 °C- 43 °C, 10 °C- 43 °C- 10 °C or 43 °C- 10 °C- 43 °C cycle for 48 hours at each temperature.

Pigment analysis

Pellets of the cells grown in liquid cultures were extracted in 90 % methanol. The chlorophyll concentration was determined spectroscopically with Jasco V530 spectrophotometer (Jasco, USA) using the molar extinction coefficient determined by Lichtenthaler [Wellburn and Lichtenthaler, *Advances In Photosynthesis Research* Vol. II (1984)]. The carotenoid content was analyzed with Smartline HPLC system (Knauer, Germany) using Jupiter HPLC column (15 µm particle size, C18, 300 Å pore size, 250 x 15 mm, Phenomenex, USA) according to the procedure previously described [Pocock et al., *Methods Mol Biol* 274, 137-48 (2004)].

Growth rate measurements

The growth rates were measured in terms of optical density at 730 nm (OD₇₃₀) in aliquots obtained from the liquid cultures every 24 hours. Cell density was estimated by turbidometry, measuring light attenuation at 730 nm after diluting

cultures to OD values less than 0.5, where turbidity is roughly proportional to cell concentration.

Isolation of thylakoid proteins and Western blot analysis

5 Thylakoid membranes were isolated according to the method previously described by Callahan [Callahan et al., *J Biol Chem* 265, 15357-60 (1990)]. Thylakoid proteins were solubilized in sample buffer [0.5 M Tris-HCl pH 6.8, 1 % SDS, 24 % glycerol, 4 % β -mercaptoethanol, 0.001 % (w/v) Bromophenol blue], incubated at room temperature for 1 hour and then separated by SDS-PAGE 12.5 %. The
10 equivalent of 2.5 μ g of chlorophyll was loaded into each well. Proteins were electroblotted to PVDF (Hybond - P, Amersham) using a Bio-Rad Mini Transblot Cell (Bio-Rad, Hercules, CA, USA). The immunodetection was carried out using a chemiluminescence kit (Pierce). Antibodies against D1, psaC and RbcL proteins were purchased from Agrisera (Umeå, Sweden).

15

Measurement of the temperature dependence of the $Q_A^-Q_B$ electron transfer rate

The temperature dependence for the $Q_A^-Q_B$ electron transfer rate was assessed by measurements the chlorophyll Chl *a* fluorescence as recently described [Shlyk-
20 Kerner et al., *Nature*, 442(7104): 827-30 (2006)]. Three days old liquid cultures were diluted by BG-11 medium to 5 μ M of chlorophyll and dark adapted for at least 30 minutes on ice prior to fluorescence measurements. Aliquots of 3 ml cell suspension were then placed into polystyrene cuvettes (Sigma) and adapted to a selected temperature for 5 minutes in the dark. The measurements of Chl *a* fluorescence were
25 performed with FL-3000 double-modulation fluorometer equipped with TR 2000 thermoregulator (P.S. Instruments Inc., Czech Republic). The re-oxidation kinetics of Q_A^- was measured as the decay of Chl *a* fluorescence following the single-turnover saturating flash (25 μ s duration, $\lambda = 630$ nm) that reduced all Q_A in the sample into Q_A^- . The variable fluorescence decay, reflecting the re-oxidation of Q_A^- was measured
30 on the logarithmic time scale to minimize the perturbation of the sample. The normalized decay kinetic curves were fitted by least-squares numerical analysis to a three-exponential function. The highest rate constant was assigned to the electron transfer from Q_A^- to Q_B , as described by Nedbal *et al.* [Nedbal et al., *Journal of*

Photochemistry and Photobiology B: Biology 48, 154-157 (1999)]. Data from at least ten individual experiments were averaged. Importantly, the fluorescence analysis as used allowed for discrimination between active and inactive PSII reaction centres [Joshi and Fragata, *Z Naturforsch [C]* 54, 35-43 (1999)]. The inactive centres
5 contribute to the longest lifetime component and become apparent at temperatures well above the physiological range [Joshi and Fragata, *supra*].

In silico analysis

Intra-molecular cavity analysis (Protein cavities): Cavity analysis was
10 conducted with VOLBL [Liang et al., *Proteins* 33, 18-29 (1998); Liang et al., *Proteins* 33, 1-17 (1998)] using the standard 1 Å-radius probe used for cavities. Proteins were analyzed without hydrogens. The addition of hydrogens caused a reduction of up to 30 % in cavity size (data not shown). Notably, the binding niche of the cofactors was surrounded by a number of cavities. As the study was confined to
15 the area within the four-helix-bundle transmembrane core of the protein, only cavities that were lined solely by atoms from the D and E helices were represented.

Evolutionary conservation of the GxxxG-like motif in RC subunits of thermophilic, thermotolerant and psychrophilic organisms: The conservation level of photosystem II RC was assessed by (a) searching the BLOCKS [Henikoff et
20 al., *Nucleic Acids Res* 28, 228-30 (2000)] database with the D1 sequence of *Synechocystis* 6803; and (b) searching sequence space with PSI-BLAST [Altschul et al., *Nucleic Acids Res* 25, 3389-402 (1997)], using both the original D1 sequence and the representative sequence (Cobbler sequence) from the Blocks database [Henikoff et al., *supra*] – block IPB000484B. In addition, newly sequenced relevant organisms
25 were searched via the Entrez Genome Project, and a Blast search (using TBLASTN2.1.2) was conducted against cyanobacteria and plant sequences that have not been fully annotated into DNA databases and in designated databases such as CyanoBase [Nakamura et al., *Nucleic Acids Research* 26, 63-67 (1998)].

Hydrogen assignment and hydrogen bond analysis: Hydrogens were added
30 to the structure using REDUCE [Word et al., *J Mol Biol* 285, 1711-33 (1999)], and intersubunit H-bonds were screened using an in-house program [Goldberg, M.Sc. thesis, Weizmann Institute of Science (2004)] following geometric criteria [Senes et al., *Proc Natl Acad Sci U S A* 98, 9056-61 (2001)] for putative bonds. The geometric

criteria included: $DH < 3.5\text{\AA}$ and $DH-A \text{ angle} > 120^\circ$ (or $DH-A \text{ angle} > 100^\circ$ if $DH < 3\text{\AA}$) where D, A, and DH were the donor atom, acceptor atom and hydrogen found on the donor atom. These constraints, particularly the allowed D/A pair distance and DH-A angle, were more relaxed than those given by Desiraju [Desiraju, *Acc Chem Res* 35, 565-73 (2002)]. The constraints were used to ensure that under the non-ideal resolution of the structures analyzed all potential H-bonds were captured. Several interactions were depicted in which the DH-A and/or angle thresholds were not satisfied. Alternatively, in some cases the H-bond flip between two positions but cannot occupy both simultaneously. These interactions were given to demonstrate the close intersubunit packing in this region.

Graphical representation of structural motifs: Structures were drawn with PyMol (Warren L. DeLano "The PyMOL Molecular Graphics System." DeLano Scientific LLC, San Carlos, CA, USA. (www dot pymol dot org). In order to replicate the result of the VOLBL [Liang et al., *Proteins* 33, 18-29 (1998); Liang et al., *Proteins* 33, 1-17 (1998)] cavity analysis program, atom radii were changed to the OPLS-derived van der Waals radii as applied in VOLBL, and found in the param dat file. In order to confine the surface representation to the cavities, pseudo atoms were inserted into the cavities. Next, the surface_carve_selection function was used to confine the surface represented to the area adjacent to these pseudo atoms. The surface was represented using a command such as "show surface, protein and (pseudo_atoms expand 3)". This resulted with surface representation of the cavity but also pieces of other cavities found around the cofactors' binding niche. In order to eliminate the latter from the drawing, the van der Waals radii of the cofactor atoms was set to a large enough number.

Non-conserved positions at D1-209 and D1-212 loci of additional, hydrogen generated cyanobacteria

1. **D1(Ser209)Ala** - (locus: YP_476691, gi: 86607929) from extremophilic cyanobacterium Yellowstone B-Prime known also as Synechococcus sp. strain JA-2-3B'a(2-13). The A-Prime strain has the same sequence in this region (locus: YP_474719, gi: 86605956).

2. **D1(Ser209)Ala** - (locus: ABA21206, gi: 17132876) - *Nostoc* (or *Anabena*) sp PCC 7120 - mesophilic, nitrogen fixing cyanobacterium.

3. **D1(Ser209)Ala** – (locus: BAB75441, gi: 75701530) – *Anabaena variabilis* ATCC 29413 – mesophilic, nitrogen fixing, hydrogen producing cyanobacterium.
4. **D1(Ser209)Ala** – (locus: EAQ69357, gi: 86168099) – *Synechococcus* sp. RS9917 – halotolerant cyanobacterium.
5. **D1(Ser209)Ala,(Ser212)Ala** - PSBA-2 (Swissprot: PSBA2_SYNEN or PSBA2_SYNEL), one of the two psbA genes from the thermophilic cyanobacterium *Synechococcus elongatus naegeli*.
6. **D1(Ser209)Ala,(Ser212)Cys** - PSBA-1 (Swissprot; PSB1_SYNVU), thermophilic cyanobacterium *Synechococcus vulcanus*.
7. **D1(Ser212)Cys** – PSBA-1, (TrEMBL; Q9S3W5) – true branching filamentous thermophilic cyanobacterium *Mastigocladus laminosus*.
8. **D1(Ser209)Ala,(Ser212)Ala** - PSBA-1 (Swissprot PSB1_SYNY3, *Synechocystis* sp. PCC 6803). Notably, this PSBA-1 is not expressed on a constitutive level but is expressed under high-light-stimulated transcription. Thus, it may represent an adaptation to stress conditions.
9. **D1(Ser209)Ala,(Leu210)Phe(Ser212)Cys** – (locus: ZP_00108645, gi: 23126757) *Nostoc punctiforme* sp. PCC 73102 – mesophilic, nitrogen fixing cyanobacterium.

20

Non-conserved positions at the D1-209 and D1-212 loci in microalgae

1. In the thermotolerant red alga *Cyanidium caldarium*, the **D1-Ser209** has been modified to **D1-Ala209** - PSBA-1 (Swissprot: PSBA_CYACA) (thrives at temperature higher than 45 °C, pH=1 and/or high salinity).
2. **D1(Ser209)Ala** - (locus: NP_848970, gi: 30468083) in thermoacidophilic unicellular red alga *Cyanidioschyzon merolae* strain 10D that is found in acidic hot springs (locus: AAM62037, gi: 21913561).
3. **D1(Ser209)Ala** - (locus: AAR30280, gi: 39753030) in thermoacidophilic unicellular red alga *Galdieria maxima* from acidic hot springs.
4. **D1(Ser209)Ala** – (locus: AAZ59080, gi: 72003278) – *Prochlorococcus marinus* str. NATL2A – dominates phytoplankton in tropical and subtropical oceans.

30

EXAMPLE 1

Protein flexibility acclimatizes photosynthetic energy conversion to the ambient temperature

Solar energy conversion in reaction centers (RC) from non-oxygenic purple bacteria and from photosystem II (PSII) of oxygenic organisms is initiated by means of a multi-step, light-induced electron transfer (ET) across the photosynthetic membrane (Figure 1A). The process ends in the reduction of a quinone molecule (Q_B) by another quinone (Q_A) accompanied by imperative reorganization and mobilization of protons. Former studies of this reaction showed that protein flexibility and a conformational change are required to set on the $Q_A \rightarrow Q_B$ ET, that the rate constant, k , is independent of the quinones' redox energies, and that the reaction should be accelerated at elevated temperatures [Okamura et al., *Biochim Biophys Acta* 1458, 148-63 (2000); Li et al., *Biochemistry* 37, 2818-29 (1998); Xu et al., *Biochemistry* 41, 2694-701 (2002)]. Following the classical Arrhenius paradigm, k is expected to increase exponentially with $1/T$ at a pace equal to $\Delta H^\ddagger/R$ where ΔH^\ddagger is the enthalpy of ET activation. However, inventors observed that this prediction was violated in intact cells of mesophilic and thermophilic cyanobacteria (Figure 2A). Here, k levels off at T_o slightly below the growth temperature of the examined organisms ($T_o = 26$ °C and $T_o = 59$ °C, for mesophilic and thermophilic strains, respectively), the rate became temperature independent throughout the entire physiological range and then slightly decreased. In all strains, k reached a similar maximum value of ~ 3000 - 3500 sec^{-1} . In the thermo-tolerant organism *Cyanidium caldarium* it reached the same value but at $T_o = 43$ °C (data not shown). T_o was significantly lower from the reported denaturation temperature of the RC (higher than 40 °C in mesophiles [Joshi et al., *Zeitschrift Fur Naturforschung C-a Journal of Biosciences* 54, 35-43 (1999)]) ruling it out as the origin for the observed phenomenon (Figure 6, further explained in detail hereinbelow).

Thus, the photosynthetic organism appeared to utilize novel physico-chemical principles to slow down the rate acceleration at elevated temperature. Remarkably, the activation enthalpies (ΔH^\ddagger) below T_o for the examined mesophile and thermophile were almost identical (Figure 2A). However, the activation entropies (ΔS^\ddagger) for the thermophilic *Thermosynechococcus elongatus* was more negative by ~ 1 kcal.mol^{-1}

suggesting that flexibility of the RC excited state was a key player in the temperature adjustment of the energy conversion rate.

