



Jun 30, 2020

# ♦ NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state – GAPDH-GlyPDH

In 1 collection

Cristina Rodrigues Gabriel Sales<sup>1</sup>, Anabela Silva<sup>2</sup>, Elizabete Carmo-Silva<sup>1</sup>

<sup>1</sup>Lancaster Environment Centre, Lancaster University, Library Avenue, Lancaster, LA1 4YQ, UK;

<sup>2</sup>Biosystems & Integrative Sciences Institute (BioISI), Science Faculty of Lisbon University, Lisbon, 1749-016, Portugal

1 Works for me

dx.doi.org/10.17504/protocols.io.bgasjsee



Cristina Rodrigues Gabriel Sales

#### ABSTRACT

This protocol uses five reactions to couple RuBP carboxylation and 3-PGA formation to NADH oxidation to measure Rubisco activity, based on Kubien et al. (2011).





During the NADH-linked assays, some Rubisco active sites might become carbamylated as the leaf extract is exposed to high  $\rm CO_2$  and  $\rm Mg^{2+}$  in the assay buffer. Therefore, these assays are not suitable for measuring Rubisco initial activity and/or activation state at different levels in the canopy or in conditions in which low intercellular  $\rm CO_2$  is promoted (e.g., low light, drought stress, cold stress); the 30 second  $\rm ^{14}CO_2$ -based assay is recommended in such situations (protocol available in this collection).

## THIS PROTOCOL ACCOMPANIES THE FOLLOWING PUBLICATION

Sales CRG, Silva AB, Carmo-Silva E. 2020. Measuring Rubisco activity: challenges and opportunities of NADH-linked microtiter plate-based and 14C-based assays. Journal of Experimental Botany, https://doi.org/10.1093/jxb/eraa289

DO

dx.doi.org/10.17504/protocols.io.bgasjsee

#### PROTOCOL CITATION

Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva 2020. NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state – GAPDH-GlyPDH. **protocols.io** dx.doi.org/10.17504/protocols.io.bgasjsee

MANUSCRIPT CITATION please remember to cite the following publication along with this protocol

G

Sales CRG, Silva AB, Carmo-Silva E. 2020. Measuring Rubisco activity: challenges and opportunities of NADH-linked microtiter plate-based and 14C-based assays. Journal of Experimental Botany, https://doi.org/10.1093/jxb/eraa289

COLLECTIONS (i)

Protocols from Sales et al. (2020) Rubisco activity: challenges and opportunities of NADH-

protocols.io

06/30/2020

Citation: Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva (06/30/2020). NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state â GAPDH-GlyPDH. <a href="https://dx.doi.org/10.17504/protocols.io.bgasjsee">https://dx.doi.org/10.17504/protocols.io.bgasjsee</a>

linked microtiter plate-based and 14C-based assays

Protocols from Sales et al. (2020) Rubisco activity: challenges and opportunities of NADH-linked microtiter plate-based and 14C-based assays

#### KFYWORDS

Enzyme activity assay, Rubisco, Crop improvement , NADH-linked assay, GAPDH-GlyPDH, Plant phenotyping, Microtiter plate

#### LICENSE

This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

#### CREATED

May 12, 2020

LAST MODIFIED

Jun 30, 2020

PROTOCOL INTEGER ID

36914

PARENT PROTOCOLS

#### Part of collection

Protocols from Sales et al. (2020) Rubisco activity: challenges and opportunities of NADH-linked microtiter plate-based and 14C-based assays

Protocols from Sales et al. (2020) Rubisco activity: challenges and opportunities of NADH-linked microtiter plate-based and 14C-based assays

# GUIDELINES

- 1. Check the "Materials" tab for a list of all the chemicals used in this protocol.
- 2. In the "Steps" tab, there is a brief description of the materials and equipment necessary for the protocol execution.
- 3. In the "Steps" tab, there is information on preparation of solutions, procedures for determining Rubisco initial and total activities, and notes to take into consideration to ensure reliable results.
- 4. The references cited are at the end of the "Materials" tab.

