3-20-2012

Modified Rubisco Large Subunit N-Methyltransferase Useful for Targeting Molecules to the Active-Site Vicinity of Ribulose-1,5-Bisphosphate

Robert L. Houtz
University of Kentucky, rhoutz@uky.edu

Click here to let us know how access to this document benefits you.

Follow this and additional works at: https://uknowledge.uky.edu/horticulture_patents

Part of the Horticulture Commons

Recommended Citation
Houtz, Robert L., "Modified Rubisco Large Subunit N-Methyltransferase Useful for Targeting Molecules to the Active-Site Vicinity of Ribulose-1,5-Bisphosphate" (2012). Horticulture Faculty Patents. 1.
https://uknowledge.uky.edu/horticulture_patents/1

This Patent is brought to you for free and open access by the Horticulture at UKnowledge. It has been accepted for inclusion in Horticulture Faculty Patents by an authorized administrator of UKnowledge. For more information, please contact UKnowledge@lsv.uky.edu.
The present invention generally relates to a modified Rubisco large subunit N-Methyltransferase (Rubisco LSMT, or RLSMT). The present invention also relates to a modified RLSMT-carbonic anhydrase (RLSMT-CA). This modified RLSMT-CA improves the efficiency of the reduction of CO₂ during photosynthesis, which may increase plant growth rates. The present invention also relates to nucleic acids encoding the modified RLSMT-CA or modified RLSMT. Also, the present invention relates to cells including the modified RLSMT-CA or modified RLSMT, plants containing the modified RLSMT-CA or modified RLSMT, and methods using compositions of the present invention. In addition, the present invention relates to antibodies conjugated to CA which may bind to Rubisco, and antibodies which bind a modified RLSMT-CA. The invention also relates to modified forms of the LS and SS of Rubisco where the modified forms are fusions with CA or biologically active fragments thereof. The present invention provides methods of altering Rubisco carboxylase activity and altering plant growth.
Fig. 1

**A**

Activity (% of zero time)

\[ y = 82.9205 \times \exp(-1.5234 \times x) + 16.8855 \]

Time (h)

**B**

Activity (% of zero time)

\[ y = 88.9804 \times \exp(-0.002837 \times x) - 1.9712 \]

Time (h)

**C**

\( k_{on} \times 10^{-4} \) (sec\(^{-1}\))

K\(_D\) = \( k_{dis} / k_{ass} \) = 0.1 nM

\( k_{dis} = 8.0 \times 10^{-7} \) sec\(^{-1}\)

\( k_{ass} = 7.368 \) M\(^{-1}\) sec\(^{-1}\)

Rubisco LSMT (nM)
Fig. 3

A

6 µg Spinach Rubisco

B

46 µg Pea Rubisco

nmoles \(^{14}\)CO\(_2\) fixed

Time (minutes)

0.0 0.5 1.0 1.5 2.0

0 5 10 15 20 25 30
Fig. 5
MODIFIED RUBISCO LARGE SUBUNIT N-METHYLTRANSFERASE USEFUL FOR TARGETING MOLECULES TO THE ACTIVE-SITE VICINITY OF RIBULOSE-1,5-BISPHOSPHATE

This work was supported by the U.S. Department of Energy grant DE-FGO2-02-ER20075, and therefore the government may have certain rights to the invention.

This application is a divisional of U.S. application Ser. No. 10/555,795, filed Nov. 22, 2006, now abandoned, which is a 371 of PCT/US2004/014593, filed May 7, 2004, which claims the benefit of U.S. Provisional Application No. 60/468,966, filed May 8, 2003.

FIELD OF THE INVENTION

The present invention generally relates to a modified Rubisco large subunit "N-Methyltransferase (Rubisco LSMT, or RLSTM). The modified RLSTM is useful for targeting molecules to Rubisco. The present invention also relates to a modified RLSTM-carnitine anhydridase (RLSTM-CA). This modified RLSTM-CA improves the efficiency of the reduction of CO2 during photosynthesis, which may increase plant growth rates. The present invention also relates to nucleic acids encoding the modified RLSTM-CA or modified CA. Also, the present invention relates to cells including the modified RLSTM-CA or modified CA, plants containing the modified RLSTM-CA or modified RLSTM, and methods using compositions of the present invention. In addition, the present invention relates to antibodies conjugated to CA which may bind to the large or small subunits of Rubisco. The present invention provides methods of altering Rubisco carboxylase activity and altering plant growth.

BACKGROUND

Ribulose-1,5-bisphosphate carboxylase-oxygenase (Rubisco) catalyzes the reduction of atmospheric CO2 during photosynthesis. This process is not efficient and methods to improve the efficiency of CO2 reduction are needed. Methods which improve the efficiency of CO2 reduction will increase plant growth rates, which will be useful to a variety of industries.


Acad. Sci. USA 85:1513-1517 (1988)). The N-terminal Met-1 and Ser-2 are removed and Pro-3 acetylated. Additionally, the LS of Rubisco from tobacco, muskmelon, pea, and several other species is post-translationally modified by trimethylation of the e-amino of Lys-14 (Houtz et al., "Post-translational modifications in the amino-terminal region of the large subunit of ribulose-1,5-bisphosphate carboxylase/oxygenase from several plant species," Plant Physiol. 98:1170-1174 (1992); Houtz et al., "Post-translational modifications in the large subunit of ribulose bisphosphate carboxylase/oxygenase," Proc. Natl. Acad. Sci. USA 86:1855-1859 (1989)). The enzyme responsible for this latter modification is a highly specific chloroplast-localized S-adenosylmethionine (AdoMet)-protein (lys) "N-methyltransferase (protein methylase III, Rubisco LSMT, EC 2.1.1.43) (Houtz et al., "Post-translational modifications in the large subunit of ribulose-bisphosphate carboxylase/oxygenase," Proc. Natl. Acad. Sci. USA 86:1855-1859 (1989)). Rubisco LSMT has been affinity purified-8000-fold from pea chloroplasts and identified as a monomeric protein with a molecular mass of ~57 kDa (Wang et al., "Affinity Purification of Ribulose-1,5-bisphosphate Carboxylase/Oxygenase Large Subunit "N-Methyltransferase,"" Protein Expression and Purification 6:528-536 (1995). Recently, Rubisco LSMT cDNAs have been cloned and sequenced from pea and tobacco (Klein et al., "Cloning and developmental expression of pea ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit N-methyltransferase," Plant Molecular Biol. 27:249-261 (1995); Ying et al., "Organization and characterization of the ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit "N-methyltransferase gene in tobacco," Plant Molecular Biology 32:663-671 (1996)). The deduced amino acid sequences of tobacco Rubisco LSMT has 64.5% identity and 75.3% similarity with the sequence of pea Rubisco LSMT, and both proteins contain several copies of a conserved imperfect leucine-rich repeat motifs (Ying et al., "Organization and characterization of the ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit "N-methyltransferase gene in tobacco," Plant Molecular Biology 32:663-671 (1996)).

Kaplan et al., in U.S. Pat. No. 6,320,101, addressed the need for increased efficiency of Rubisco CO2 reduction by over-expressing a gene in a cell. However, the efficiency of CO2 reduction in this method is not optimal and there is a need for additional methods of altering Rubisco carboxylase activity. There is a need for improved methods of targeting molecules to Rubisco.

SUMMARY

The present invention relates to a modified Rubisco large subunit "N-Methyltransferase (RLSTM). The modified RLSTM may include a Rubisco large subunit or biologically active fragment thereof linked to a second molecule or biologically active fragment thereof. The modified RLSTM may have the ability to bind a Rubisco.

In a further embodiment, upon binding of the modified RLSTM to Rubisco, the concentration of the second molecule is increased at the active site of Rubisco. Upon binding of the modified RLSTM to Rubisco, the concentration of another molecule may be increased at the active site of Rubisco. The second molecule may be carbonic anhydrase. The second molecule may be a phosphatase. The second molecule may be a 3-phosphoglycerate kinase, glyceraldehyde-3-phosphate dehydrogenase, Rubisco Activase, Phosphoribulokinase, polylysine, or oxygenase, for example. The other molecule
may be CO₂. The second molecule may alter Rubisco activity. The another molecule may alter Rubisco activity.

The present invention relates to the polypeptide sequences of modified RLSMTs, the nucleic acid sequences encoding modified RLSMTs, expression vectors including the nucleic acid sequence encoding a modified RLSMT operably linked to an expression control sequence, host cells expressing one or more modified RLSMTs of the present invention, and plants and photosynthetic organisms containing host cells expressing one or more modified RLSMTs of the present invention. The growth of the plant containing the host cell may be altered.

Aspects of the present invention relate to methods of increasing the efficiency of CO₂ reduction in a cell. This increased efficiency of CO₂ reduction will result in increased plant productivity.

An embodiment of the present invention is a modified Rubisco large subunit "N-Methyltransferase-carbonic anhydrase (RLSMT-CA). The modified RLSMT-CA of the present invention may be linked to a carbonic anhydrase (CA). The modified RLSMT-CA may have the ability to bind a Rubisco. The carboxylase activity of Rubisco may be altered upon binding of Rubisco to the modified RLSMT-CA.