In search for motifs that account for the observed differences in the activation entropies, inventors examined the sequences of the two major protein subunits by sequence alignment of the *D* helices in RC found in all photosystem II RCs and in purple bacteria (Figures 1A-C).

Rather than searching in proximity to the cofactor-binding sites, inventors confined the search to the four transmembrane-helices that physically hold the electron transfer cofactors. Two sites, D1-209 and D1-212, were found to undergo consistent changes between mesophilic, thermotolerant and thermophilic organisms including cyanobacteria, algae and green plants. These sites, positioned in a GxxxG-like sequence motif (gray background, Figure 1C), where 'G' denotes small residues such as Gly, Ala, Ser, Cys and Thr, were found at the closest contact of the two major protein subunits (D1 and D2, Figure 1C). This motif and the structurally homologous motif in purple bacteria was found at the center of the transmembrane (TM) domain, where the D and E helices of the two major RC subunits of photosystem II (D1 and D2) and bacterial RC (L and M) maintain a conserved intersubunit hydrogen bonding (ISHB) network (Figures 1A-B, 5A-E and Table 2). ISHB was suggested to provide local flexibility while maintaining overall stability.

20

Table 2

Potential ISHBs Involving the Transmembrane Regions of the D and E in the RC of Purple Bacteria and Photosystem II

Organism	PDB/Ref	Res. (Å)	HB Donor	HB Acceptor	D-A	DH-A	Angle
Bacterial RC <i>R. sphaeroides</i>	1m3x ⁸⁷	2.55	M-Ala213CA	L-Ser237OG	3.7	2.9	131
			L-Asn183ND2	M-Ser212O	3.3	2.3	152
			M-Gly220CA	L-His230O	3.5	2.9	117
	1pcr ⁸⁸	2.65	L-Asn183ND2	M-Ser212O	3.3	2.4	148
			M-Ala213CA	L-Asn183OD1	4.3	3.2	176
				L-Ser237OG	4.0	3.1	132
	1aij ⁸⁹	2.2	L-Asn183ND2	M-Ser212O	3.1	2.22	148
			M-Ala213CA	L-Asn183OD1	4.4	3.3	165
			M-Ala217CA	L-Ser237OG	3.4	2.9	111
			M-Gly220CA	L-His230O	3.6	3.0	119
	1aig ⁸⁹	2.6	L-Ser237OG	M-Ala213O	2.8	1.85	165
			M-Ala213CA	L-Asn183OD1	3.1	2.2	140
L-Ser237OG				3.4	2.6	126	
<i>B. viridis</i>	1prc ⁶⁶	2.3	M-Ala217N	L-Ser237OG	3.3	2.8	107
			M-Ser271OG	L-Ala184O	2.9	2.03	144

		41						
			L-Asn183ND2	M-Cys210O	3.4	2.57	146	
			L-Ala184CA	M-Cys210SG	4.2	3.3	140	
			L-Gly188CA	M-Asn147OD1	3.7	3.0	120	
				M-Ser271OG	3.4	2.8	114	
	<i>T. tepidum</i>	leys ⁹⁰	2.2	L-Asn191ND2	2.8	1.9	145	
				M-Ala212CA	4.5	3.4	177	
				M-Ala272CA	4.1	3.3	126	
Photosystem II RC	<i>T. elongatus</i>	Is51 ⁷	3.5	D1-Gly208CA	D2-Cys211SG	4.3	3.3	149
				D1-Cys212SG	D2-Gly207O	3.4 ^a		
				D1-Cys212CA	D2-Cys211SG	4.2 ^a		
				D1-Ala213CA	D2-Met271O	3.8	2.8	143
				D1-Gly216CA	D2-His268O	3.3	2.5	130
				D1-Leu275CA	D2-Cys211SG	3.8 ^a		
				D1-Ala276CA	D2-Cys211O	3.6	2.9	123
				D2-Cys211SG	D1-Gly208O	4.0 ^a		
				D2-Cys211N	D1-Leu275O	3.5 ^a		
				D2-Ala212CA	D1-Cys212SG	3.6 ^a		
	D2-Gly215CA	D1-Leu275O	4.2	3.5	130			
		D1-His272O	3.2	2.7	110			
	<i>T. elongatus</i>	2axt ⁸	3.0	D1-Ala209CA	D2-Gly207O	4.0	3.0	152
				D1-Gly208CA	D2-Cys211SG	3.9	3.2	127
				D1-Ala276CA	D2-Ala212O	4.1	3.4	124
				D2-Ala212CA	D1-Leu275O	3.9	2.9	147
				D1-Cys212SG	D2-Met271O	3.2 [~]	1.9	178
				D1-Ala213CA	D2-Gly207O	3.2 ^a		
				D2-Gly215CA	D2-Met271O	3.6	2.7	138
				D1-Gly216CA	D1-His272O	3.3	2.9	101
D2-Cys211SG				D2-His268O	3.1	2.7	99	
D2-Met271CA				D1-Gly208O	3.1 ^a			
	D1-Leu275O	4.0	2.8	156				
	D1-Cys212SG	3.8 ^a	3.2	113				

Thus, changes in the amino acids within the identified GxxxG-like motif may result in modification of the RC localized flexibility and consequently, in the entropy of activation. Indeed, Ser occupies the D1-212 and D1-209 in mesophilic organisms including the cyanobacterium *Synechocystis* 6803. The D1-209 is modified to Ala in the thermotolerant red alga *Cyanidium caldarium*, and in thermophilic cyanobacteria, Cys or Ala populates the D1-212 while Ala occupies the D1-209 site (Figure 1C).

Two sizable cavities augmented by several small ones complement the ISHB cluster (Figures 1A-B, 5A-E and Table 3). The largest cavity (47 Å³) lay at the interface of the two subunits and extends from the D1-212 site toward the histidines that ligate the non-heme iron. A second cavity (32 Å³) occupies the other side of the D1-212 residue and is lined by two flexible side-chains that are in common to the largest cavity. Consequently, movements or changes in the size of the D1-212 residue may alter the volume and shape of either cavity or merge the two cavities into one. In the structurally homologous domain in RC from purple bacteria, a large cavity

encompassing over 100 Å³ was found in this region. As previously demonstrated for bacteriorhodopsin [Friedman et al., *Biophys J* 85, 886-96 (2003)], such cavities in the inner core of a protein may facilitate the conformational rearrangement required for enhancing the protein's local flexibility during discrete functional steps.

5

Table 3**Cavities found between the D and E helices of bacterial RC and PSII RC**

PDB Code / 212 residue	Molecular Surface (Å³)	Atoms lining the cavity
Wild type – photosystem II (D1-212 = Cys)		
2axt	46.8	D1: Phe-211-O, Cys-212-CA, His-215-CB, Leu-275-CB/CD1/CD2 D2: Leu-210-O, Cys-211-CA/C/CB/SG, His-214-CB/CD2, Gly-215-N, Met-271-CE
	31.8	D1: G-208-CA, F-211-CB/CD1, F-274-O/CD2, L-275-CA/O/CD2, W-278-CB/CD1, P-279-CD D2: C-211-SG
	7.9	D1: C-212-SG D2: G-207-CA, L-210-CD1, M-274-CB/CG1, P-275-CD
	6.2	D1: C-212-SG D2: L-210-CD1, M-271-CA, V-274-CB, P-275-CD
	6.3	D1: H-272-CD2, L-275-CD1 D2: H-214-CD2, G-215-N/CA
Photosystem II RC – Rotamer-library based D1-212 mutation		
all residues	40.2	D1: F-211-C/O, RESIDUE-212-CA, H-215-CB, L-275-CB/CD1/CD2 D2: C-211-CA/C/CB/SG, H-214-CB/CD2, G-215-N, M-271-CE * In the Gln and Glu mutants the OE1 atom is involved causing a decrease of 0.1, and 0.2 Å ³ in cavity volume, respectively. In Arg and Trp the cavity is 39.7 Å ³ due to similar reasons.
	28.8	D1: G-208-CA, F-211-CD1, F-274-O/CD2, L-275-CA/O/CD2, W-278-CB/CD1, P-279-CD D2: C-211-SG
Gly	48.7	D1: A-209-CA/O, G-212-CA/C D2: G-207-CA/C/O, L-210-CB/CD1, C-211-N, M-271-CA/O/CB/CG/CE, P-275-CB/CG/CD
Ala/Cys / Ser	21.9	D1: A-212-CB D2: G-207-CA, L-210-CB/CD1, M-271-CA/O/CB/CG/CE, P-275-CD * In Cys / Ser the atoms are the same but cavity is 0.1 Å ³ smaller
Bacterial Reaction Center (Largest cavity only)		
1ajj ¹⁶	108	L subunit: Leu187-O/CD1, His190-CB/CD2, Gly191-N/CA, M subunit: Leu215-O/CG/CD1/CD2, Phe216-CA/CD1/CE1/CZ, His219-CB/CG/CD2/NE2, Ile265-C/O/CG1/CG2/CD1, His266-CA/CD2, Ala269CB, U108-C1M
2prc ⁶⁶	135	L subunit: Leu187-O/CD1/CD2, His190-CB/CD2, Gly191-N/CA, M subunit: Leu213-C/O/CB/CD1/CD2, Phe214-CA/CD1/CE1/CZ, His217-CB/CG/ND1/CD2/CD1/NE2, Val263-C/O/CG1/CG2, His264-CA/O/CG/CD2, Gly26-7N/CA/C, Trp268-N, 7MQ501-C3M

Having two adjacent structural motifs associated with flexibility, inventors tentatively concluded that replacement of Ser by Cys or Ala within the GxxxG-like motif accounts for the observed changes in the localized protein flexibility and subsequently the temperature adjustment of the RC activity. *In-silico* rotamer-library based saturated mutagenesis was performed on the D1-212 site in the available structures of thermophilic cyanobacteria conservatively allowing only the D1-212 to re-pack. Under such constraints, the volume of the larger cavities was reduced by about 10-20 % and the small ones merged into one. This new cavity was evident only in the mutants that had small residues (Gly, Ala, Cys, Ser; Table 3). In all other mutants the cavity was blocked by the D1-212 side-chain (Figures 3A-B and Table 3). Dissociation of ISHB between residues of the GXXXG-like motif should increase the localized flexibility provided that side chains are free to undergo reorganization. However, this freedom of motion was decreased when the packing values of the involved residues are decreased, shifting T_o to higher values. Thus, Ala and Cys (packing value = 0.471 and 0.463, respectively) were expected to correspond to higher T_o values compared to Ser (packing value = 0.484). Unlike the non-bulky residues (termed 'class I'), the bulky ones (termed 'class II') partly blocked the intersubunit cavity (Figure 3A-B and Table 3). The latter had the lowest packing values and were expected to elevate T_o to a non-physiological range where PSII RC denaturation takes over.

Inventors directly challenged the *in-silico* predictions by introducing mutations in photosystem II RC of *Synechocystis* 6803. The D1-S212 residue was replaced by each of the remaining 19 amino acids and the effect on the photosynthetic parameters in intact cells was investigated. Thirteen photoautotrophic mutant strains were isolated (see Materials section). Mutants with aromatic and positively charged residues, all assigned to class II, failed to grow photoautotrophically possibly due to impaired RC folding. Mutants containing class I residues (Gly, Cys, Ala, Thr, Asn, and Asp) presented similar patterns of $Q_A^- \rightarrow Q_B$ ET temperature dependence (Figure 2B). Yet, T_o was elevated by more than 6 °C and more than 7 °C in the D1-S212C and D1-S212A mutants, respectively. The rate constant reached similar maximum values (approximately $3-4 \times 10^3 \text{ sec}^{-1}$) in the different class I mutants at their T_o and above (Figure 2B). Obtained photoautotrophic mutants with class II residues (Pro, Val, Ile, Leu, Glu, and Gln) presented significantly lower values for the ET rate

constant and their k values increased in a monotonic, monophasic fashion with increasing temperature through most of the physiological range (Figure 4A). ΔH^\ddagger was similar in all class I residues but ΔS^\ddagger at room temperature was more negative by 0.15 and 0.3 kcal.mol⁻¹ for the Cys and Ala-substituted strains (Figures 2B and 4B) depicting the significance of D1-212 in modulating the RC localized flexibility and T_o . For class II mutants, ΔH^\ddagger was usually similar to the wild type and class I mutants (below T_o) but ΔS^\ddagger was more negative by up to ~ 1.8 kcal.mol⁻¹ at room temperature (Figure 4B) thus indicating decreased local flexibility of the transition state involved in the $Q_A^- \rightarrow Q_B$ ET in class II mutants.

10 Mutagenesis of the D1-S209 residue resulted in only eight photoautotrophic mutants: six of class I (Gly, Ala, Thr, Asn, and Asp), and only two from class II (Val and Pro), possibly because substitution with residues of very low packing values impaired the PSII RC functional assembly. All photoautotrophic mutants exhibited levelling of the $Q_A^- \rightarrow Q_B$ ET rate at elevated temperatures (data not shown). The T_o temperature of the D1-S209A mutant elevated by more than 7 °C while maintaining a similar maximal rate constant as the wild type. This finding supports the suggested significance of the D1-209 loci in modifying the local flexibility of PSII RC and thereby the value of T_o .