# MATERIALS

NAME	CATALOG #	VENDOR
Bicine	B3876	Sigma Aldrich
Magnesium chloride hexahydrate (MgCl2.6H20)	M2393	Sigma Aldrich
Ethylenediaminetetraacetic acid disodium salt dihydrate (EDTA)	E1644	Sigma Aldrich
Benzamidine	B6506	Sigma Aldrich
ε-Aminocaproic acid	A2504	Sigma Aldrich
Sodium hydroxide (NaOH)	S5881	Sigma Aldrich
2-Mercaptoethanol	M6250	Sigma Aldrich
DL-Dithiothreitol (DTT)	43819	Sigma Aldrich
Phenylmethanesulfonyl fluoride (PMSF)	P7626	Sigma Aldrich
Protease inhibitor cocktail	P9599	Sigma Aldrich
D-Ribulose 1.5-bisphosphate sodium salt hydrate (RuBP)	83895	Sigma Aldrich

protocols.io
2
06/30/2020

Citation: Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva (06/30/2020). NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state â GAPDH-GlyPDH. https://dx.doi.org/10.17504/protocols.io.bgasjsee

NAME	CATALOG #	VENDOR
Sodium bicarbonate (NaHCO3)	S6014	Sigma Aldrich
$\beta\textsc{-Nicotinamide}$ adenine dinucleotide reduced disodium salt hydrate (NADH)	N8129	Sigma Aldrich
Phosphocreatine disodium salt hydrate	P7936	Sigma Aldrich
Adenosine 5'-triphosphate disodium salt hydrate (ATP)	A3377	Sigma Aldrich
Creatine phosphokinase from rabbit muscle (PCK)	C3755	Sigma Aldrich
Glyceraldehyde-3-phosphate dehydrogenase from rabbit muscle (GAPDH)	G2267	Sigma Aldrich
3-Phosphoglyceric phosphokinase from bakers yeast (PGK)	P7634	Sigma Aldrich
α-Glycerophosphate dehydrogenase/Triosephosphate isomerase (GlyPDH/TPI)	G1881	Sigma Aldrich
Ethanol absolute 99.8 %	10437341	Fisher Scientific

#### MATERIALS TEXT

- Kane HJ, Wilkin JM, Portis AR, Andrews TJ (1998). Potent inhibition of ribulose-bisphosphate carboxylase by an oxidized impurity in ribulose-1,5-bisphosphate. Plant Physiology 117: 1059-1069. https://doi.org/10.1104/pp.117.3.1059
- Kubien DS, Brown CM, Kane HJ (2011). Quantifying the amount and activity of Rubisco in leaves. Methods in Molecular Biology (Clifton, N.J.) 684: 349-362. https://doi.org/10.1007/978-1-60761-925-3\_27
- Lampinen J, Raitio M, Perälä A, Oranen H, Harinen R (2015).
  Microplate based pathlength correction method for
  photometric DNA quantification assay. Application
  Laboratory, Sample Preparation & Analysis, Thermo Fisher
  Scientific, Vantaa, Finland.
  https://assets.thermofisher.com/TFSAssets/LCD/Application-Notes/AN-SkanIT-Microplate-BasedPathlength-Correction-Technical-Note
- Sales CRG, Degen GE, Silva AB, Carmo-Silva E (2018).
  Spectrophotometric determination of Rubisco activity and activation state in leaf extracts. In: Covshoff S, ed. Methods in Molecular Biology. New York: Humana Press 1770: 239-250. https://doi.org/10.1007/978-1-4939-7786-4\_14

protocols.io
3
06/30/2020

Citation: Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva (06/30/2020). NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state â GAPDH-GlyPDH. <a href="https://dx.doi.org/10.17504/protocols.io.bgasjsee">https://dx.doi.org/10.17504/protocols.io.bgasjsee</a>

Sharwood RE, Sonawane BV, Ghannoum O, Whitney SM (2016). Improved analysis of C4 and C3 photosynthesis via refined in vitro assays of their carbon fixation biochemistry. Journal of Experimental Botany 67: 3137-3148. https://doi.org/10.1093/jxb/erw154

Wong C-H (1980). Practical enzymatic syntheses of ribulose 1,5-bisphosphate and ribose 5-phosphate. Journal of the American Chemical Society 102: 7938-7939. https://doi.org/10.1021/ja00547a023

#### SAFETY WARNINGS

Before using the protocol always check the Safety Data Sheet (SDS) for each chemical.