The present invention relates to the polypeptide sequences of modified RLSMT-CAs, the nucleic acid sequences encoding modified RLSMT-CAs, expression vectors including the nucleic acid sequence encoding a modified RLSMT-CA operably linked to an expression control sequence, host cells expressing one or more modified RLSMT-CAs of the present invention, and plants and photosynthetic organisms containing host cells expressing one or more modified RLSMT-CAs of the present invention. The growth of the plant containing the host cell may be altered.

The modified RLSMT-CAs of the present invention include Rubisco large subunit "N-Methyltransferase(s), which have been linked to CA by a chemical. The modified RLSMT-CAs of the present invention also include fusion proteins. The fusion proteins are encoded by a nucleic acid. The nucleic acid may include a first nucleic acid encoding a Rubisco large subunit "N-Methyltransferase or biologically active fragment thereof. The nucleic acid may further include a second nucleic acid encoding a carbonic anhydrase or biologically active fragment thereof. The order of these sequences may be reversed.

In a further embodiment, the modified RLSMT-CA may include an antibody or biologically active fragment thereof. The antibody may be linked to a CA or a biologically active fragment thereof.

The present invention further relates to an antibody which binds to a Rubisco large subunit "N-Methyltransferase. The antibody may be linked to or bind another molecule. The other molecule may be carbonic anhydrase. The Rubisco carboxylase activity may be altered upon binding of Rubisco to the RLSMT linked or bound to the antibody linked to or bound to the carbonic anhydrase.

The present invention further relates to a modified Rubisco subunit linked to a carbonic anhydrase (RSCA). The Rubisco subunit may be a Rubisco large subunit (LS) or a Rubisco small subunit (SS). The modified Rubisco subunit may assemble into Rubisco.

In a further embodiment, upon assembly of the modified LS or SS into Rubisco, the concentration of a first molecule is increased at the active site of Rubisco. Upon assembly of the modified LS and/or SS into Rubisco, the concentration of a second molecule may be increased at the active sites of Rubisco. The first molecule may be carbonic anhydrase. The second molecule may be CO₂. The first or second molecule may alter Rubisco activity.

The present invention relates to the polypeptide sequences of modified RSCA, the nucleic acid sequences encoding modified RSCA, expression vectors including the nucleic acid sequences encoding modified RSCA operably linked to an expression control sequence, host cells expressing one or more modified RSCAs of the present invention, and plants and photosynthetic organisms containing host cells expressing one or more modified RSCA of the present invention. The growth of the plant containing the host cell may be altered.

The present invention further relates to an antibody which binds to the LS or SS of Rubisco. The antibody may be linked to or bind another molecule. The other molecule may be carbonic anhydrase. The Rubisco carboxylase activity may be altered upon binding of the antibody linked to or bound to the carbonic anhydrase.

A method of altering Rubisco carboxylase activity is provided including providing a modified RLSMT-CA, modified RSCA, modified LSMT, a nucleic acid encoding a modified RLSMT-CA, modified RSCA, modified LSMT, or an antibody conjugated to CA.

A method of altering plant growth is provided including providing a modified RLSMT-CA, modified RSCA, modified LSMT, a nucleic acid encoding a modified RLSMT-CA, modified RSCA, modified LSMT, or an antibody conjugated to CA.

A particular embodiment is related to a method of delivering a molecule to the Rubisco active site. A Rubisco large subunit or small subunit is provided. A RLSMT is provided, which binds the Rubisco large or small subunit, wherein the Rubisco large and small subunits are integral components of Rubisco. The molecule to be delivered is bound to the RLSMT.

BRIEF DESCRIPTION OF DRAWINGS

The objects and advantages of the invention will be understood by reading the following detailed description in conjunction with the drawings in which:

FIGS. 1A-C illustrate an estimation of the binding affinity of wild-type pea Rubisco LSMT for PVDF-immobilized des (methyl) spinach Rubisco.

FIGS. 2A-C illustrate the fractionation of human carbonic anhydrase II (HCAII)-Rubisco LSMT conjugate, Rubisco, Rubisco LSMT, and HCAII by chromatography on Sepharose 6B.

FIGS. 3A-B illustrate the effect of HCAII conjugated to Rubisco LSMT on carboxylase activity of spinach and pea Rubisco.

FIG. 4 illustrates kinetic analyses of Rubisco, Rubisco complexed with hCA, Rubisco complexed with hCA in the presence of acetazolamide (Aza).

FIG. 5 illustrates the purification of RLSMT-CA fusion protein.

FIG. 6A is the nucleic acid sequence of the RLSMT-CA (SEQ ID No.: 1). FIG. 6B is the amino acid sequence of the RLSMT-CA fusion protein (SEQ ID No.: 2).

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Unless otherwise stated, the following terms used in the specification and claims have the meanings given below:

By "RLSMT" is meant a Rubisco large subunit "N-Methyltransferase."
By “RLSMT-CA” is meant a Rubisco large subunit “N-Methyltransferase-carbonic anhydrase.” By “RSCA” is meant a Rubisco subunit-carbonic anhydrase.

By “modified RLSMT-CA” is meant contains modifications of molecules not found in the wild-type RLSMT or wild-type CA.

By “Biologically active fragment” of a carbonic anhydrase is meant a polypeptide fragment of a carbonic anhydrase which has the ability to facilitate the equilibrium between dissolved HCO₃⁻ and CO₂.

By “Biologically active fragment” of Rubisco large subunit “N-Methyltransferase” is meant a polypeptide fragment of a Rubisco large subunit “N-Methyltransferase” which has the ability to bind to Rubisco.

Polymer (reacted with itself) is a repiledon, such as a plasmid, phage or other vector that may be attached to a second molecule or biologically active fragment thereof linked to a second molecule or biologically active fragment thereof. The modified RLSMT has the ability to bind to Rubisco. The modified RLSMT may result in an increase in the concentration of the second molecule near Rubisco. The second molecule may be carbolic anhydrase, a phosphatase, a 3-phosphoglycerate kinase, glyceraldehyde-3-phosphate dehydrogenase, Rubisco Activase, Phosphoribulokinase, polyisine, or oxygenase, for example. The increased concentration of the second molecule may increase the concentration of another molecule, such as a nucleotide or amine, near Rubisco. For example, the modified molecule may be an enzymatic product of the second molecule. For example, the modified molecule may be a molecule that binds the second molecule.

In a particular embodiment, the Rubisco large subunit “N-methyltransferase” is linked to the second molecule by a chemical. Representative chemicals useful for linking these molecules are well-known to one of skill in the art. In this example the linkage is through free amine groups using glutaraldehyde. The use of any reactive molecule capable of cross-linking two proteins is also contemplated in this embodiment. Non-limiting examples of such cross-linking reagents are APS—(p-Azidophenyl glyoxal monohydrate)—specific for arginine residues, BMHE—(Bismaleimidohexane)—specific for sulfhydryl groups, DSP—(Luman’s Reagent)—([Di-bis[succinimidylpropionate]])—specific for amines, EGS—([Ethylene glycol-bis[succinimidylsuccinate]])—specific for amines, EDC—([1-ethyl-3-[3(dimethylaminopropyl)]carboimide Hydrochloride])—specific for amines and carboxyls, MBS—([Maleimidobenzylation-Hydroxysuccinimide ester])—specific for sulfhydryls and amines, HPG—([p-Hydroxyphenylglyoxylate])—specific for arginine residues, SPDP—([N-Succinimidyl-3-[2-pyridyldithio]propionate])—reactive with amines and sulfhydryls, Sulfo-SAPB—([Sulfo-succinimidyl-4-[p-azidophenyl]butyrate])—amines and photoreactive, and BSOCOES—([Bis[2-(succinimidooxycarbonyloxyethyl)sulfonyl])—reactive with amines.

In a particular embodiment, the modified RLSMT is a fusion protein. The fusion protein may be encoded by a nucleic acid sequence, wherein the nucleic acid sequence comprises a first nucleic acid sequence encoding a Rubisco large subunit “N-methyltransferase” and/or a second nucleic acid sequence encoding a biologically active fragment thereof, and wherein the nucleic acid sequence further comprises a second nucleic acid sequence encoding a polypeptide. Methods of molecular biology useful in these methods are well-known to one of skill in the art. In a particular embodiment, the second nucleic acid encodes carbonic anhydrase. In a particular embodiment, the second nucleic acid encodes a phosphatase. The second nucleic acid may encode a 3-phosphoglycerate kinase, glyceraldehyde-3-phosphate dehydrogenase, Rubisco Activase, Phosphoribulokinase, polyisine, or oxygenase, for example. The order of these sequences may be reversed.