Collectively, the identified structural motifs (ISHBs in a GxxxG-like sequence motif and adjacent cavities) in purple bacteria and photosystem II RCs, appeared to function as key modulators in controlling the activation entropy of the $Q_A^- \rightarrow Q_B$ ET rate and subsequently the value of T_o .

Non Arrhenius behaviour of enzymatic rate constants has been frequently considered to reflect inactivation and denaturation of the examined enzymes. However, T_o seems independent of the PSII RC denaturation which was introduced at significantly higher temperatures (Figure 6). In search for an alternative explanation for the temperature dependence of k , inventors looked at the non-Arrhenius behaviour in hydrogen transfer enzymes. The thermostability of the PSII in the wild type and the mutants was deduced from the amplitude of the fastest fluorescence decay component that is directly proportional to the concentration of photosystem II RC active in Q_A^- to Q_B ET. As shown in Figure 6, no significant change in the amplitude of the wild type fluorescence (D1- S212) was observed until approximately 37 °C and $T_{1/2}$ (50 % inactivation) was achieved at approximately 45 °C. Similar curves with

practically the same $T_{1/2}$ were obtained for Ala and Cys. Thus, T_o (25 °C, Figures 2A, 4A) appear independent of the thermostability for the wild type and class I mutants of *Synechocystis* 6803. Class II mutants may show similar temperature dependence to class I mutants (Val) or much lower thermostability (Gln) with well defined Arrhenius behaviour throughout the entire range of measurements (Figure 4B). Collectively, these findings exclude the involvement of photosystem II RC inactivation or denaturation in the non-Arrhenius behaviour of mesophiles, thermotolerant, thermophiles and class I mutants described.

In conclusion, inventors have shown that changes in locally flexible domains in membrane proteins provided the means for enzyme adaptation to the ambient temperature. Furthermore, inventors described and validated protein motifs that modulate this adaptation independently of denaturation processes.

EXAMPLE 2

Temperature dependence of the electron transfer rate in the different single and double mutants of the *Synechocystis* pp. 6803

The observed pattern of temperature dependence for the viable mutants (Table 4, below) of the *Synechocystis* mutants was the same as previously described for mutation with class I amino acid residues. Namely, the increase of electron transfer rate with temperature until an optimal temperature termed T_o was reached (where the rate constant reached a plateau and leveled off, Figure 7). In the D1-S212C and D1-SS209/212AC mutants, T_o shifted to higher temperature (approximately 32-33 °C) compared to the one observed for the wild type (approximately 25 °C). The D1-S212P presented the same T_o (approximately 25 °C) as a wild type, and in the D1-SS209/212AA, T_o was down shifted by 2.5 °C (Figure 7). The maximum electron transfer rate for the D1-S212C was the same as the one measured for the wild type wild type ($\sim 3-4 \times 10^3 \text{ s}^{-1}$). All other mutants presented significantly lower rate constants (Figure 7).

Table 4

Viable and non-viable mutations of D1-212 and 209 sites

Viable and non-viable mutations at D1-212 site																		
C	E	T	I	P	N	V	A	L	Q	M	D	G	F	W	Y	R	K	H
4	1	2	5	3	1	4	4	10	4	3	10	15	0	0	0	0	0	0
Viable and non-viable mutations at D1-212 site																		
C	G	T	P	N	V	A	D	E	I	L	Q	M	F	W	Y	R	K	H
2	4	2	10	11	1	3	8	0	0	0	0	0	0	0	0	0	0	0

Table 4: describes statistical data for amino acid residues obtained in the degenerated mutagenesis. Optional amino acids that could be obtained by the combinatorial mutagenesis approach / amount of mutants obtained for each AA residue. Each transformation yielded around 300 colonies under selection conditions. Randomly selected colonies (Table 4) were picked up for each mutation and their genomic DNA was amplified by PCR with P3 and P5 oligonucleotides (Table 1). The type of mutation was checked by restriction pattern with NsiI and by directed sequencing of that PCR product. For D1-Ser212, a total of 13 functional mutant strains and 6 non-functional mutants were obtained. For D1-Ser209 a total of 9 functional mutant strains and 10 non-functional mutants were obtained.

EXAMPLE 3

Thermotolerance of the mutants

To determine the effect of mutation on high temperature tolerance, cultures of *Synechocystis* 6803 wild type (Δ KS), the single D1-S212C, D1-S212P and double D1-SS209/212AA, D1-SS209/212AC mutants were grown under standard (30 °C) and under high (43 °C) temperatures for six days. The phenotypes of both the mutants and Δ KS did not show a significant difference at 30 °C (Figure 8A). However, bleaching of the wild type, D1-S212P and D1-SS209/212AA mutants was observed after 6 days of incubation at 43 °C (Figure 8B) suggesting growth arrest, cell death and chlorophyll (Chl) degradation. This observation was reflected in the chlorophyll accumulations rates (Figures 8E-F). At 30 °C, Δ KS and all mutants showed similar chlorophyll accumulation rates. At the high temperature limit (43 °C) and after 6 days of growth, D1-S212C and D1-SS209/212AC mutants presented several fold higher chlorophyll content than wild type and other mutants.

EXAMPLE 4**Growth rate of the wild type and the mutants at elevated temperatures**

The growth rate of the *Synechocystis* sp. PCC6803 wild type and mutant strains provided a direct estimate of the biomass production under different
5 constrains. Figure 8C depicts the time dependent turbidity of cell cultures grown at the physiological temperature (30 °C). All strains showed a significant growth rate where the biomass appears to increase by a factor of 14-16 after 6 days of growth. Figure 8D shows that at 43 °C the wild type and the D1-SS209/212AA, stop growing and consequently die after 2 days. However, the biomass of the D1-S212C and D1-
10 S212P increased at a rate of 6-6.5 fold after 6 days. Importantly, the growth arrest of the wild type and the D1-SS209/212AA are in line with the rapid decline of their chlorophyll content and the depletion of Rubisco and D1 proteins (Figures 8F, 8H and 8J).

15

EXAMPLE 5**Stability of D1 and Rubisco is maintained in D1-S212C and D1-SS209/212AC**

D1 and Rubisco are two essential proteins for photosynthesis and biomass formation, respectively [Haldimann and Feller, *Plant Cell and Environment* 28, 302-
20 317 (2005); Kouril et al., *Photosynthesis Research* 81, 49-66 (2004)]. Grown under standard temperatures (30 °C) their expression was the same for both wild type and mutants (Figures 8G and 8I). However, expression of these proteins was arrested upon incubation at 43 °C for 6 days for wild type (Δ KS), D1-SS209/212AA and D1-S212P mutants (Figures 8H and 8J). Yet, D1-S212C and D1-SS209/212AC mutants
25 present normal synthesis and accumulation of these proteins at both temperatures (Figures 8G-J). While the former is essential for light reactions, the latter catalyses the carbon dioxide uptake and fixation to form the carbohydrates. This example proves the viability of maintaining high biomass production in thermally adapted cyanobacteria after some mutations. Notably, Rubisco is known to undergo
30 inactivation at increased temperatures [Berry and Bjorkman, *Ann. Rev. Plant Physiol.* 31, 491-543 (1980)] underscoring the major inhibition of large scale biomass production at elevated temperatures in mesophiles. The psaC protein, which is part of

PSI protein complex, while present at 30 °C (Figure 8K) was depleted in the wild type and D1-SS209/212AA mutant at 43 °C (Figures 8L).

EXAMPLE 6

Thermoplasticity tests

5 The thermoplasticity of wild type and mutants was examined by subjecting the strains to high/low temperature cycles. First, cell cultures were acclimated to 43 °C - 30 °C - 43 °C cycle for 48 hours at each temperature (Figures 9A, 9C and 9E). Similarly all the cultures were acclimated at high temperature and then abruptly
10 shifted to low temperature (43 °C - 10 °C - 43 °C cycle) for 48 hours at each temperature (Figures 9B, 9D and 9F). As depicted in Figure 9A, only D1-S212C and D1-SS209/212AC mutants survived at the high temperature after 43°C- 30°C- 43°C cycle, while Δ KS and the other two mutant strains were completely bleached. The D1-S212C mutant showed the highest survival rate and the highest biomass
15 production as reflected in the growth rates and chlorophyll accumulation curves (Figures 9C and 9E, respectively). The D1-SS209/212AC mutant had comparable biomass production after periodic temperature changes (Figures 9C and 9E). All the other strains did not endure the treatment (Figures 9C and 9E). Acclimation at 43 °C - 10 °C - 43 °C cycle resulted in an impairment of mainly the wild type strain (Figures
20 9B, 9D and 9F) although the D1-S212P mutant also seems to be impaired. As shown in Figures 10A-C (10 °C - 43 °C -10 °C cycle) a very slow growth rate was observed at 10 °C. As the temperature was elevated to 43°C, the wild type and D1-SS209/212AA mutants showed significant decline while no increase was recorded in the remaining strains. As the temperature dropped back to 10 °C, the growth of all
25 the strains was practically arrested but a clear decline of the wild type was obvious.

EXAMPLE 7

Enhancement of D1 stability and Rubisco activation in the wild type and different mutants under high and low temperatures

30 Following the 43 °C - 30 °C - 43 °C cycle, D1 protein levels were high in S212C while they were found to be expressed at lower levels in the D1-SS209/212AC mutants (Figure 9G). In the other strains, D1 protein levels were essentially depleted (Figure 9G). The Rubisco and psaC protein were found to be stably expressed and at

higher concentrations in D1-S212C and in D1-SS209/212AC mutants compared to the other strains which did not present detectable amounts (Figures 9I and 9K).

The outcome of the second temperature cycle (43 °C - 10 °C - 43 °C cycle) was very different as shown in Figures 9F, 9H, 9J and 9L. At 43 °C (during the first 5 48 hours) the chlorophyll content was increased for all the cultures then at subsequent two days at low temperature (10 °C) the chlorophyll quantity was slightly dropped in all strains and after the shift to 43 °C all the cultures maintain their chlorophyll levels (Figure 9F). D1-S212C mutant had the best chlorophyll accumulation rates and higher biomass compared to all other strains at the mentioned conditions (Figure 9F). 10 Although the phenotypes of the strains were not very different from each other as it is shown by their color (Figure 9B), the D1-S212C and D1-SS209/212AC mutants maintained the highest chlorophyll content as well as the highest biomass production (Figure 9F). After the temperature treatment in this cycle, the steady content of Rubisco and psaC proteins was not changed in the wild type and all the mutants 15 (Figures 9J and 9L). The D1 protein was severely depleted in the wild type but not in the mutant strains (Figure 9H).

D1 protein levels were barely detectable in wild type, D1-SS209/212AA and D1-S212P mutants grown at 10 °C - 43 °C - 10 °C conditions (Figure 10D), while, detectable D1 levels were demonstrated for D1-S212C and D1-SS209/212AC mutants 20 (Figure 10D). Rubisco levels were not detected in wild type and D1-SS209/212AA strains, however, similar Rubisco levels were detected in D1-S212C, D1-S212P and D1-SS209/212AC mutants (Figure 10E). In contrast, all the strains exhibited similar and stable levels of psaC protein of PSI (Figure 10F).

25 It is appreciated that certain features of the invention, which are, for clarity, described in the context of separate embodiments, may also be provided in combination in a single embodiment. Conversely, various features of the invention, which are, for brevity, described in the context of a single embodiment, may also be provided separately or in any suitable subcombination.

30

Although the invention has been described in conjunction with specific embodiments thereof, it is evident that many alternatives, modifications and variations will be apparent to those skilled in the art. Accordingly, it is intended to embrace all

such alternatives, modifications and variations that fall within the spirit and broad scope of the appended claims. All publications, patents and patent applications and GenBank Accession numbers mentioned in this specification are herein incorporated in their entirety by reference into the specification, to the same extent as if each
5 individual publication, patent or patent application or GenBank Accession number was specifically and individually indicated to be incorporated herein by reference. In addition, citation or identification of any reference in this application shall not be construed as an admission that such reference is available as prior art to the present invention.

WHAT IS CLAIMED IS:

1. An isolated polynucleotide comprising a nucleic acid sequence encoding a polypeptide of a Type II reaction center of a photosynthetic organism, said nucleic acid sequence being capable of imparting said type II reaction center with an activity under a temperature range different than that of said type II reaction center endogenous to said photosynthetic organism.

2. The isolated polynucleotide of claim 1, wherein said polypeptide of said type II reaction center is selected from the group consisting of D1, D2, L and M.

3. The isolated polynucleotide of claim 1, wherein said polypeptide is D1.

4. The isolated polynucleotide of claim 1, wherein said activity comprise solar energy conversion activity.

5. The isolated polynucleotide of claim 3, wherein said D1 polypeptide comprises an amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 and/or at a position corresponding to 212 of said SEQ ID NO: 36.

6. The isolated polynucleotide of claim 5, wherein said amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 comprises a Serine to Alanine mutation.

7. The isolated polynucleotide of claim 5, wherein said amino acid sequence alteration at a position corresponding to 212 of SEQ ID NO: 36 comprises a Serine to Cysteine mutation.

8. The isolated polynucleotide of claim 1, wherein said D1 polypeptide comprises an amino acid sequence alteration at any amino acid position corresponding to 208-212 of SEQ ID NO: 36.

9. The isolated polynucleotide of claim 1, wherein said type II reaction center is rendered more thermotolerant as compared to said type II reaction center endogenous to said photosynthetic organism.

10. The isolated polynucleotide of claim 1, wherein said type II reaction center is rendered more thermoplastic as compared to said type II reaction center endogenous to said photosynthetic organism.

11. The isolated polynucleotide of claim 1, wherein said photosynthetic organism is selected from the group consisting of a higher plant, a photosynthetic bacteria and an algae.

12. The isolated polynucleotide of claim 1, wherein said photosynthetic organism is selected from the group consisting of a mesophile, a thermophile and a psychrophile.

13. The isolated polynucleotide of claim 1, wherein said temperature range comprises 10 °C – 43 °C.

14. The isolated polynucleotide of claim 1, wherein said temperature range comprises an upwards shift.

15. The isolated polynucleotide of claim 14, wherein said upwards shift is by at least 6 °C.

16. The isolated polynucleotide of claim 1, wherein said temperature range comprises a downwards shift.

17. The isolated polynucleotide of claim 16, wherein said downwards shift is by at least 2.5 °C.

18. The isolated polynucleotide of claim 1, wherein said temperature range comprise a wider temperature range.

19. The isolated polynucleotide of claim 5, wherein an amino acid at position 209 or 212 is selected from the group consisting of glycine, cysteine, alanine, threonin, asparagine, aspartanine, proline, valine, isoleucine, leucine, glutamine and glutamic acid.

20. A nucleic acid construct comprising the polynucleotide of claim 1.

21. The nucleic acid construct of claim 20, further comprising a cis-regulatory element.

22. The nucleic acid construct of claim 21, wherein said cis-regulatory element is a promoter.

23. A cell of a photosynthetic organism comprising the nucleic acid sequence of claim 1.

24. An isolated polypeptide comprising an amino acid sequence encoding a polypeptide of a type II reaction center of a photosynthetic organism, said nucleic acid sequence being capable of imparting said type II reaction center with an activity under a temperature range different than that of said type II reaction center endogenous to said photosynthetic organism.

25. The isolated polypeptide of claim 24, wherein said polypeptide is selected from the group consisting of D1, D2, L and M.

26. The isolated polypeptide of claim 24, wherein said polypeptide is D1.

27. The isolated polypeptide of claim 24, wherein said activity comprise solar energy conversion activity.

28. The isolated polypeptide of claim 26, wherein said D1 comprises an amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 and/or at a position corresponding to 212 of said SEQ ID NO: 36.

29. The isolated polypeptide of claim 28, wherein said amino acid sequence alteration at a position corresponding to 209 of SEQ ID NO: 36 comprises a Serine to Alanine mutation.

30. The isolated polypeptide of claim 28, wherein said amino acid sequence alteration at a position corresponding to 212 of SEQ ID NO: 36 comprises a Serine to Cysteine mutation.

31. The isolated polypeptide of claim 26, wherein said D1 comprises an amino acid sequence alteration at any amino acid position corresponding to 208-212 of SEQ ID NO: 36.

32. The isolated polypeptide of claim 24, wherein said type II reaction center is rendered more thermotolerant as compared to said type II reaction center endogenous to said photosynthetic organism.

33. The isolated polypeptide of claim 24, wherein said type II reaction center is rendered more thermoplastic as compared to said type II reaction center endogenous to said photosynthetic organism.

34. The isolated polypeptide of claim 24, wherein said photosynthetic organism is selected from the group consisting of a higher plant, a cyanobacteria and an algae.

35. The isolated polypeptide of claim 24, wherein said photosynthetic organism is selected from the group consisting of a mesophile, a thermophile and a psychrophile.

36. The isolated polypeptide of claim 24, wherein said temperature range comprises 10 °C – 43 °C.

37. The isolated polypeptide of claim 24, wherein said temperature range comprises an upwards shift.

38. The isolated polypeptide of claim 37, wherein said upwards shift is by at least 6 °C.

39. The isolated polypeptide of claim 24, wherein said temperature range comprises a downwards shift.

40. The isolated polypeptide of claim 39, wherein said downwards shift is by at least 2.5 °C.

41. The isolated polypeptide of claim 24, wherein said temperature range comprise a wider temperature range.

42. The isolated polypeptide of claim 28, wherein an amino acid at position 209 or 212 is selected from the group consisting of glycine, cysteine, alanine, threonin, asparagine, aspartanine, proline, valine, isoleucine, leucine, glutamine and glutamic acid.

43. The isolated polypeptide of claim 24, wherein said amino acid sequence is selected from the group consisting of SEQ ID NO: 37, 38, 50, 52 and 56.

44. A photosynthetic organism comprising an exogenous nucleic acid sequence encoding a polypeptide of a type II reaction center of the photosynthetic organism, said nucleic acid sequence imparting said type II reaction center with an activity under a temperature range different than that of said type II reaction center endogenous to the photosynthetic organism.

45. The photosynthetic organism of claim 44, selected from the group consisting of a higher plant, a cyanobacteria and an algae.

46. The photosynthetic organism of claim 44, selected from the group consisting of a mesophile, a thermophile and a psychrophile.

47. The photosynthetic organism of claim 44, wherein said exogenous nucleic acid sequence further renders said organism more thermoplastic than an identical photosynthetic organism not comprising said exogenous nucleic acid sequence.

48. The photosynthetic organism of claim 44, wherein said exogenous nucleic acid sequence further renders said organism capable of expressing more Rubisco than an identical photosynthetic organism not comprising said exogenous nucleic acid sequence.

49. The photosynthetic organism of claim 44, wherein said exogenous nucleic acid sequence further renders said organism capable of expressing more psaC than an identical photosynthetic organism not comprising said exogenous nucleic acid sequence.

50. The photosynthetic organism of claim 44, wherein said exogenous nucleic acid sequence further renders said organism capable of growing faster than a photosynthetic organism not comprising said exogenous nucleic acid sequence.

51. The photosynthetic organism of claim 44, wherein said exogenous nucleic acid sequence further renders said organism capable of accumulating more chlorophyll than an identical photosynthetic organism not comprising said exogenous nucleic acid sequence.

52. A method of improving biomass/vigor/yield of a photosynthetic organism comprising introducing into the photosynthetic organism the isolated polynucleotide of claim 1, thereby increasing biomass, vigor and/or yield of the photosynthetic organism.

53. The method of claim 52, further comprising growing said photosynthetic organism under abiotic stress conditions.

54. The method of claim 53, wherein said abiotic stress conditions comprise heat, cold and alternating temperatures.

55. A method of increasing activity of a type II reaction center of a photosynthetic organism under non-physiological temperature, the method comprising introducing into the photosynthetic organism the isolated polynucleotide of claim 1, thereby increasing the activity of a type II reaction center under a non-physiological temperature.

56. A method of identifying mutations which impart a photosynthetic organism with a photosynthetic activity under a non-physiological temperature, the method comprising:

(i) subjecting the photosynthetic organisms to a mutagen, so as to obtain mutated photosynthetic organisms;

(ii) identifying an organism of said mutated photosynthetic organisms exhibiting at least one parameter associated with enhanced photosynthetic activity under non-physiological temperature as compared to corresponding wild type photosynthetic organisms; and

(iii) identifying mutations in a polypeptide of a type II reaction center of said organism, said mutations being correlated with photosynthetic activity under a non-physiological temperature.

57. The method of claim 56, wherein said at least one parameter associated with enhanced photosynthetic activity is selected from the group consisting of biomass, growth rate, rubisco level, chlorophyll content, psaC level and thermoplasticity.

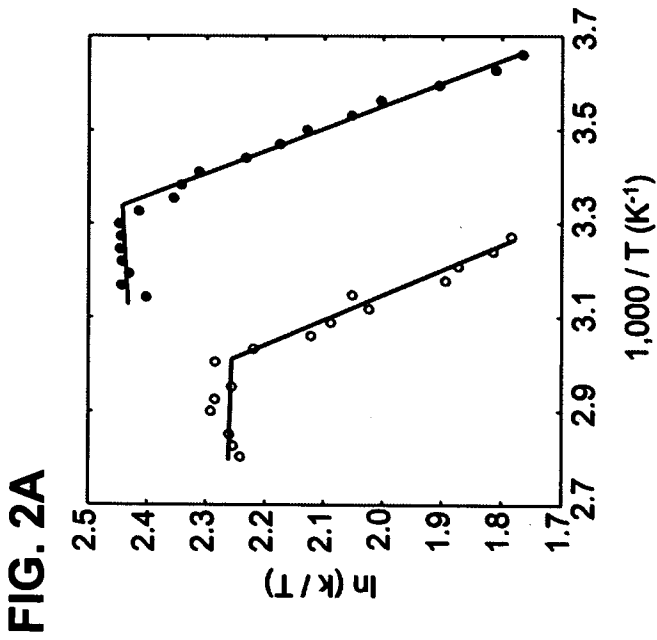
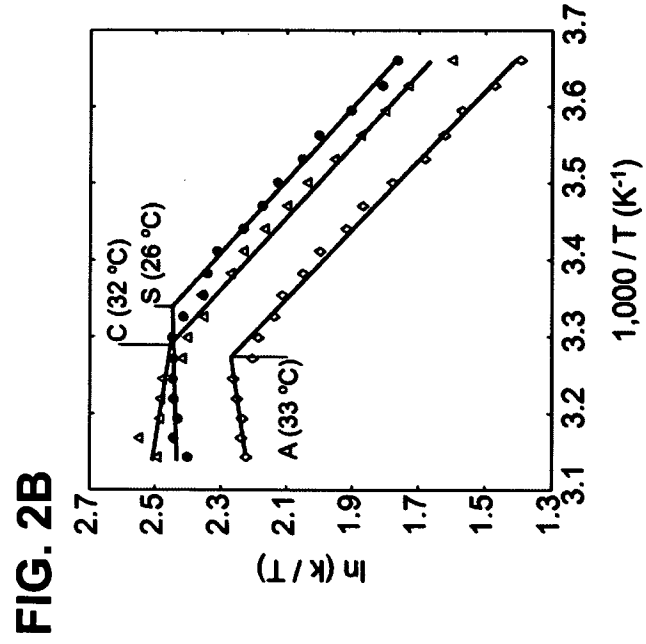
58. The method of claim 56, wherein said polypeptide of a type II reaction center is selected from the group consisting of D1, D2, L and M.

59. A method of increasing hydrogen production in a photosynthetic organism, the method comprising introducing into the photosynthetic organism

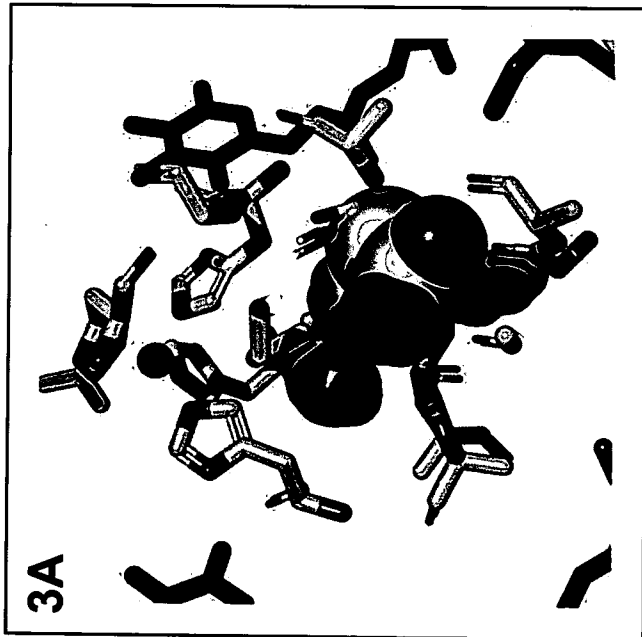
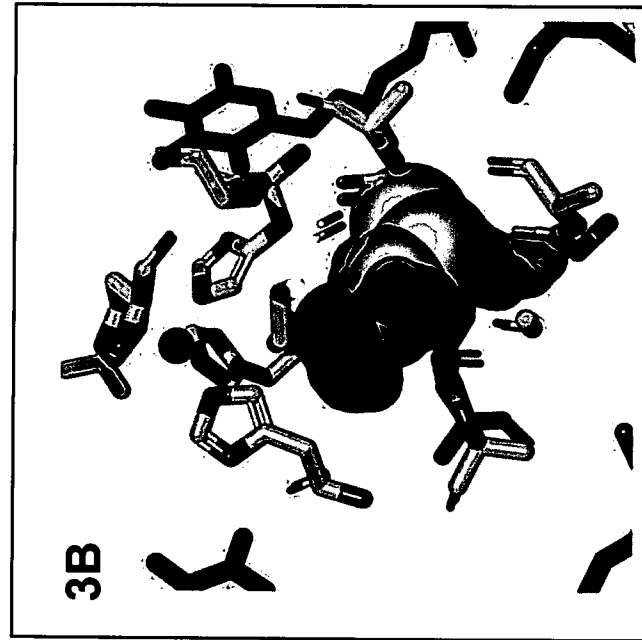
capable of generating hydrogen the isolated polynucleotide of claim 1, thereby increasing hydrogen production in the photosynthetic organism.