A particular embodiment of the present invention relates to a modified Rubisco large subunit “N-methyltransferase” (RLSMT). The modified RLSMT contains a Rubisco large subunit “N-methyltransferase” or biologically active fragment thereof linked to a second molecule or biologically active fragment thereof. The modified RLSMT has the ability to bind to Rubisco. The modified RLSMT may result in an increase in the concentration of the second molecule near Rubisco. The second molecule may be carbolic anhydrase, a phosphatase, a 3-phosphoglycerate kinase, glyceraldehyde-3-phosphate dehydrogenase, Rubisco Activase, Phosphoribulokinase, polyisine, or oxygenase, for example. The increased concentration of the second molecule may increase the concentration of another molecule, such as a nucleotide or amine, near Rubisco. For example, the modified molecule may be an enzymatic product of the second molecule. For example, the modified molecule may be a molecule that binds the second molecule.

By “vector” is meant a repiledon, such as a plasmid, phage or other vector that may be attached to a second molecule or biologically active fragment thereof linked to a second molecule or biologically active fragment thereof. The modified RLSMT has the ability to bind to Rubisco. The modified RLSMT may result in an increase in the concentration of the second molecule near Rubisco. The second molecule may be carbolic anhydrase, a phosphatase, a 3-phosphoglycerate kinase, glyceraldehyde-3-phosphate dehydrogenase, Rubisco Activase, Phosphoribulokinase, polyisine, or oxygenase, for example. The increased concentration of the second molecule may increase the concentration of another molecule, such as a nucleotide or amine, near Rubisco. For example, the modified molecule may be an enzymatic product of the second molecule. For example, the modified molecule may be a molecule that binds the second molecule.

The present invention relates to a delivery compound to the active-site vicinity of Rubisco. The presence and/or increased concentration of compounds and enzymes near the active-site vicinity of Rubisco may result in an increased in the carboxylase activity or specificity factor of Rubisco, for example. Increased carboxylase activity and specificity factor result in increased biomass production and thus translate into increased food and fiber production.

As described herein, localization of CA to Rubisco substantially increases the efficiency of CO₂ fixation by Rubisco at low CO₂. The delivery of enzymes such as Rubisco Activase may increase the efficiency of Rubisco Activase activity, an enzyme known to be essential for Rubisco activity. Also, delivery of phosphatases specific for degradation of carboxyarabinitol-1-phosphate would also be useful since this compound is a known natural inhibitor of Rubisco activity that occurs in many plants. Furthermore, the rapid metabolism of phosphoglyceric acid, the immediate product of the carboxylase reaction, or 2-phosphoglycolate the immediate product of the oxygenase reaction, by targeting enzymes capable of metabolizing these products may also facilitate higher carboxylase activity or increases in the specificity factor of Rubisco. Other polypeptides such as polyclins and other amino acid polymers may also be beneficial by delivering additional carbon dioxide to Rubisco sequestered as carbonates on the amino acid amine groups.

The interaction between pea Rubisco LSMT and des(methyl) forms of Rubisco is exceptionally tight and specific (see FIG. 1). The interaction in general at least involves regions from the carboxy-terminal region of the LS of Rubisco and the amino-terminal region of pea Rubisco LSMT as measured by the susceptibility of Cys-119 and Cys-188 in pea Rubisco LSMT and Cys-459 in the LS of spinach Rubisco to cross-linking by 1,6-Bismaleimidohexane (BMHE, a homobifunctional sulfhydryl-specific cross-linking reagent). The interaction between pea Rubisco LSMT and des(methyl)spinach Rubisco appears to be stoichiometric with 1:4 monomers of pea Rubisco LSMT binding to each Rubisco holoenzyme. However, the binding has synergistic characteristics with decreasing binding affinity with increasing number of bound pea Rubisco LSMT monomers. Other regions of Rubisco LS and/or Rubisco SS may also be involved in Rubisco LSMT binding.

Given the tight binding of Rubisco LSMT to Rubisco, the present invention utilizes the Rubisco LSMT to deliver molecules to the active-site vicinity of Rubisco. A number of molecules may be delivered to Rubisco in this fashion. This provides utility for increasing the local concentration of a molecule at the active-site vicinity of Rubisco. Modified Rubisco Large Subunit “N-methyltransferase”
In a particular embodiment, the Rubisco large subunit is linked to the second molecule by a chemical. Representative chemicals useful for linking these molecules are well-known to one of skill in the art.

In a particular embodiment, the modified Rubisco large subunit is a fusion protein. The fusion protein may be encoded by a nucleic acid sequence, wherein the nucleic acid sequence comprises a first nucleic acid sequence encoding a Rubisco large subunit or biologically active fragment thereof, and wherein the nucleic acid sequence further comprises a second nucleic acid sequence encoding a polypeptide. Methods of molecular biology useful in these methods are well-known to one of skill in the art. In a particular embodiment, the second nucleic acid sequence encodes carbonic anhydrase. The order of these sequences may be reversed.

A particular embodiment of the present invention relates to a modified Rubisco small subunit. The modified RSS contains a Rubisco small subunit or biologically active fragment thereof linked to a first molecule or biologically active fragment thereof.

In a particular embodiment, the Rubisco small subunit is linked to the second molecule by a chemical. Representative chemicals useful for linking these molecules are well-known to one of skill in the art.

In a particular embodiment, the modified Rubisco small subunit is a fusion protein. The fusion protein may be encoded by a nucleic acid sequence, wherein the nucleic acid sequence comprises a first nucleic acid sequence encoding a Rubisco small subunit or biologically active fragment thereof, and wherein the nucleic acid sequence further comprises a second nucleic acid sequence encoding a polypeptide. Methods of molecular biology useful in these methods are well-known to one of skill in the art. In a particular embodiment, the second nucleic acid sequence encodes carbonic anhydrase. The order of these sequences may be reversed.

A particular embodiment of the present invention relates to a modified carbonic anhydrase. The modified carbonic anhydrase contains a carbonic anhydrase or biologically active fragment thereof linked to a first molecule or biologically active fragment thereof.

In a particular embodiment, the carbonic anhydrase is linked to the second molecule by a chemical. Representative chemicals useful for linking these molecules are well-known to one of skill in the art.

In a particular embodiment, the modified carbonic anhydrase is a fusion protein. The fusion protein may be encoded by a nucleic acid sequence, wherein the nucleic acid sequence comprises a first nucleic acid sequence encoding a carbonic anhydrase or biologically active fragment thereof, and wherein the nucleic acid sequence further comprises a second nucleic acid sequence encoding a polypeptide. Methods of molecular biology useful in these methods are well-known to one of skill in the art. The order of these sequences may be reversed. Preferably, the second molecule is RLSMT, Rubisco LS, Rubisco SS, or a protein that binds either Rubisco LS or Rubisco SS. The second molecule may be Rubisco LSMT. Most preferably, the carboxylase activity of Rubisco is altered upon binding of the modified carbonic anhydrase to Rubisco LS or Rubisco SS.


Rubisco large subunit N-methyltransferases useful in the present invention may be derived from any plant or photosynthetic organism.

Upon binding of the modified RLSMT to Rubisco, the concentration of the first molecule may be increased at the active site of Rubisco. The first molecule may be carbonic anhydrase or biologically active fragments thereof. The first molecule may alter Rubisco activity.

Upon binding of the modified RLSMT to Rubisco, the concentration of a second molecule may be increased at the active site of Rubisco. The second molecule may be the product of a reaction catalyzed by the first molecule. The second molecule may be CO2. The second molecule may alter Rubisco activity.

In a particular embodiment, the modified Rubisco large subunit N-methyltransferase is in a complex with carbonic anhydrase. The modified Rubisco large subunit N-methyltransferase may be linked to carbonic anhydrase or a biologically active fragment thereof. The modified RLSMT-CA has the ability to bind Rubisco. Rubisco activity is altered when bound to the RLSMT-CA complex.

In a particular embodiment, the Rubisco large subunit N-methyltransferase is linked to the carbonic anhydrase by a chemical. Representative chemicals useful for linking these molecules are well-known to one of skill in the art.

In a particular embodiment, the modified RLSMT-CA is a fusion protein. The fusion protein is encoded by a nucleic acid sequence, wherein the nucleic acid sequence includes a first nucleic acid sequence encoding a Rubisco large subunit N-methyltransferase or biologically active fragment thereof, and wherein the nucleic acid sequence further includes a second nucleic acid sequence encoding a carbonic anhydrase or biologically active fragment thereof. The order of these sequences may be reversed.

In a particular embodiment, the Rubisco complex with carbonic anhydrase includes an antibody. The antibody may be linked to a carbonic anhydrase. Preferably, the antibody may bind to Rubisco LS or Rubisco SS.

In a particular embodiment, the Rubisco large subunit N-methyltransferase-CA complex is within a cell. The cell may be prokaryotic or eukaryotic. The cell may be a plant cell or plant tissue.

In a particular embodiment, the Rubisco large subunit N-methyltransferase-CA complex is in a photosynthetic organism.