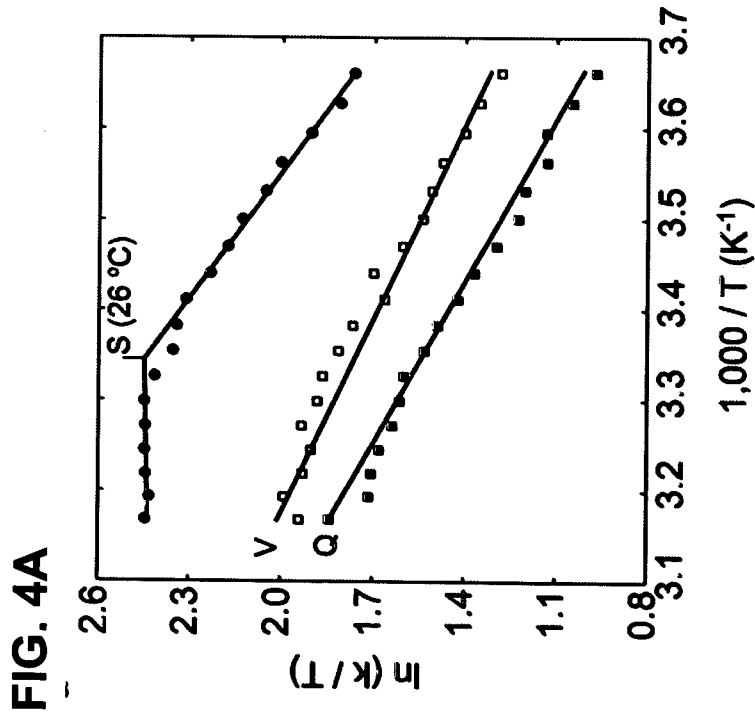
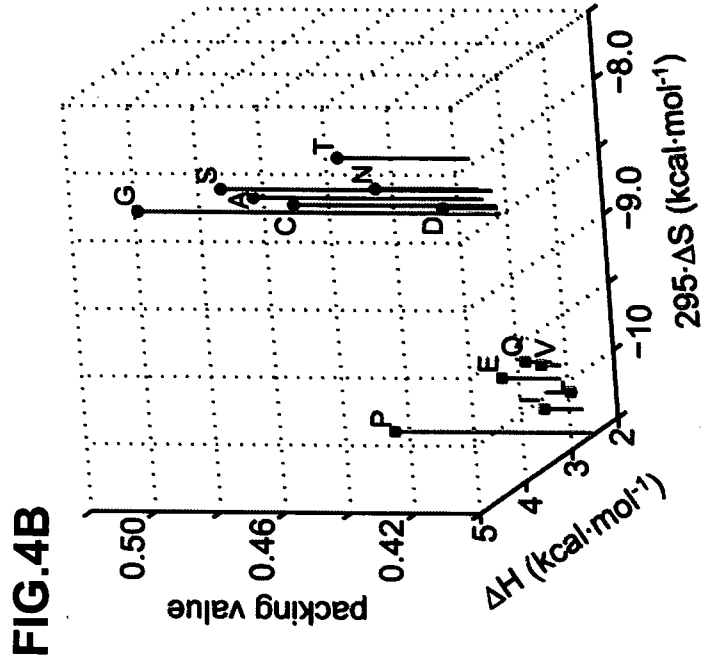
Figure 1C

L	RCEL_RHOSH	178	SFFFTNALALALHGA	SEQ	ID NO:	11
L	RCEL_RHOVI	178	SFLFVNAMALGLHGG	SEQ	ID NO:	12
L	RCEL_CHLAU	218	TGLFASTWLLACHGS	SEQ	ID NO:	13
D1	PSB1_SYNVD	203	AGVFGGALFCAMHGS	SEQ	ID NO:	14
D1	PSB2_SYNEL	203	AGVFGGALFAMHGS	SEQ	ID NO:	15
D1	PSB2_SYNY3	203	AGVFGGSLFSAMHGS	SEQ	ID NO:	16
D1	PSBA_CYACA	203	AGVFGGALFAMHGS	SEQ	ID NO:	17
D1	PSBA_CHLRE	203	AGVFGGSLFSAMHGS	SEQ	ID NO:	18
D1	PSBA_SPIOL	203	AGVFGGSLFSAMHGS	SEQ	ID NO:	19
D1	PSBA_VIGUN	203	AGVFGGSLFSAMHGS	SEQ	ID NO:	20
D2	1s51_d	202	AGVLGGALLCAIHGA	SEQ	ID NO:	21
D2	NP_681245	202	AGVLGGALLCAIHGA	SEQ	ID NO:	22
D2	PSBD_SYNY3	202	AGILGGAMPCAIHGA	SEQ	ID NO:	23
D2	PSBD_CYACA	201	AGILGGALLCAIHGA	SEQ	ID NO:	24
D2	PSBD_CHLRE	203	AGVLGAALLCAIHGA	SEQ	ID NO:	25
D2	PSBD_SPIOL	203	AGVLGAALLCAIHGA	SEQ	ID NO:	26
M	RCEM_RHOVI	205	GFAYGCGLLFAHGA	SEQ	ID NO:	27
M	RCEM_RHOSH	207	AFLYGSALLFAMHGA	SEQ	ID NO:	28
M	RCEM_CHLAU	197	FLLGSLTLLLAMHAG	SEQ	ID NO:	29

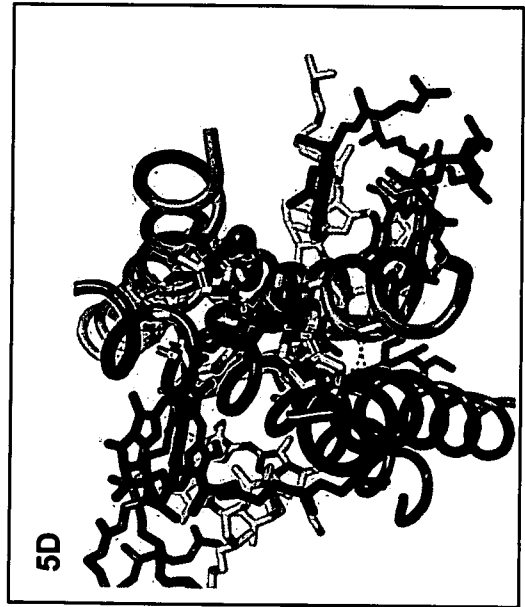
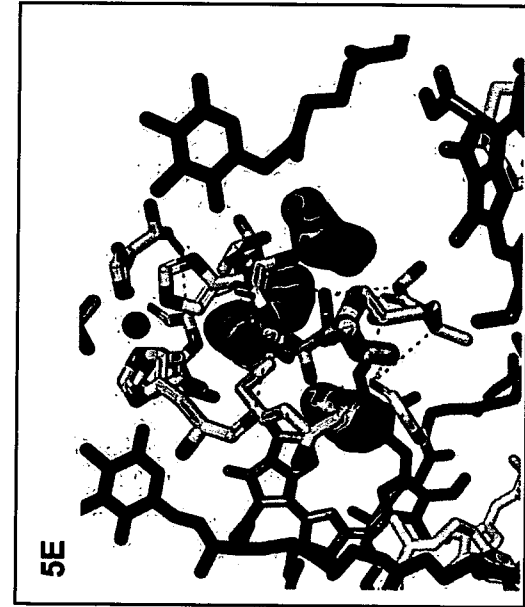
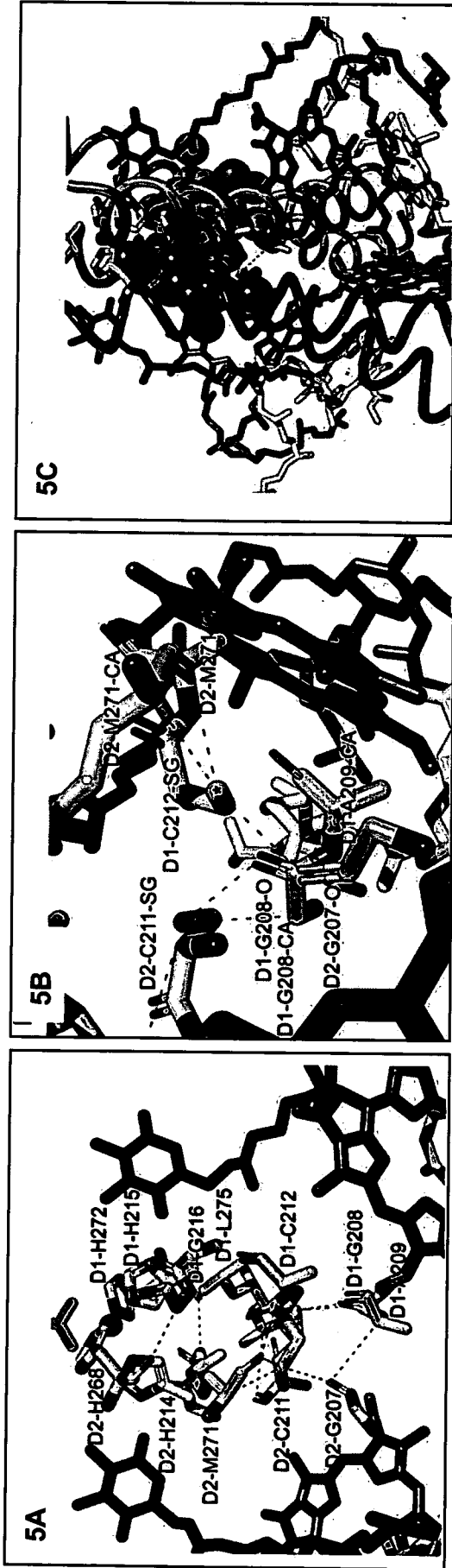


Figures 3A-B





Figures 5A-E



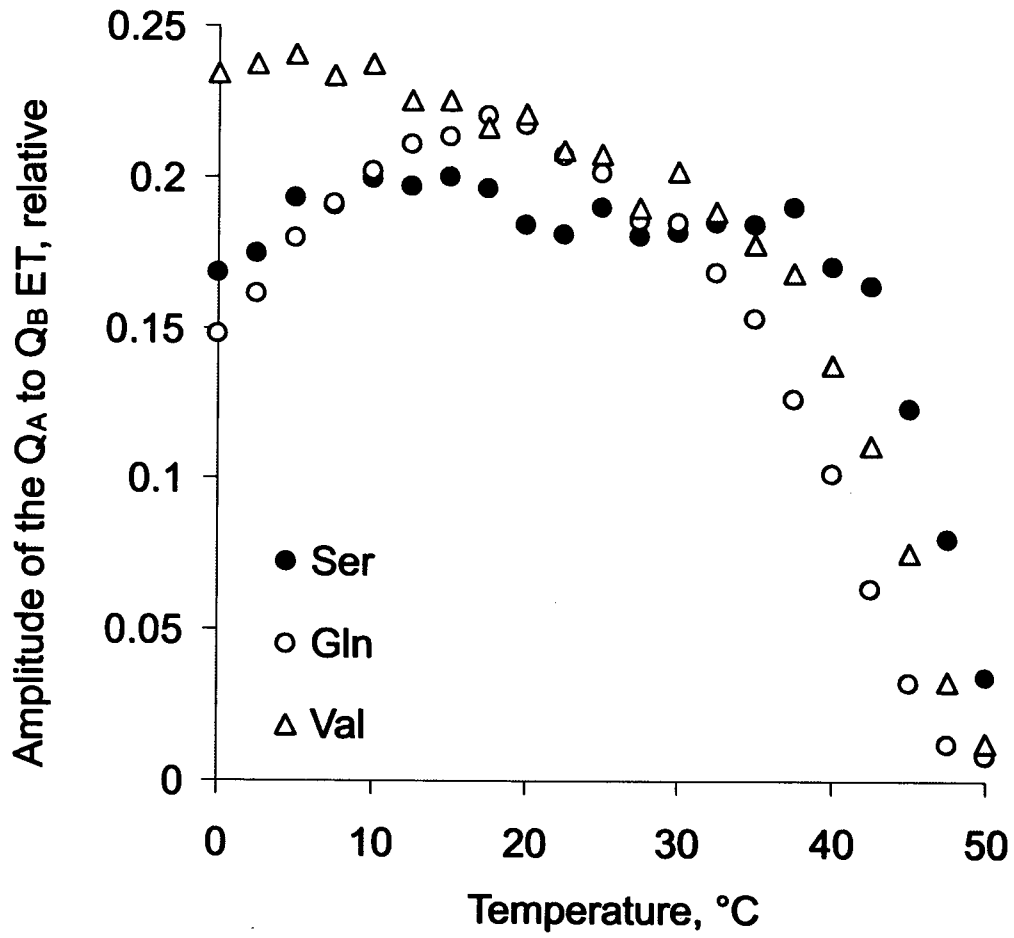
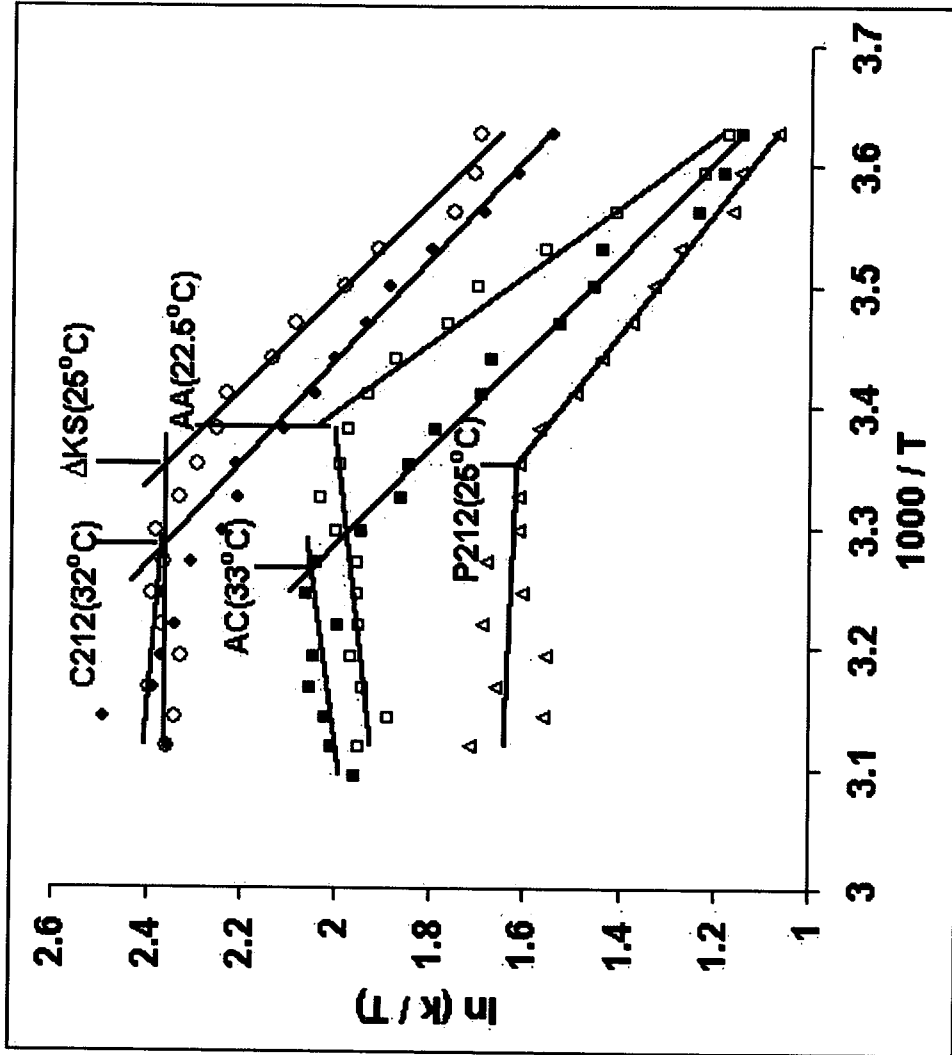