Rubisco Large Subunit N-methyltransferase Carbonic Anhydrase

An aspect of the present invention addresses the need to increase the rate of CO2 fixation by Rubisco, which is an inefficient process in wild-type plants. An increased rate of CO2 fixation by Rubisco will increase plant productivity, which is useful in terms of food, fiber, and biomass and is applicable to various aspects of agriculture.
At atmospheric levels of CO₂, the carboxylase activity of Rubisco is limited. Additional limitations occur as a consequence of competitive inhibition by O₂. However, many plants possess mechanisms to alleviate this constraint, and increase the rate of CO₂ fixation by compartmentalization of Rubisco in conjunction with accumulation of inorganic carbon or CO₂. An important part of this mechanism is carbonic anhydrase that facilitates equilibrium between dissolved HCO₃⁻ and CO₂. Given the tight and specific binding of Rubisco LSMT to (de)s(ethyl) forms of Rubisco, and the absence of any negative consequences from this binding on the catalytic activity of Rubisco, studies were undertaken to target carbonic anhydrase (CA) to Rubisco utilizing Rubisco LSMT. The results from these experiments demonstrate that localization of CA to Rubisco substantially increases the efficiency of CO₂ fixation by Rubisco at low CO₂.

A particular embodiment of the present invention relates to a modified Rubisco large subunit "N-methyltransferase-carboxic anhydrase (RLSMT-CA) complex, wherein the Rubisco large subunit "N-methyltransferase is linked to carbonic anhydrase. This RLSMT-CA complex has the ability to bind to Rubisco, and Rubisco carboxylase activity is altered when bound to the RLSMT-CA complex.

The reduction in the apparent affinity for CO₂ by the direct placement of CA to the active-site vicinity of Rubisco circumvents the kinetic constraints and enables Rubisco to maintain nearly saturated rates of carboxylase activity at air levels of CO₂.

Methods of the Present Invention

A method of altering Rubisco carboxylase activity is provided including modifying CA by providing a modified RLSMT-CA, a nucleic acid sequence encoding a RLSMT-CA, or an antibody conjugated to CA.

A method of altering plant growth is provided including modifying the RLSMT-CA by providing a modified RLSMT-CA, a nucleic acid sequence encoding a RLSMT-CA, or an antibody conjugated to CA.

A particular embodiment is related to a method of delivering a molecule to the vicinity of the Rubisco active site. A Rubisco large or small subunit is provided. A RLSMT is provided, which binds the Rubisco large or small subunit. The molecule to be delivered is bound to the RLSMT.

Antibody

Also, antibodies including both polyclonal and monoclonal antibodies, and drugs that modulate the production or activity of the modified RLSMT-CAs, modified RLSMTs, Rubisco LSs, Rubisco SSs, and/or their biologically active fragments or subunits may be useful in the production or modification of modified RLSMT-CAs, modified RLSMTs, Rubisco LSs, or Rubisco SSs. Antibodies may be conjugated to another molecule, such as a carbonic anhydrase. For example, RLSMT-CAs, modified RLSMTs, Rubisco LSs, Rubisco SSs, or fragments or subunits thereof may be used to produce both polyclonal and monoclonal antibodies, to the RLSMT-CAs, modified RLSMTs, Rubisco LSs, Rubisco SSs, or fragments or subunits thereof, in a variety of cellular media, by known techniques such as the hybridoma technique utilizing, for example, fused mouse spleen lymphocytes and myeloma cells. Likewise, small molecules that mimic or antagonize the activity of the RLSMT-CAs, modified RLSMTs, Rubisco LSs, or Rubisco SSs, of the invention may be discovered or synthesized, and may be used in diagnostic and/or therapeutic protocols.

The present invention also extends to the development of antibodies against the modified RLSMT-CAs, modified RLSMTs, Rubisco LSs, Rubisco SSs, including naturally raised and recombinantly prepared antibodies. For example, the antibodies could be used to modulate Rubisco activity, or in the production or modulation of a modified RLSMT-CA or modified RLSMT.

The general methodology for making monoclonal antibodies by hybridomas is well known. Methods for producing monoclonal anti-modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco antibodies are also well-known in the art. See Nimam et al., Proc. Natl. Acad. Sci. USA, 80:4949-4953 (1983). Typically, the modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS or a peptide analog is used either alone or conjugated to an immunogenic carrier, as the immunogen in the before described procedure for producing anti-modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco monoclonal antibodies. The culture is maintained under conditions and for a time period sufficient for the hybridoma to secrete the antibodies into the medium. The antibodies are then purified for the ability to produce an antibody that immunoreacts with the modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, Rubisco or peptide analog. The antibody-containing medium is then collected. The antibody can then be further isolated by well-known techniques. Immortal, antibody-producing cell lines can also be created by techniques other than fusion, such as direct transformation of B lymphocytes with oncogenic DNA, or transfection with Epstein-Barr virus. See, e.g., Using Antibodies: A Laboratory Manual, Harlow, Ed and Lane, David (Cold Spring Harbor Press, 1999).

Media useful for the preparation of these compositions are both well-known in the art and commercially available and include synthetic culture media, inbred mice and the like. An exemplary synthetic medium is Dulbecco's minimal essential medium (DMEM; Dulbecco et al., Virol. 8:396 (1959)) supplemented with 4.5 g/ml glucose, 20 mm glutamine, and 20% fetal calf serum. A preferred inbred mouse strain is the Balb/c.

Methods for producing polyclonal anti-polypeptide antibodies are well-known in the art. See U.S. Pat. No. 4,493,795 to Nestor et al. A monoclonal antibody, and immunologically active fragments thereof, can be prepared using the hybridoma technology described in e.g., Using Antibodies: A Laboratory Manual, Harlow, Ed and Lane, David (Cold Spring Harbor Press, 1999), which is incorporated herein by reference. Briefly, to form the hybridoma from which the monoclonal antibody composition is produced, a myeloma or other self-perpetuating cell line is fused with lymphocytes obtained from the spleen of a mammal hyperimmunized with a modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco. Splenocytes are typically fused with myeloma cells using polyethylene glycol (PEG) 6000 MW. Fused hybridomas are selected by their sensitivity to HAT (hypoxanthine, aminopterin, thymidine) supplemented media. Hybridomas producing a monoclonal antibody useful in practicing this invention are identified by their ability to immunoreact with the present modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco and their ability to alter specified modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco activity in target cells.

Panels of monoclonal antibodies produced against modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco peptides can be screened for various properties, i.e., isotype, epitope, affinity, and the like. High affinity antibodies are also useful when immunoaffinity purification of native or recombinant modified RLSMT-CA, modified RLSMT, Rubisco LS, Rubisco SS, or Rubisco is possible.
Cells, Tissues, and Substrates
The cells of the present invention may express at least one modified Rubisco Large Subunit *N*-methyltransferase. The cells may further comprise a conjugate of a carbohydrate anhydride and Rubisco Large Subunit *N*-methyltransferase. Useful cells include enkaryotic and prokaryotic cells, including, but not limited to, bacterial cells, yeast cells, fungal cells, insect cells, nematode cells, plant cells, and animal cells. Suitable animal cells include, but are not limited to, HEK-293 cells, U2OS cells, HeLa cells, COS cells, and various primary mammalian cells.

In a preferred embodiment of the present invention, the photosynthetic organism is a C3 plant. Non-limiting examples of C3 plants include spinach, wheat, rice, and barley.

The cells of the present invention may express one modified Rubisco large subunit *N*-methyltransferase that results in enhanced carbon fixation upon expression in a plant.

A substrate may have deposited thereon a plurality of cells of the present invention. The substrate may be any suitable biologically substrate, including but not limited to, glass, plastic, ceramic, semiconductor, silica, fiber optic, diamond, biocompatible monomer, or biocompatible polymer materials.

Nucleic Acids Encoding Modified Rubisco
The present invention further includes nucleic acid molecules that encode modified Rubisco large subunit *N*-methyltransferase. It should be appreciated that also within the scope of the present invention are DNA sequences encoding modified Rubisco large subunit *N*-methyltransferase.

The modified Rubisco large subunit *N*-methyltransferase may be generated by molecular biological techniques standard in the genetic engineering art, including but not limited to, polymerase chain reaction (PCR), restriction enzymes, expression vectors, plasmids, and the like. By way of example, vectors may be designed which contain a signal sequence. PCR amplified DNA fragments of a modified Rubisco large subunit *N*-methyltransferase to be modified may be digested by appropriate restriction enzymes and sub-cloned into the vector. Modifications of the DNA may be introduced by standard molecular biological techniques as described above.

Transformation
In order to prepare the integration of foreign genes into higher plants a high number of cloning vectors are available, containing a replication signal for *E. coli* and a marker gene for the selection of transformed bacterial cells. Examples for such vectors include, but are not limited to, pBR322, pUC series, M13 mp series, pACYC184. The desired sequence may be integrated into the vector at a suitable restriction site. The obtained plasmid is used for the transformation of *E. coli* cells. Transformed *E. coli* cells are cultivated in a suitable medium and subsequently harvested and lysed. The plasmid is recovered. Restriction analysis, gel electrophoresis and other biochemical-molecular biological methods may be used to analyze the obtained plasmid DNA. After each manipulation the plasmid DNA may be cleaved and the obtained DNA fragments may be linked to other DNA sequences. Each plasmid DNA may be cloned into the same or into other plasmids.