Fig. 6

Figure 7



30 °C



Fig. 8A

43 °C

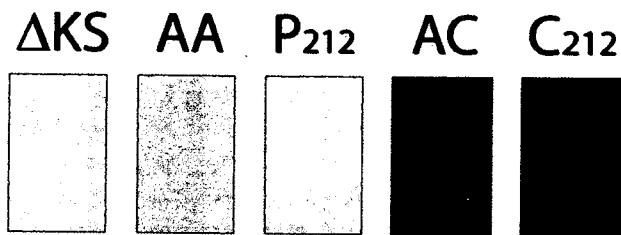


Fig. 8B

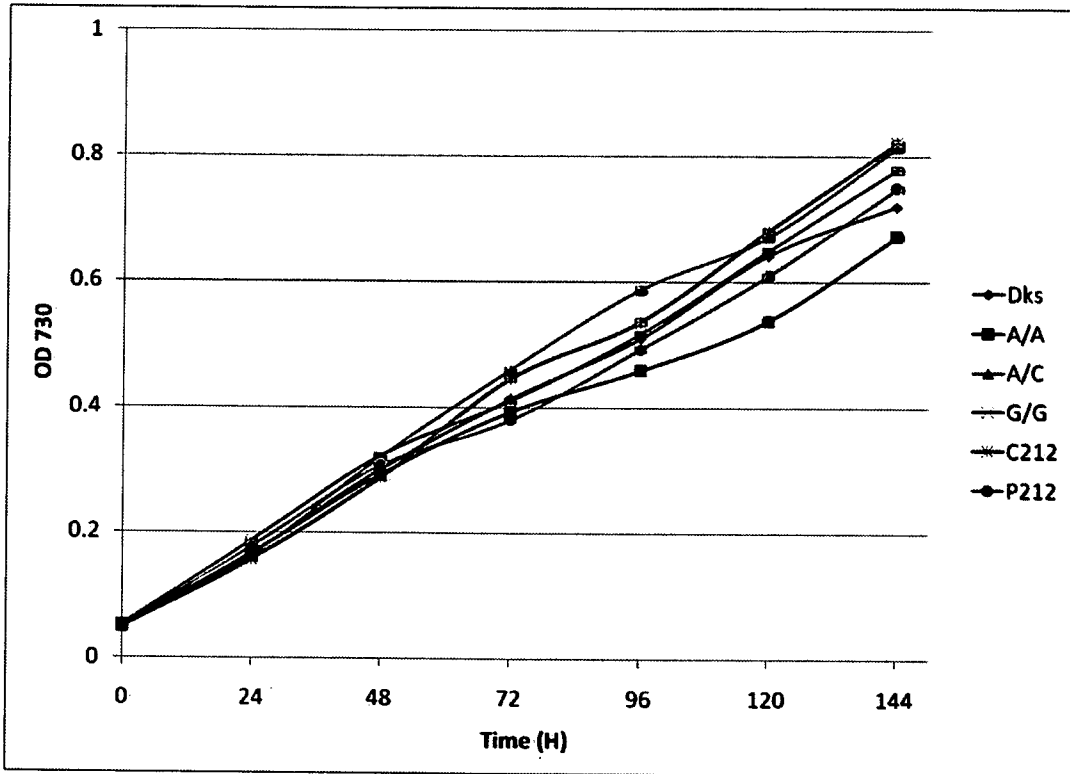


Fig. 8C

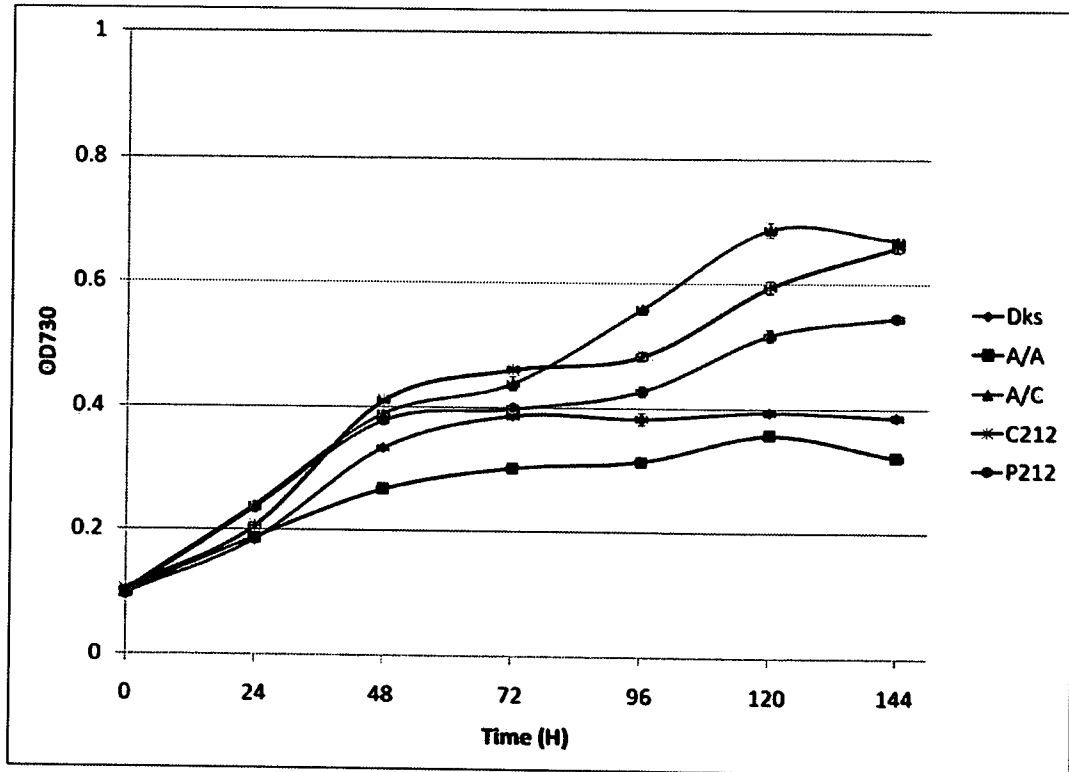


Fig. 8D

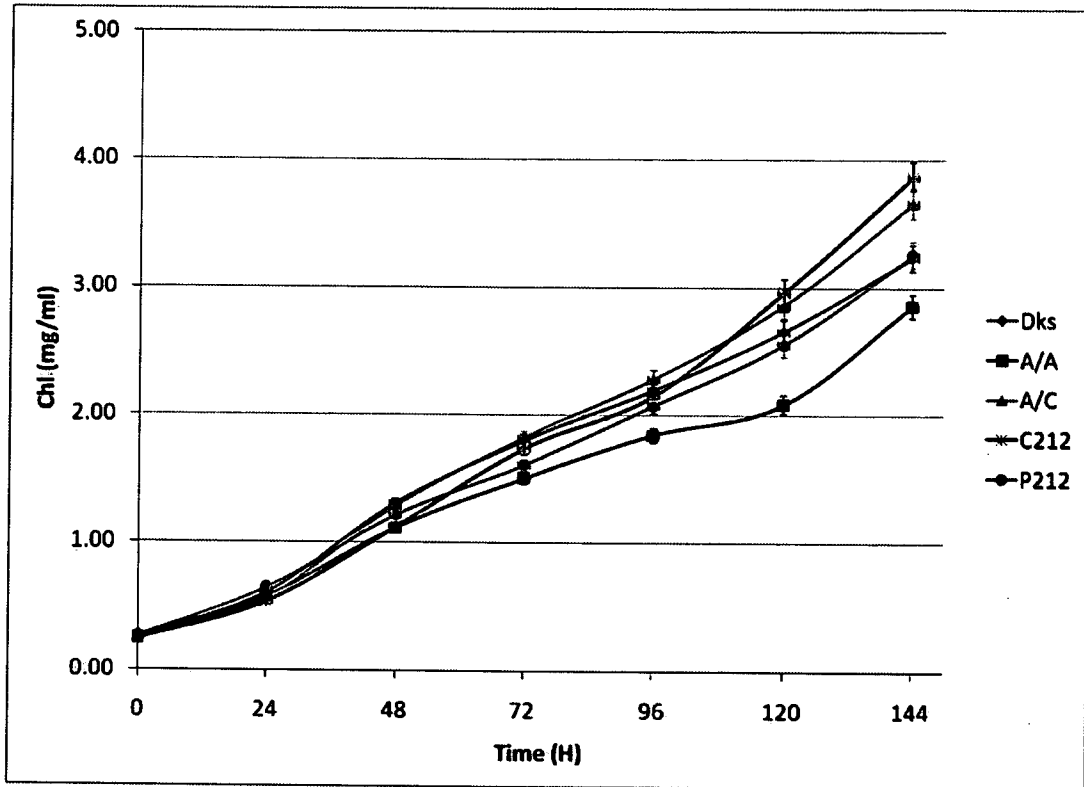


Fig. 8E

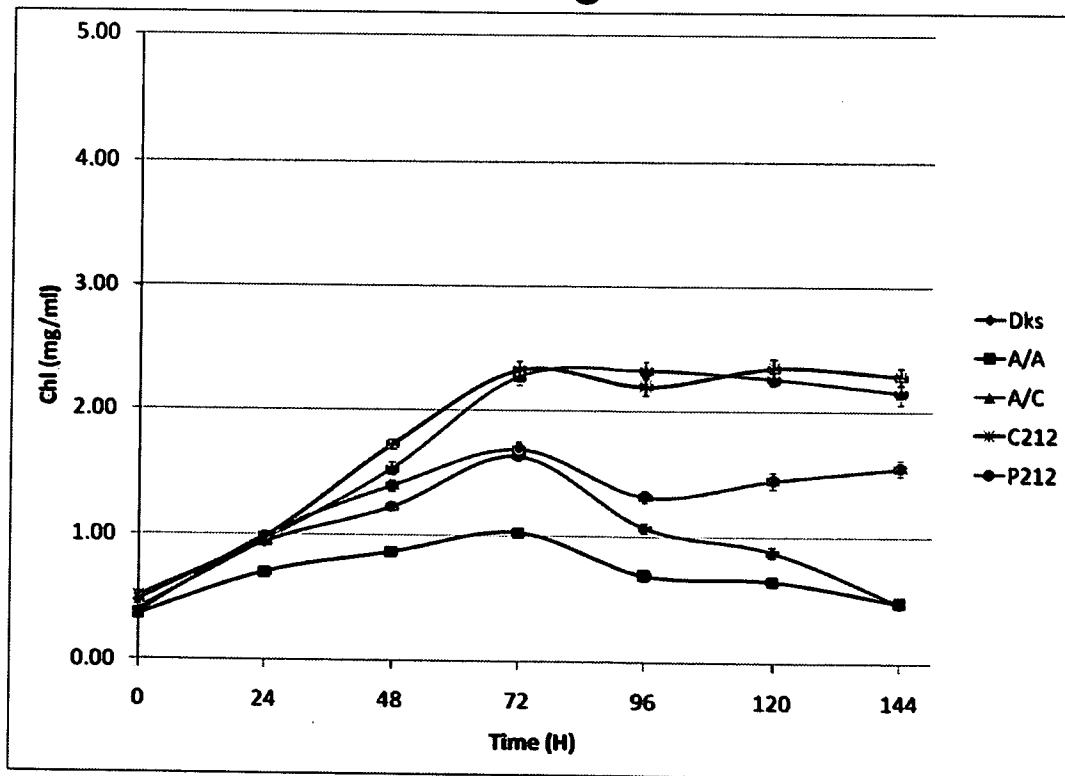
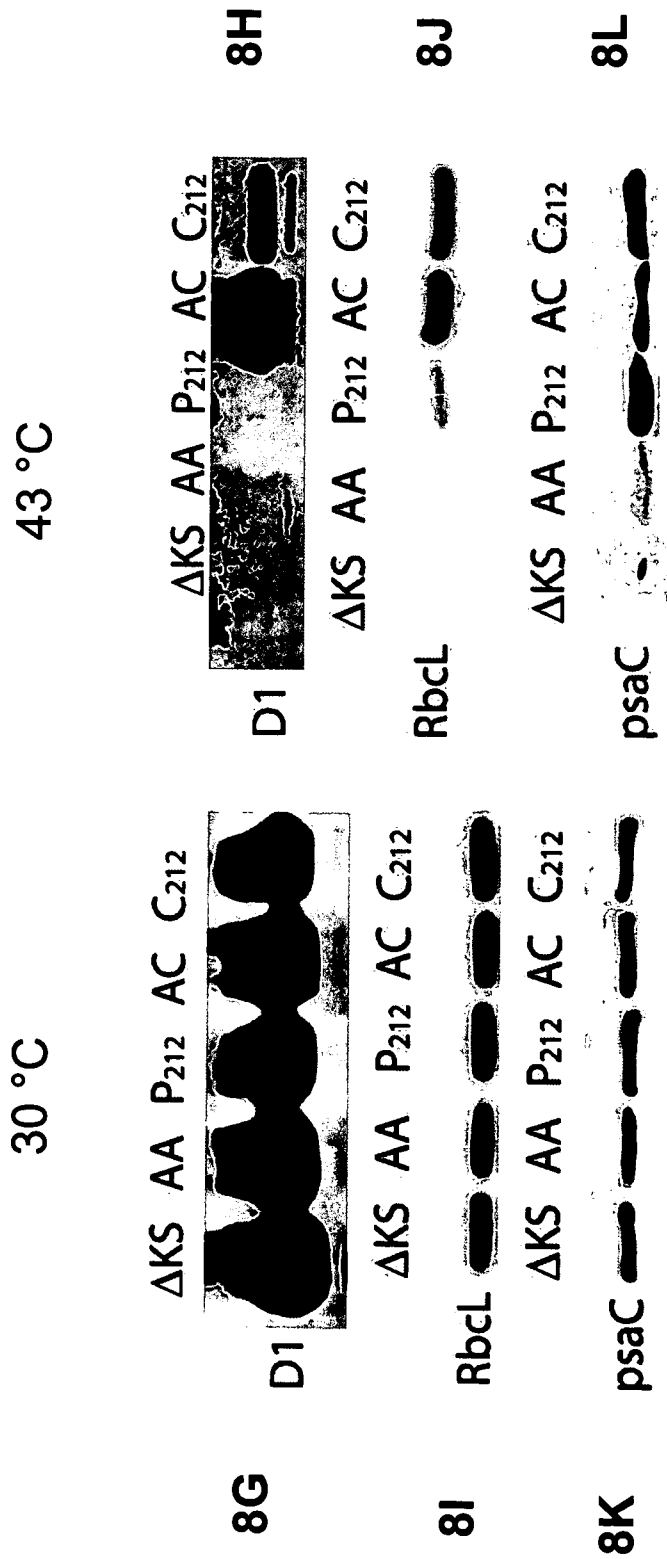


Fig. 8F

12/19



Figures 8G-L

Plasticity test 43 °C - 30 °C - 43 °C

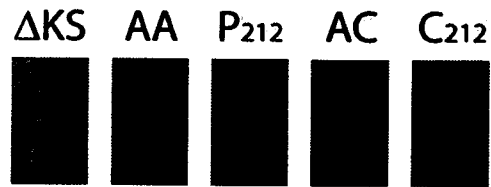


Fig. 9A

Plasticity test 43 °C – 10 °C – 43 °C

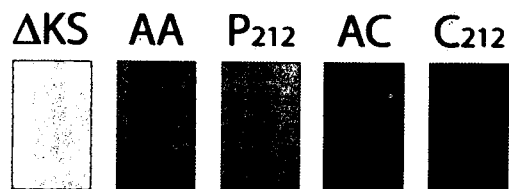


Fig. 9B

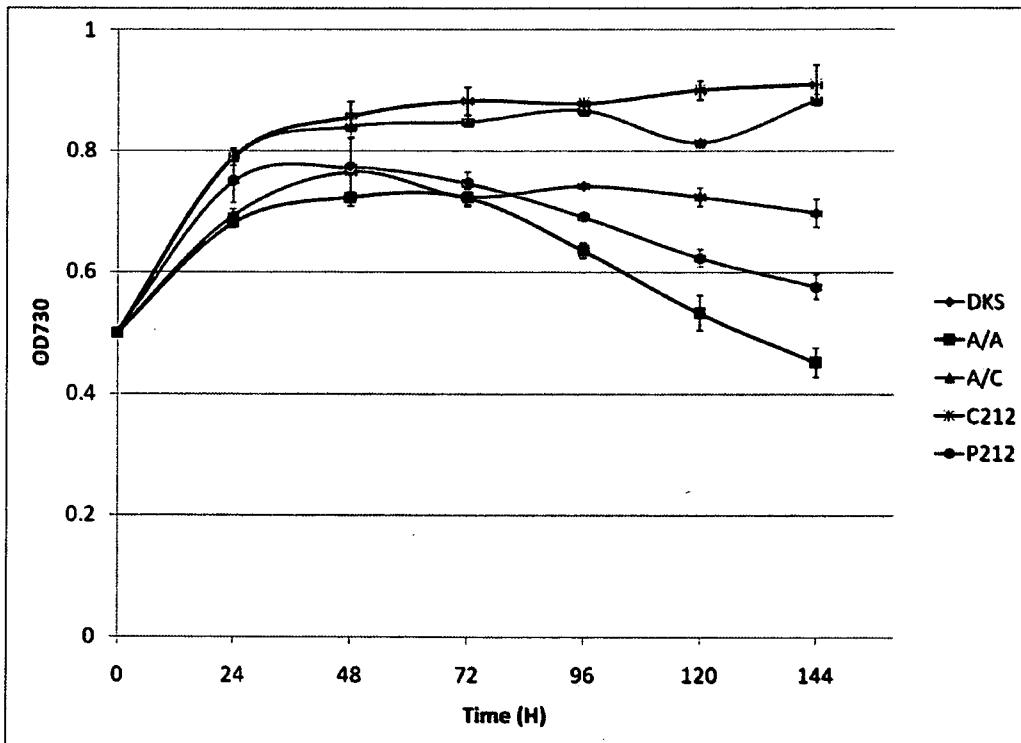


Fig. 9C

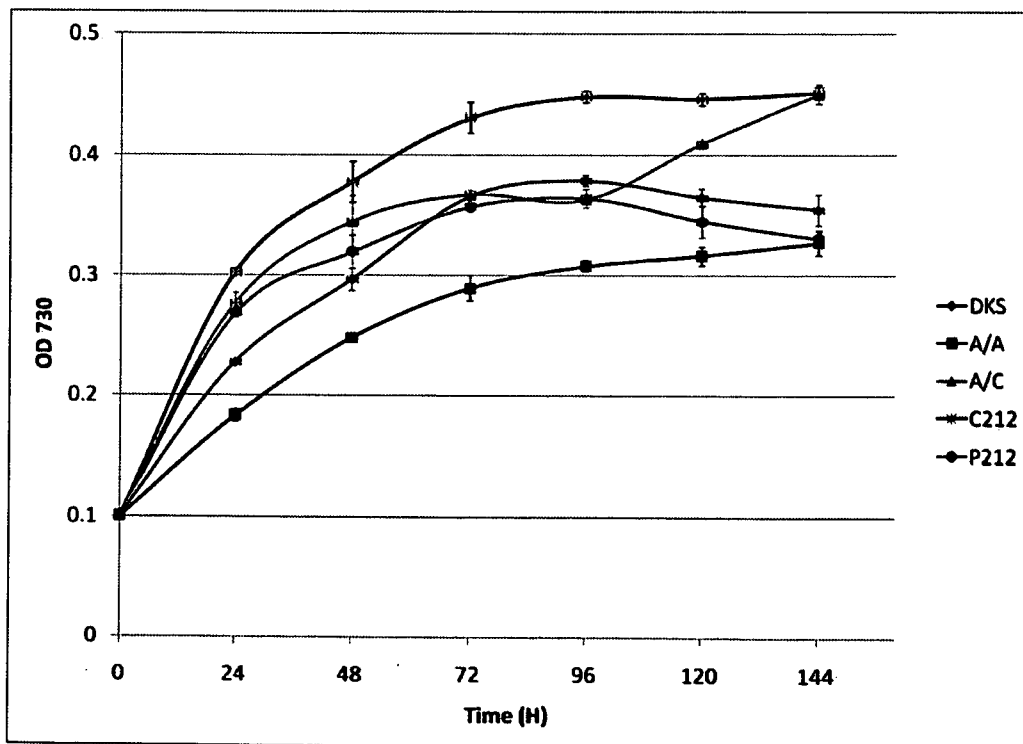


Fig. 9D

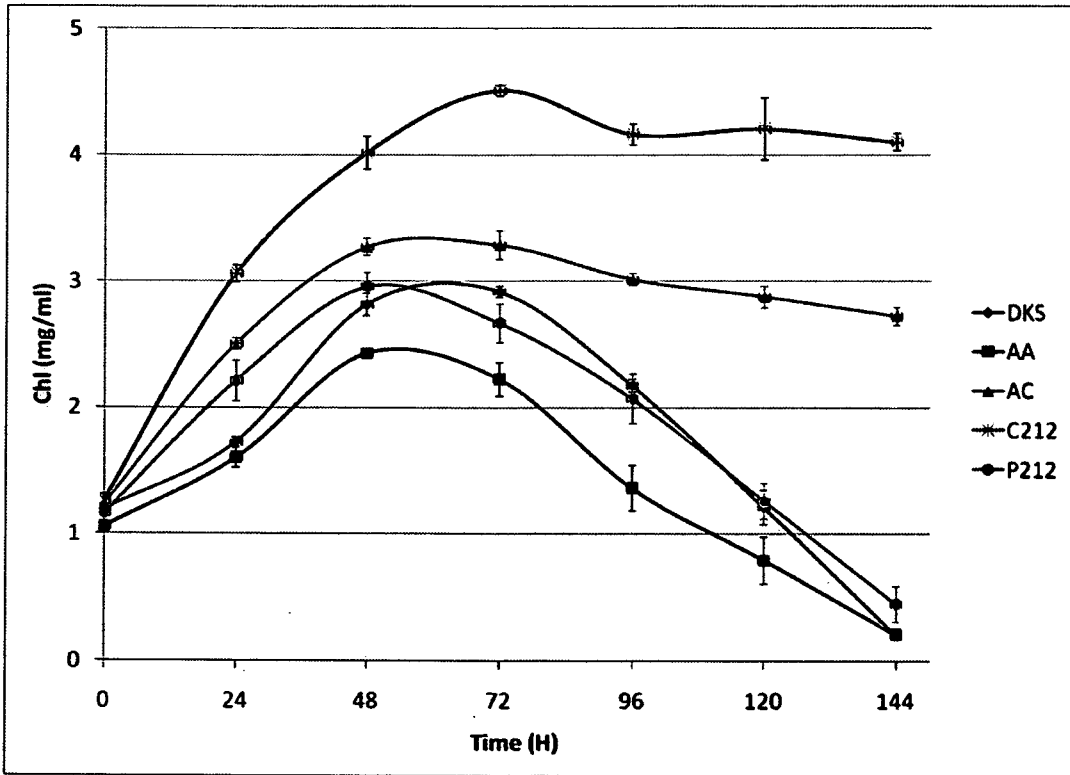


Fig. 9E

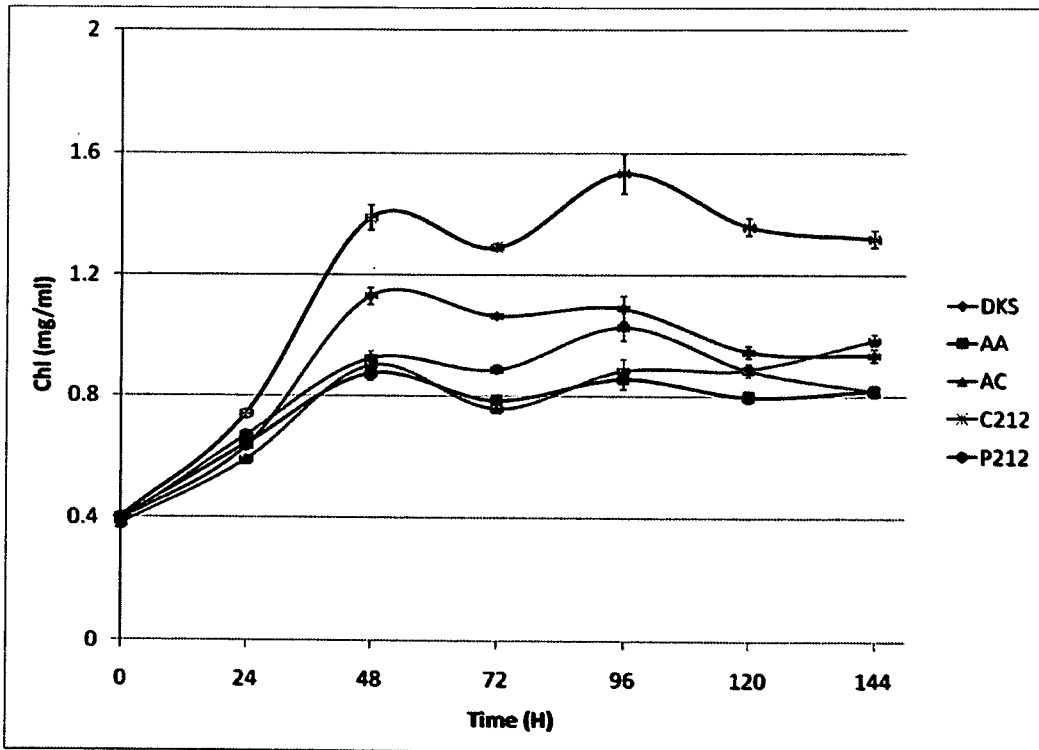
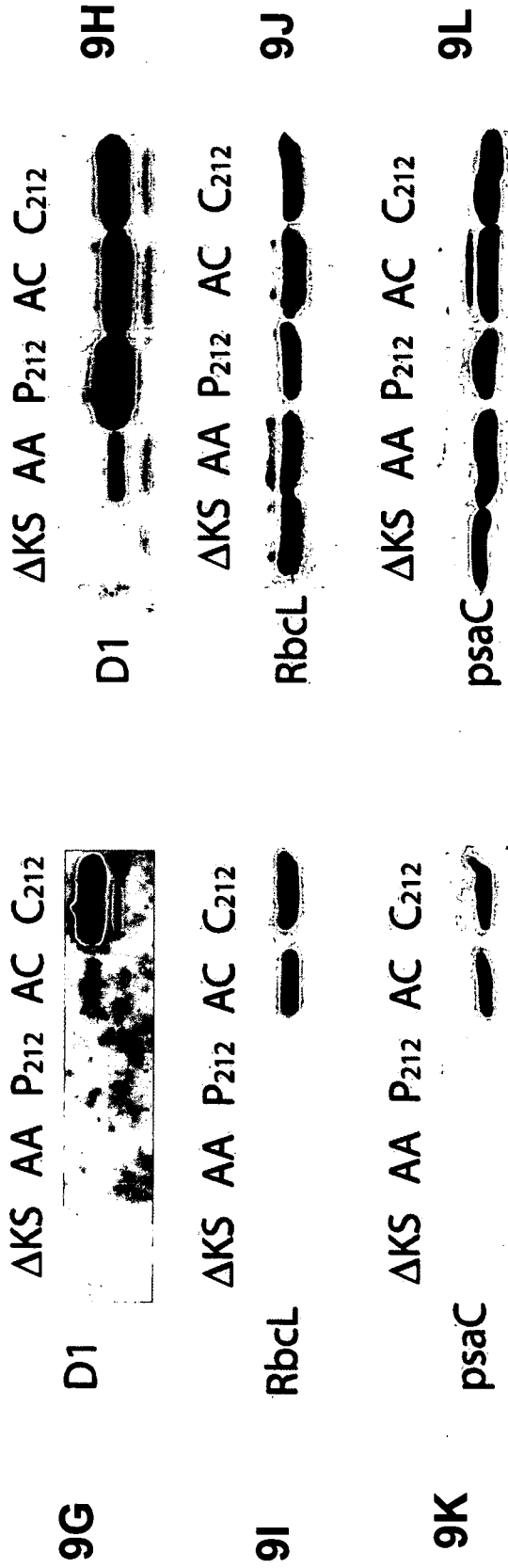