In order to integrate DNA into plant host cells a wide range of techniques are available. These techniques comprise the transformation of plant cells with T-DNA by using *Agrobacterium tumefaciens* or *Agrobacterium rhizogenes* as transformation medium, the fusion of protoplasts, the injection and the electroporation of DNA, the integration of DNA by means of the biolistic method as well as further possibilities.

Several modes of delivery of the DNA are known to those of skill in the art. For example, DNA may be delivered into the nuclear genome, or the chloroplast genome. If delivered into the nuclear genome, the construct will have to an N-terminal sequence included in the construct capable of targeting the fusion to the chloroplast. This technique well known to one of skill in the art, may use the transit sequence for the SS of Rubisco, which is known to be chloroplast-localized, but is a nuclear-encoded gene. Alternatively, recent advances also allow for delivery of the fusion construct directly into the chloroplast genome, which obviates the necessity for a chloroplast targeting sequence. These methods are described, for example, in Karlin, Neumann, G. A. and Tobin, E. M., EMBO J. 5 (1986) 9-13; Liu, Y. Y., Kaderbhai, N., and Kaderbhai, M. A., Biochem. J. 351 Pt 2 (10-15-2000) 377-384; Mishkind, M. L., Wessler, S. R., and Schmidt, G. W., J. Cell Biol. 100 (1985) 226-234; Huang, F. C., Klaus, S. M., Herz, S., Zou, Z., Koop, H. U., and Golds, T. J., Mol. Genet. Genomics 268 (2002) 19-27; Madoka, Y., Tomizawa, K., Mizoi, J., Nishida, L., Nagano, Y., and Sasaki, Y., Plant Cell Physiol 43 (2002) 1518-1525.

In the case of injection and electroporation of DNA into plant cells, plasmids such as pUC derivatives may be used. However, in cases in which whole plants are to be regenerated from transformed cells, a selectable marker gene should be present.

Depending on the method used for integrating desired genes into the plant cell, further DNA sequences such as Ti-and Ri-plasmid T-DNA may be necessary.

If *Agrobacteria* are used for the transformation, the DNA which is to be integrated must be cloned into special plasmids, namely either into an intermediate vector or into a binary vector. Due to sequences homologous to the sequences within the T-DNA, the intermediate vectors may be integrated into the Ti- or Ri-plasmid of the *Agrobacterium* due to homologous recombination. This also contains the vir-region necessary for the transfer of the T-DNA. Intermediate vectors cannot replicate in *Agrobacteria*. By means of a helper plasmid the intermediate vector may be transferred to *Agrobacterium tumefaciens* (conjugation). Binary vectors may replicate in *E. coli* as well as in *Agrobacteria*. They contain a selectable marker gene as well as a linker or polylinker which is framed by the right and the left T-DNA border region. They may be transformed directly into the *Agrobacterium* (Holsters et al. Mol. Genet. Genet. 163 (1978), 181-187). The *Agrobacterium* acting as host cell should contain a plasmid carrying a vir-region. The vir-region is necessary for the transfer of the T-DNA into the plant cell. Additional T-DNA may be present. The *Agrobacterium* transformed in such a way is used for the transformation of plant cells.


For transferring the DNA into the plant cells, plant explants may suitably be co-cultivated with *Agrobacterium tumefaciens* or *Agrobacterium rhizogenes*. From the infected plant material (e.g. pieces of leaves, stem segments, roots, but also protoplasts or suspension-cultivated plant cells) whole plants may then be regenerated in a suitable medium which may contain antibiotics or bioxides for the selection of transformed cells. The plants obtained in such a way may then be examined as to whether the integrated DNA is present or not. Other possibilities in order to integrate foreign DNA by using the biolistic method or by transforming protoplasts are known.
to the skilled person (cf., e.g., Willmitzer, L., 1993 Transgenic plants. In: Biotechnology, A Multi-Volume Comprehensive Treatise. H. J. Rehm, G. Reed, A. Puhler, P. Stadler, editors), Vol. 2, 627-659, VCH Weinheim-New York-Basel-Cambridge). Once the introduced DNA has been integrated in the genome of the plant cell, it usually continues to be stable and also remains within the descendants of the originally transformed cell. It usually contains a selectable marker which confers resistance against a biocide or against an antibiotic such as kanamycin, G 418, bleomycin, hygromycin or phosphinotricine, etc. to the transformed plant cells. The individually selected marker should therefore allow for a selection of transformed cells to cells lacking the integrated DNA. The transformed cells grow in the usual way within the plants (see also McCormick et al., 1986, Plant Cell Reports 5: 81-84).

The resulting plants can be cultivated in the usual way and cross-bred with plants having the same transformed genetic heritage or another genetic heritage. The resulting hybrid plants will have the corresponding phenotypic properties. Two or more generations should be grown in order to ensure whether the phenotypic feature is kept stably and whether it is transferred. Furthermore, seeds should be harvested in order to ensure that the corresponding phenotype or other properties will remain.

Although the present invention has been described in detail with reference to examples above, it is understood that various modifications can be made without departing from the spirit of the invention, and would be readily known to the skilled artisan. Additionally, the invention is not to be construed to be limited by the following examples.

EXAMPLES

Example 1

Materials and Methods


Example 2

Binding of Rubisco LSMT and des(methyl) Forms of Rubisco

Purified Rubisco LSMT and hCAII (1:8 M ratio) reacted in buffer A with 0.5% glutaraldehyde for 2 hours at room temperature (approx. 22° C.). The glutaraldehyde was quenched by the addition of Tris pH 0.4 M, then dialyzed exhaustively in the microdializer (MWCO 8000) against buffer B at 4° C. Conjugated protein was then incubated in buffer B with purified spinach Rubisco (1 Rubisco:2 conjugate molar ratio) for 4 hours on ice. The multi-enzyme complex (conjugate) was subjected to Sepharose CL 6B gel permeation chromatography (1x45 cm column) and resolved at 0.1 mL/min into 0.5 mL fractions. The Sepharose CL 6B column was calibrated according to manufacturer’s instructions (FIG. 4). Fractions were analyzed for Rubisco LSMT, Rubisco, and hCAII activities as described above.

A human form of carbonic anhydrase (hCAII) was chemically cross-linked with Rubisco large subunit (LS) N-Methyltransferase (Rubisco LSMT). The complex between hCAII and Rubisco LSMT retained the highly specific and tight-binding affinity (KD=0.1 nM) for des(methyl) forms of Rubisco characteristic of Rubisco LSMT. Binding of this complex to Rubisco, and localization of carbonic anhydrase to the active-site vicinity of Rubisco, resulted in a significant increase in the carboxylation efficiency of Rubisco manifested as a decrease in the K<sub>m</sub> (CO₂) from 13.4 μM to 3.44 μM (FIG. 4). Since the level of dissolved CO₂ in equilibrium with air (~360 ppm) is approximately 15 μM, it has been widely recognized for decades that a major limitation to plant productivity and biomass production is the kinetic constraint for CO₂ fixation by Rubisco. At air levels of CO₂, the enzymatic velocity for carboxylation by Rubisco is only half of the potential velocity, and additionally is further reduced by competitive inhibition by oxygen. These two factors reduce the carboxylation efficiency of Rubisco, to the point that nearly all crop plants show a remarkable stimulation of growth and productivity when cultured under atmospheres enriched in CO₂. The reduction in the apparent affinity for CO₂ by the direct placement of CA to the active-site vicinity of Rubisco circumvents these kinetic constraints and enables Rubisco, to maintain nearly saturated rates of carboxylation at air levels of CO₂ (see FIG. 3, panel A and inset). It is contemplated in the present invention that these results can be duplicated using gene fusion techniques, and introduced into transgenic plants. The potential exists to increase plant productivity by 50-100% in all C3 plants (a form of biochemical plant classification based on photosynthesis that represents the majority of crop plants).

FIG. 1. Estimation of the binding affinity of wild-type pea Rubisco LSMT for PVDF-immobilized des(methyl) spinach Rubisco. FIG. 1 A shows the loss in Rubisco LSMT activity at 4° C. from pea chloroplast lysates over time during incubation with PVDF-immobilized Rubisco. The concentration of Rubisco LSMT was estimated at 58 m from activity measurements assuming a specific activity of purified pea Rubisco LSMT of 306 nmoles/min/mg protein. FIG. 1B shows the loss in bound Rubisco LSMT activity at 4° C. from PVDF-immobilized spinach Rubisco over time as determined by activity measurements of bound Rubisco LSMT. Immunological analyses confirmed that the loss in Rubisco LSMT activity was due to disassociation and not catalytic inactivation. FIG. 1C shows an affinity plot of data from FIG. 1A and FIG. 1B estimating the affinity constant KD for the binding of pea Rubisco LSMT to spinach Rubisco.