Fig. 9F

Plasticity test 43 °C - 10 °C - 43 °C

Plasticity test 43 °C - 30 °C - 43 °C



Figures 9G-L

Plasticity test 10 °C - 43 °C - 10 °C

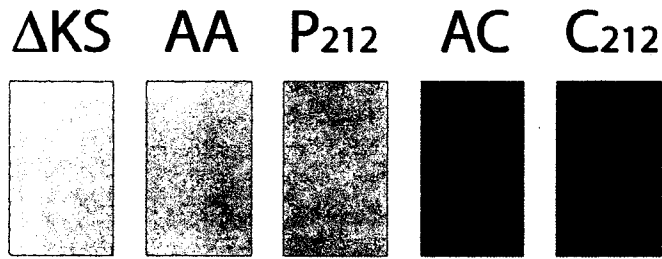


Fig. 10A

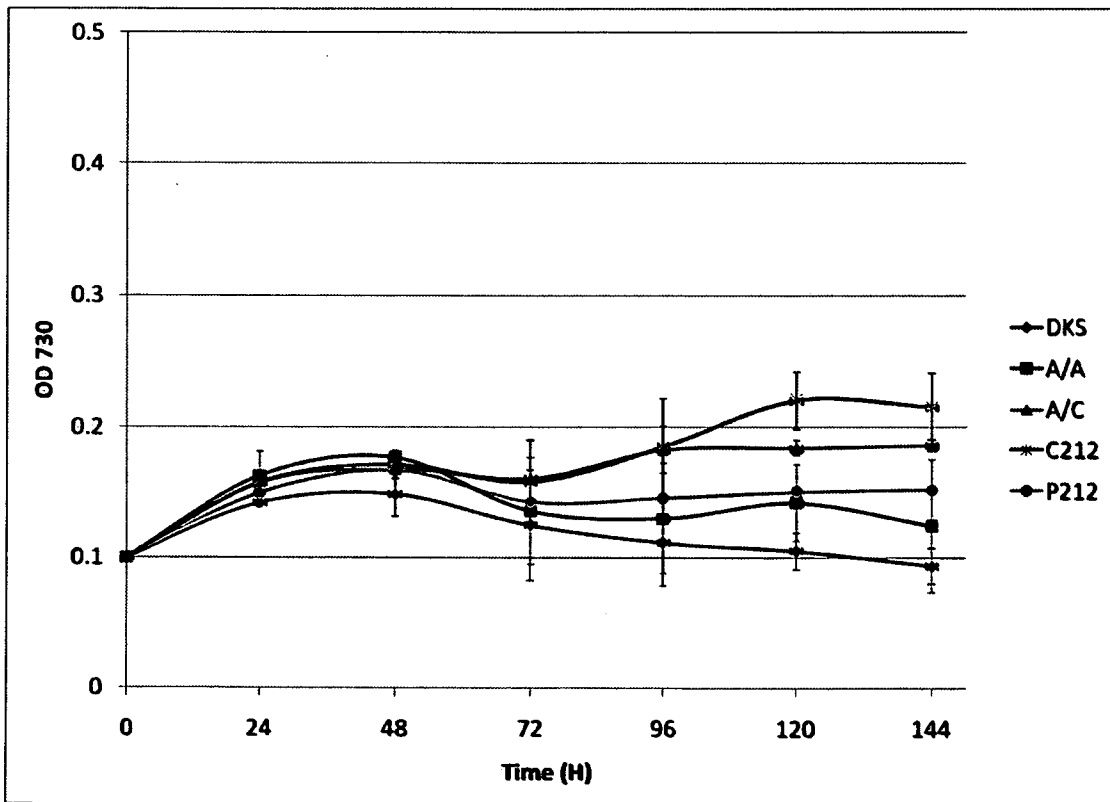


Fig. 10B

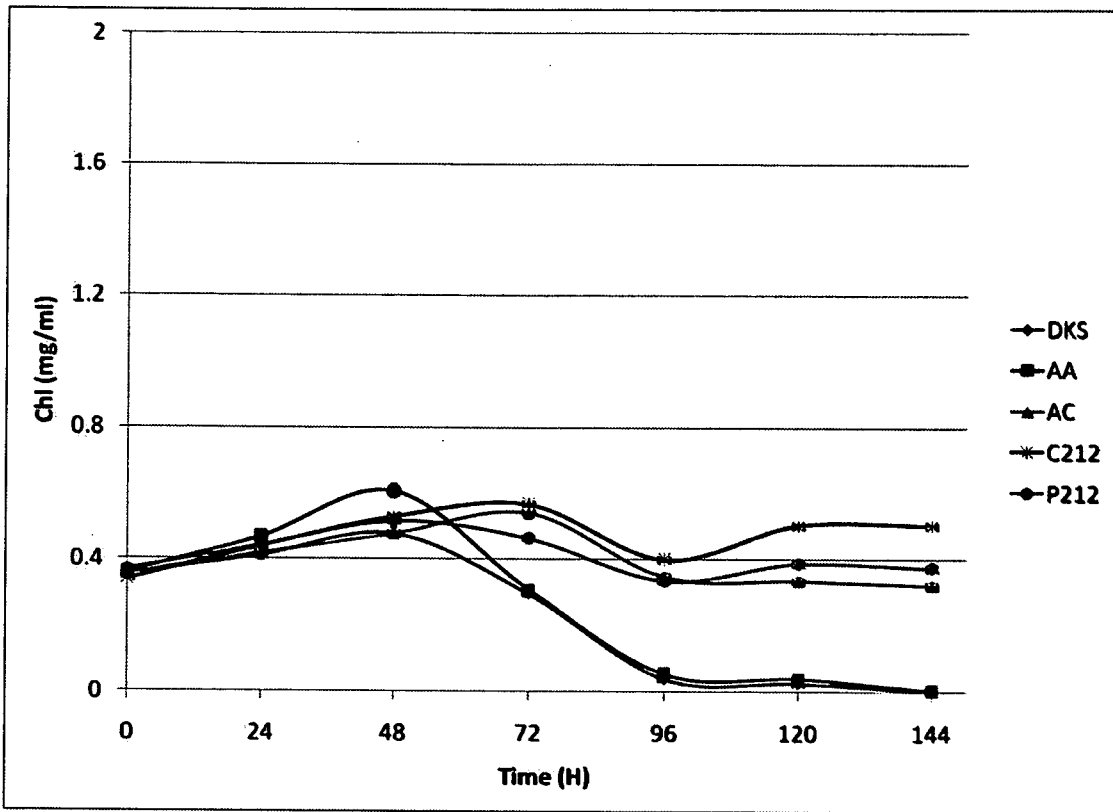


Fig. 10C

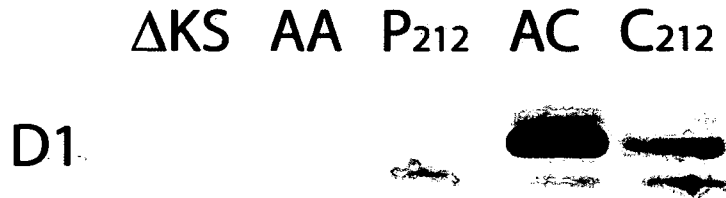


Fig. 10D

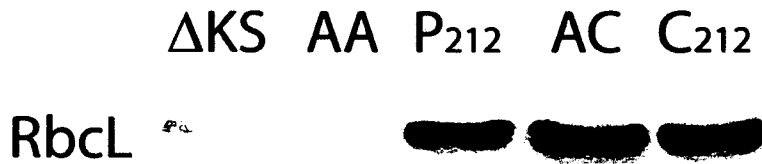


Fig. 10E

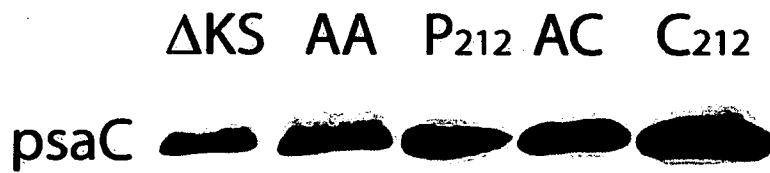


Fig. 10F