Example 3

Purification of Rubisco LSMT and des(methyl) Forms of Rubisco

Several studies have suggested that the activity of CA could be essential for optimum rates of photosynthetic CO₂ fixation in C3 plants at low levels of CO₂. Ample evidence exists for an essential role of CA in the CCM (carbon concentrating mechanism) in C4 plants, and aquatic species with CCMs. Other studies have identified an association of CA with Calvin cycle enzymes and photosystem II complexes, as
as more defined localizations to pyrenoids and carboxysomes. However, direct experimental evidence of whether or not a close association of CA with Rubisco results in increased rates of CO$_2$ fixation is unavailable. The possibility that CA placed in direct proximity to Rubisco influences the rate or extent of CO$_2$ fixation at low CO$_2$ by creating catalytically and binding competent conjugates between HCAII (human carbon anhydrase II) and pea Rubisco LSMT was explored in the present invention. This complex has a relatively large molecular mass, binds to des(methyl) spinach Rubisco, retains CA, Rubisco LSMT, and Rubisco activity, and is immunologically reactive with Rubisco LSMT antibody (Figs. 2A, B, and D). The complex does not bind to pea Rubisco (Fig. 2A).

FIG. 2. Fractionation of human carbonic anhydrase II (HCAII)-Rubisco LSMT conjugate, Rubisco, Rubisco LSMT, and HCAII by chromatography on Sepharose 6B. Human carbonic anhydrase II enzyme (HCAII), kindly provided by P. J. Lapins, was induced according to Tanhauser, et al., Gene 117: 113-117 (1992), purified according to Whitley, et al., Anal. Biochem. 57: 467-476 (1974), and assayed according to Cronk, et al., Protein Sci. 10: 911-922 (2001) (2.22 μmoles/min/mg protein). 3 Wilbur-Anderson (W-A) units/μg. Spinach and pea Rubisco, and pea Rubisco LSMT were purified according to McCurry, et al., Methods Enzymol. 90 Pt E: 515-521 (1982) and Zheng, et al., Protein Expr. Purif. 14:104-112 (1998) respectively. Pea Rubisco LSMT and HCAII were conjugated at a 1:8 molar ratio (240 μM:1.92 mM) in 0.1 M potassium phosphate buffer, pH 6.8, by incubation for 2 hours at 25°C with (0.5% v/v) glutaraldehyde. Excess glutaraldehyde was quenched by the addition of Tris buffer, pH 8.2, to 0.4 M and the reaction diazoyzed exhaustively against 50 mM Bicine, pH 8.2. The conjugated reaction products were incubated with 10 μg of per (A) or spinach (B) Rubisco for 2 hours at 25°C. The solution (0.5 mL) was applied to a Sepharose 6B gel permeation column (1.5 x 45 cm) and eluted at 0.1 mL/min in 50 mM Bicine pH 8.2. Protein concentration was determined by the Bradford method. Pooled fractions were enzymatically assayed for HCAII according to Cronk, et al., Protein Sci. 10: 911-922 (2001) (A: pool 1, 1.69 W-A; B: pool 1, 1.46 W-A; B: pool 2, 1.73 W-A); pea Rubisco LSMT (A: pool 1, 26600 μmoles/min/mg protein; B: pool 1, 26600 μmoles/min/mg protein) and Rubisco carboxylase (A: pea pool 2, 0.3 μmoles/min/mg protein; B: spinach pool 1, 2.2 μmoles/min/mg protein). C, Coomassie-blue stained SDS-PAGE (15%; upper panel) and pea Rubisco LSMT immunoanalysis (lower panel) of pooled fractions and enzymes described in B. A duplicate gel as in the upper panel was electroblotted to a PVDF membrane, and probed with protein A-purified anti-pea Rubisco LSMT antibody (1:10000). The pea Rubisco LSMT:HCAII conjugate was of such high molecular mass that the immunoreactive band represents a complex that remained stacked just above the resolving gel such that a Coomassie-blue stained band is not visible in the upper panel.

Example 4

Effect of HCAII Conjugated to Rubisco LSMT on Carboxylase Activity of Spinach and Pea Rubisco

Kinetic analyses were performed as described previously for Rubisco Lorimer, G. H., Badger, M. R., and Andrews, T. J. Biochemistry 15, 529-536.1976. Pierce, J. W., McCurry, S. D., Mulligan, R. M., and Tolbert, N. E. Methods Enzymol. 89 Pt D, 47-55.1982. Assays were conducted under CO$_2$ free conditions with CO$_2$-free buffers as described below, and controlled normalized Rubisco and hCAH activities, unless otherwise indicated. The assays contained, unless otherwise indicated, 100 mM Bicine adjusted to pH 8.2 with a saturated NaOH solution, 20 mM MgCl$_2$, 2 mM dithiothreitol, 0.5 mM RuBP, and NaH$_2$CO$_3$ (1.0 mM/mmol). Reactions were initiated by the addition of activated Rubisco, or conjugate, injected into an 8 mL serum capped vial (first flushed with nitrogen). Activation of Rubisco or conjugate was performed at 2 mg/mL by incubation at 30°C for 15 min in 50 mM Bicine, pH 8.2, containing 20 mM MgCl$_2$ and 15 mM NaH$_2$CO$_3$. Assay times, enzyme masses, volumes, or specific activities were adjusted to limit substrate consumption to 10% or less. The assays were terminated by the addition of 2N HCl (0.4 volumes), and the liquid evaporated to dryness under a stream of air at 50°C. Radioactivity was determined with a liquid scintillation analyzer (Packard 2000 CA), after addition of 0.2 mL H$_2$O and 4.5 mL of counting solvent (Bio-Safe II, RPI). Enzyme activity was plotted versus substrate concentration and the data fitted to the Michaelis-Menten equation using Sigma Plot version 8.0.

Under anaerobic conditions and utilizing CO$_2$-free buffers, the carboxylase activity of spinach Rubisco (6 μg, 15 μM CO$_2$, 1 mM HCO$_3$-, pH 8.2, 0.5 mM RuBP) remains linear until the ~4 μmoles of free CO$_2$ (250 μL assay volume) is consumed yielding a velocity of 0.83 μmoles/min/mg protein (1/5 V$_{max}$, 2.2 μmoles/min/mg protein), and then ceases as CO$_2$ is totally depleted (Fig. 3A). In contrast an equal amount of spinach Rubisco (6 μg) with 0.89 W-A (Wilbur-Anderson units) of CA bound (equivalent to ~300 μg of native unconjugated CA) under identical conditions exhibits linear CO$_2$ fixation for 2 min, at a velocity of 2.08 μmoles/min/mg protein. Addition of an equal amount of CA (300 ng) to control assays has no effect on the profile of CO$_2$ fixation (Fig. 3A). Increasing amounts of free CA to 10-fold more than bound (~3 μg, ~10-fold molar excess over Rubisco) resulted in linear CO$_2$ fixation rates over the entire 2 min assay, at a velocity of 0.83 μmoles/min/mg protein. However, a 100-fold increase (also an ~100-fold molar excess over Rubisco) in the amount of free CA resulted in CO$_2$ fixation rates equal to that observed with 300 ng of bound CA. These differences are not observed at saturating levels of CO$_2$, and CA inhibitors abolished the effect of CA on carboxylase activity (Fig. 3C). Carboxylase assays at saturating and low CO$_2$ showed no difference in activity between native Rubisco and Rubisco with bound pea Rubisco LSMT alone. Thus, the effect of CA on the carboxylase activity of Rubisco appears to be a consequence of localization through pea Rubisco LSMT binding to Rubisco. An ideal control to substantiate this observation would be an identical conjugate with CA activity that did not bind to Rubisco. The conjugate between pea Rubisco LSMT and CA retains Rubisco LSMT activity, and catalytic methylation of des(methyl) spinach Rubisco by Rubisco LSMT results in release of Rubisco LSMT. We therefore added 100 μM AdoMet to the Rubisco complex containing bound CA prior to the initiation of carboxylase assays, and the effect of bound CA was abolished (Fig. 3C). Furthermore, Rubisco because it contains a trimethyllysyl residue at position 14 in the LS, and similar experiments with pea Rubisco demonstrated that the pea Rubisco LSMT CA conjugate did not bind to pea Rubisco (Fig. 2A), and no stimulation of carboxylase activity at low CO$_2$ was observed with pea Rubisco (Fig. 2B). The decline in Rubisco carboxylase activity after 1 min was not due to decarboxylaition and inactivation of Rubisco, because addition of saturating levels of CO$_2$ after 2 min (15 mM HCO$_3$-) to control assays resulted in velocities of 2.1 μmoles/min/mg protein.
FIG. 3. Assays were conducted under O2 free conditions with CO2-free buffers according to Lorimer, et al., Biochemistry 15: 529-536 (1976), altered as described below, and contained equal Rubisco activity and HCAII activity, unless otherwise indicated. The assays contained 100 mM Bicine adjusted to pH 8.2 with a saturated NaOH solution, 20 mM MgCl2, 2 mM dithiothreitol, 0.5 mM RuBP, and 15 mM NaH13CO3 (499-573 dpm/n mole). Reactions were initiated by the addition of 6 μg of activated Rubisco, (activated at 2 mg/mL by incubation at 30°C for 15 min in 50 mM Bicine, pH 8.2, containing 20 mM MgCl2 and 15 mM NaH13CO3, (499-573 dpm/Nmole), injected into an 8 mL serum-capped vial (first flushed with nitrogen). The assays were conducted in a volume of 0.25 mL, terminated by the addition of 0.1 mL of 2 N HCl, and the liquid evaporated to dryness under a stream of air at 50°C. Radioactivity was determined with a liquid scintillation analyzer (Packard 2000 CA), after the addition of 0.5 mL H2O and 4.5 mL of scintillation cocktail (Bio-Safe II, RPI). A. Acid-stable incorporation of 13CO2 with time with spinach Rubisco (6 μg; Vmax=2.2 μmoles CO2/min/mg protein) with and without pea Rubisco LSMT: HCAII conjugate (6 μg Rubisco, 0.9 W-A, equivalent to 0.3 μg HCAII) or 0.3 (1x), 3.0 (10x), and 30 μg (100x) of free HCAII. Inset, Spinach Rubisco with pea Rubisco LSMTP: HCAII conjugate preincubated with AdoMeth (100 μM) or HCA II inhibitors (1.5 mM ampicillin and 0.5 mM acetazolamide). B. Same condition as A but with pea Rubisco (46 μg, Vmax=0.3 μmoles CO2/min/mg protein) with and without pea Rubisco LSMTP: HCAII conjugate (0.9 W-A, equivalent to 0.3 μg HCAII) or 0.3 μg HCAII (1x).

FIG. 4 illustrates the kinetic analyses of Rubisco, Rubisco complexed with hCA, and Rubisco complexed with hCA in the presence of acetazolamide (Az). A mathematical consideration of the potential proximity effect of CA in these assays suggests that a maximum of ~41-fold could be observed in terms of a decrease in the diffusion distance between CO2 formed at the active-site of soluble CA compared to bound Rubisco assuming a mean hydrodynamic radius of ~50 Å for the distance of bound CA from the active-site of Rubisco (structural dimensions of HCAII a=45 Å, b=42 Å, c=73 Å). At 6 μg/250 μL, the concentration of Rubisco is 45 nM (~534,000 daltons). Free concentrations of CA at 0.3, 3, and 30 μg/250 μL represent 41, 410, and 4100 nM (~29,000 daltons). The molecular density of Rubisco at 45 nM is 2.7x1010 molecules/mm3. Therefore, the solution volume occupied by a single molecule of Rubisco at 45 nM is 3.7x1010 Å3, and a sphere containing this volume has a radius of 2067 Å. If CA bound to Rubisco is within ~50 Å (averaged dimensions of a CA tetrahedral) of the active-site then the ratio of the maximum distance between bound and free CA is ~41-fold. While the effectiveness of bound CA was mimicked by 50 μg of unbound CA, and 200s the efficiency of bound CA is 100-fold greater, this effect has not been titrated and may therefore be overestimated, thus reducing the actual efficiency increase. Nevertheless, clearly the possibility exist that CA targeted to Rubisco has increased efficiency for supporting carboxylase activity at low CO2, by facilitating equilibrium between HCO3— and CO2 in direct proximity to the active site of Rubisco. However, the nearly saturated levels of carboxylase activity in the presence of conjugated bound CA, and the 100-fold molar excess of free CA, is not readily explainable. This effect would presumably be manifested by a lowering of the ag Kd (CO2), but again how this might be accomplished is not clear. Perhaps the CA catalyzed equilibrium between HCO3— and CO2 in proximity to Rubisco results in a temporary increase in CO2 above that which would be present with free CA, and the effects of a 100-fold molar excess of CA mimic this effect through such a large enzyme concentration. The possibility that such a large molar excess of free CA relative to Rubisco resulted in some non-specific protein association was addressed by incubation of CA and Rubisco at these ratios and subsequent gel-filtration on Sephrose 6B as shown in FIG. 2. The results indicated no association between Rubisco and CA even at a 100:1 molar ratio.

Example 5

Construction and Purification of RLSMT-CA Fusion Protein

The RLSMT Carbonic Anhydrase (RLSMT-CA) fusion was constructed in a pET1 vector from Stratagene. The nucleic acid sequence encoding carbonic anhydrase (Accession number NM_000007) was fused to the carboxyl terminus of Pea RLSMT (Accession number Q43088), and cloned into the pET vector using pET vector protocols from Stratagene and other standard molecular biology protocols. SEQ ID No.: 1 is the nucleic acid sequence of the RLSMT-CA. SEQ ID No.: 2 is the amino acid sequence of the RLSMT-CA fusion protein. Again, using standard molecular biology protocols, this pET-RLSMT-CA vector was transformed into E. coli. E. coli expressing the pET-RLSMT-CA vector were cultured under conditions suitable for expression of the RLSMT-CA fusion protein. ~130 ml of this culture was pelleted, washed, and lysed using standard protocols. This lysate was resuspended in 5 ml standard buffer. 2.5 ml of this solution was loaded onto ~0.5 ml packed Spinach-Rubisco-conjugated activated Thiol-Sepharose 4B (T). 2.5 ml of this solution was loaded onto ~1.0 ml packed Spinach-Rubisco-conjugated Cyanogen bromide-Activated Sepharose (C). The resin was washed with 40 ml of 0.4 M NaCl in standard buffer. RLSMT-CA eluted with AdoMet: about ~3 h with 100 μM AdoMet in standard buffer with 100 μg/ml β-lactoglobulin (0.5 ml for T and 1.0 ml for C); concentrated with YM50. A Coomassie blue stained SDS-PAGE gel illustrating the elution of the RLSMT-CA is shown in FIG. 5. The Rubisco LSMTP-CA fusion protein eluted in lanes 5 and 9 contained 22.3 μmoles/min/μl of Rubisco methyltransferase activity and 0.007-0.034 Abs/sec/μl of carbonic anhydrase activity.

While the invention has been described and illustrated herein by references to various specific material, procedures and examples, it is understood that the invention is not restricted to the particular material combinations of material, and procedures selected for that purpose. Numerous variations of such details can be implied as will be appreciated by those skilled in the art.

US 8,138,309 B2
<223> OTHER INFORMATION: RLSTM-CA fusion

<400> SEQUENCE: 1

atggcaaaat cagtagcctc tgtaggaacc gaaccatcac tgcttcccagc agttczaacc  60
ttcggagagt ggctacagga agaagcgttc atcaactcaca agacccectg gaaagctgat 120
gtggcagcct agaagtagtg atgggtcgca tctaaggacca ttttagtgatt cgcttg180
ttcgaggac caaagaacct cctgatcatc cttgcaagctt ccagagtggc gccgagctgg 240
agaaggtgcc gtaagtggaag ccaactgttgg tctgtaatac ttcctctatt aagagagagg 300
tcaggggaga aattgttttg gaagcaact atctgattac tggcacaagga aactgattct 360
actataatt ggtgacaagga agaactccaa gacgctcaag gttcctcaact ttggaaaaac 420
acagttgctg tgaagaaata tgaagaagaat cagattttga agaatacacta agaatactat 480
tcctcataa agacgcttttt tcgcggactc gtagcgctgg attagttcctt ttggggcattt 540
ggaggtccgt gatacgcgct ctgctgcgtaaat ccactttctg tggtagtcgt 600
atgggagacg tcgtaaacaag ctagtcagga gttactacag aggacatgtg tcrraggtt 660
aaagagcgcg cctgcctttct tctcttcggt atcctttttt tctgaaagag ccctcccttc 720
gtcaagggcg ggaagaaggt atatatacaca tctgatttgga cacaagacca ccggcacgatg 780
gctctaagct acggctctag tgaaccaacta gaaactgcag atgcatacaca cctgacgcgt 840
gagatactcg agctgcgacc cttttttctg gacaacacta agctggctgtg gcgtcaagttgt 900
ctggcataca cagctctcct gtagatctctc tataatcgcct ctctcctccct tccttctct 960
ccatataca gactatgtaag gctagggggtt acgaccggctt cctattgaga atcactgtcc 1020
agagacacca tattggggcctgctg tggctcagcct gccgctcaatg ggcgactacta 1080
tgccaaagcg tcgtaaagctc ctagctgaactc gttctcagct gttacataca aaccctgtcga 1140
caaggttgcag gtaagtggga ccggaactctca gttcgaaggc tttggactgc agtgggaata 1200
agagagaggg aagagatggt cctgcagcagcc atgtcgaggttctgtcgacga gaaagactgt 1260
agogtgacgc aagatagaga tttcatcaagaa agaggacctca aggtcctttg atcttccgga 1320
ggaaataggg atacetctgcc aacactaggac aacttgccttc ctagaaggcg aggggaggcc 1380
gcggcgcgac gcacgctcaag tcgcctacac cttgggtcag ccggacaacca cggacgctag 1440
cacggcata agaacttccac aattgcaag ggagacgccc agtcctctcc tcgatcccaac 1500
aacttccacg ccaacatgtaa ccctttcccttg agcctccgcttg tcgcttcccaag tcgacaacca 1560
aacttccatg ggcctctcaaa taaatgtgattc tggcttcagcc cttgctggaaa ctagctcagc 1620
gcagagaagc gtagctcaaggg aacccctcttg gatggcaact cagagttgag tcagttccac 1680
ccctctctcc gttggcagtcg tcgacctggct tcgagagcaat cttgtgatgaa ccgagatgct 1740
gtgcgaacc gcgtgctggct cactgtggaac aacaattag gggtttttgg gaagagtgttg 1800
cagcgaccagc gcgtgtacgcg cttgggtctt acgcagatgtg cttgctggataa aagagcctat 1860
ggctgcctaa aggtgtgttg aagttggttg tcctttaaaa ccagggcctaa gagtgtgctac 1920	ttcactcaat tcagtcctgc ccgcttcctct ccgactatta cttggtacatc cagctccaca 1980
ggctgcctaa ccccccctcc ttctgtgggt gggttgtgtg agatgacctc ggagaacacc 2040
atcgctctca gcgcggagcgc cttgctggaaa tccgttaaaa ctccttaaagc ttgggggctgt 2100
ggaccccaag aagattgtct ggaaacaacttg cgyccagcto acgcttaaga gaaagggcga 2160
atcacaagctt cctctcataa a 2181
<table>
<thead>
<tr>
<th>Position</th>
<th>Amino Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Met</td>
</tr>
<tr>
<td>5</td>
<td>Ala</td>
</tr>
<tr>
<td>10</td>
<td>Lys</td>
</tr>
<tr>
<td>15</td>
<td>Ser</td>
</tr>
<tr>
<td>20</td>
<td>Val</td>
</tr>
<tr>
<td>25</td>
<td>Ala</td>
</tr>
<tr>
<td>30</td>
<td>Ser</td>
</tr>
<tr>
<td>35</td>
<td>Val</td>
</tr>
<tr>
<td>40</td>
<td>Thr</td>
</tr>
<tr>
<td>45</td>
<td>Ala</td>
</tr>
<tr>
<td>50</td>
<td>Leu</td>
</tr>
<tr>
<td>55</td>
<td>Lys</td>
</tr>
<tr>
<td>60</td>
<td>Arg</td>
</tr>
<tr>
<td>65</td>
<td>Leu</td>
</tr>
<tr>
<td>70</td>
<td>Arg</td>
</tr>
<tr>
<td>75</td>
<td>Asp</td>
</tr>
<tr>
<td>80</td>
<td>Ile</td>
</tr>
<tr>
<td>85</td>
<td>Ser</td>
</tr>
<tr>
<td>90</td>
<td>Val</td>
</tr>
<tr>
<td>95</td>
<td>Arg</td>
</tr>
<tr>
<td>100</td>
<td>Leu</td>
</tr>
<tr>
<td>105</td>
<td>Arg</td>
</tr>
<tr>
<td>110</td>
<td>Ser</td>
</tr>
<tr>
<td>115</td>
<td>Gln</td>
</tr>
<tr>
<td>120</td>
<td>Pro</td>
</tr>
<tr>
<td>125</td>
<td>Asp</td>
</tr>
<tr>
<td>130</td>
<td>Lys</td>
</tr>
<tr>
<td>135</td>
<td>Leu</td>
</tr>
<tr>
<td>140</td>
<td>Thr</td>
</tr>
<tr>
<td>145</td>
<td>Tyr</td>
</tr>
<tr>
<td>150</td>
<td>Val</td>
</tr>
<tr>
<td>155</td>
<td>His</td>
</tr>
<tr>
<td>160</td>
<td>Leu</td>
</tr>
<tr>
<td>165</td>
<td>Pro</td>
</tr>
<tr>
<td>170</td>
<td>Asp</td>
</tr>
<tr>
<td>175</td>
<td>Phe</td>
</tr>
<tr>
<td>180</td>
<td>Ala</td>
</tr>
<tr>
<td>185</td>
<td>Phe</td>
</tr>
<tr>
<td>190</td>
<td>Gly</td>
</tr>
<tr>
<td>195</td>
<td>Arg</td>
</tr>
<tr>
<td>200</td>
<td>Ser</td>
</tr>
<tr>
<td>205</td>
<td>Leu</td>
</tr>
<tr>
<td>210</td>
<td>Val</td>
</tr>
<tr>
<td>215</td>
<td>Pro</td>
</tr>
<tr>
<td>220</td>
<td>Met</td>
</tr>
<tr>
<td>225</td>
<td>Gly</td>
</tr>
<tr>
<td>230</td>
<td>Phe</td>
</tr>
<tr>
<td>235</td>
<td>Ser</td>
</tr>
<tr>
<td>240</td>
<td>Leu</td>
</tr>
<tr>
<td>245</td>
<td>Tyr</td>
</tr>
<tr>
<td>250</td>
<td>Asp</td>
</tr>
<tr>
<td>255</td>
<td>Pro</td>
</tr>
<tr>
<td>260</td>
<td>Ala</td>
</tr>
<tr>
<td>265</td>
<td>Gly</td>
</tr>
<tr>
<td>270</td>
<td>Asp</td>
</tr>
<tr>
<td>275</td>
<td>His</td>
</tr>
<tr>
<td>280</td>
<td>Ser</td>
</tr>
<tr>
<td>285</td>
<td>Arg</td>
</tr>
<tr>
<td>290</td>
<td>Phe</td>
</tr>
<tr>
<td>295</td>
<td>Arg</td>
</tr>
<tr>
<td>300</td>
<td>Ala</td>
</tr>
<tr>
<td>305</td>
<td>Tyr</td>
</tr>
<tr>
<td>310</td>
<td>Leu</td>
</tr>
<tr>
<td>315</td>
<td>Pro</td>
</tr>
<tr>
<td>320</td>
<td>Gly</td>
</tr>
<tr>
<td>325</td>
<td>Leu</td>
</tr>
<tr>
<td>330</td>
<td>Arg</td>
</tr>
<tr>
<td>335</td>
<td>Phe</td>
</tr>
<tr>
<td>340</td>
<td>Ser</td>
</tr>
<tr>
<td>345</td>
<td>Leu</td>
</tr>
<tr>
<td>350</td>
<td>Val</td>
</tr>
<tr>
<td>355</td>
<td>Asp</td>
</tr>
<tr>
<td>360</td>
<td>Ala</td>
</tr>
<tr>
<td>365</td>
<td>Cys</td>
</tr>
<tr>
<td>370</td>
<td>Ser</td>
</tr>
<tr>
<td>375</td>
<td>Ala</td>
</tr>
<tr>
<td>380</td>
<td>Lys</td>
</tr>
</tbody>
</table>
The invention claimed is:

1. A modified Rubisco large subunit "N-methyltransferase (RLSMT), comprising a Rubisco large subunit "N-methyltransferase linked to a second molecule and wherein the modified RLSMT has the ability to bind a Rubisco.

2. The modified RLSMT of claim 1, wherein the Rubisco large subunit "N-methyltransferase is linked to the second molecule by a chemical.

3. The modified RLSMT of claim 2, wherein the chemical is glutaraldehyde.

4. The modified RLSMT of claim 2, wherein the modified RLSMT is a fusion protein, wherein the fusion protein is encoded by a nucleic acid, wherein the nucleic acid comprises first nucleic acid encoding a Rubisco large subunit "N-methyltransferase, and wherein the nucleic acid further comprises a second nucleic acid encoding a polypeptide.

5. The modified RLSMT of claim 1, wherein the second molecule is a phosphatase, 3-phosphoglycerate kinase, glyc-
eraldehydes-3-phosphate dehydrogenase, Rubisco Activase, Phosphoribulokinase, polylysine, or oxygenase.

7. The modified RLSMT of claim 1, wherein the modified RLSMT is a fusion protein, wherein the fusion protein is encoded by a nucleic acid.

8. A modified Rubisco large subunit "N-methyltransferase of claim 1, wherein said second molecule is a carbonic anhydrase, wherein the modified RLSMT (RLSMT-CA) has the ability to bind a Rubisco large subunit.

9. The modified RLSMT of claim 8, wherein the Rubisco large subunit "N-methyltransferase is linked to the carbonic anhydrase (CA) by a chemical.

10. The modified RLSMT of claim 9, wherein the chemical is glutaraldehyde.

11. The modified RLSMT of claim 8, wherein the modified RLSMT is a fusion protein, wherein the fusion protein is encoded by a nucleic acid, wherein the nucleic acid comprises a first nucleic acid encoding a Rubisco large subunit "N-methyltransferase, and wherein the nucleic acid further comprises a second nucleic acid encoding a carbonic anhydrase.

12. The modified RLSMT of claim 8, wherein the modified RLSMT further comprises an antibody.

13. The modified RLSMT of claim 11, wherein the antibody is linked to a CA.

14. The modified RLSMT of claim 8 comprising SEQ ID No.: 2.