BEFORE STARTING

#### MATERIAL & EQUIPMENTS (for list of chemicals check "Materials" tab)

- Leaf sample frozen in -80°C
- Centrifuge for microtubes (speed 14000 g, 4 °C; VWR, Mega Star 600R)
- Microtiter plate reader (BMG Labtec, SpectroStarNano)
- 96-well microtiter plate with clear flat bottom (Thermo Scientific, 442404)
- Pipette set
- Mortar and pestle
- 1.5 mL microtubes

### REAGENTS & SOLUTIONS

#### **1 REAGENTS & SOLUTIONS TO PREPARE BEFOREHAND**



- Powder chemical stocks stored at -20°C: let warm up to room temperature on desiccant before opening container.
- Expensive chemicals purchased in very small amounts (mg), for which concentration in assay is not
  critical (e.g. in excess): trust quantity stated by the supplier and add ultrapure H<sub>2</sub>O / solvent to container
  for final concentration (e.g. PCK).
- Protein and substrate solutions should typically be stored at -80°C.
- Freeze proteins in LN<sub>2</sub> before storing at -80°C. Store in small aliquots to prevent multiple freeze-thaw cycles. If using in consecutive days, protein solutions can be kept at 4°C.
- $\,\blacksquare\,$  Buffers and solutions will last longer if filtered through 0.22  $\mu m$  membrane.

# 1.1 Basic extraction buffer (1x)

[M]50 Milimolar (mM) Bicine-NaOH pH8.2
[M]20 Milimolar (mM) MgCl<sub>2</sub>.6H<sub>2</sub>O
[M]1 Milimolar (mM) EDTA
[M]2 Milimolar (mM) Benzamidine

protocols.io
4
06/30/2020

Citation: Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva (06/30/2020). NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state â GAPDH-GlyPDH. <a href="https://dx.doi.org/10.17504/protocols.io.bgasjsee">https://dx.doi.org/10.17504/protocols.io.bgasjsee</a>

```
[M]5 Milimolar (mM) E-Aminocaproic acid
```

Dissolve in ultrapure H<sub>2</sub>O; adjust pH to 8.2 with NaOH; degas the solution bubbling with nitrogen (5 min/10 mL), then add:

[M]50 Milimolar (mM) 2-Mercaptoethanol

• Adjust for the final volume; it can be dispensed in aliquots (e.g. 50 mL Falcon tubes).

```
8 -20 °C (storage)
```

- 1.2 [M] 1 Molarity (M) DTT
  - Dissolve in ultrapure H<sub>2</sub>O. § 4 °C (storage)
- 1.3 [M]100 Milimolar (mM) PMSF
  - Dissolve in ethanol 99%. § 4 °C (storage)
- 1.4 Plant protease inhibitor cocktail

```
§ -20 °C (storage)
```

1.5 [M]20 Milimolar (mM) RuBP

```
§ -20 °C (storage)
```



High purity RuBP (≥99%) is required to avoid interference in measurable activity due to the presence of RuBP-analogs that inhibit carboxylation (Kane et al., 1998; Sharwood et al., 2016). It is available commercially or it can be produced enzymatically from AMP-5′ monohydrate and ATP disodium salt (Wong, 1980).

## **2 STOCK COMPONENTS FOR THE ASSAY BUFFER**

```
[M]1 Molarity (M) Bicine-NaOH pH8
```

Dissolve in ultrapure H<sub>2</sub>O; adjust pH to 8.2 with NaOH; filter through 0.22 μm membrane for long shelf life.

```
8 4 °C (storage)
```

```
[M]1 Molarity (M) MgCl<sub>2</sub>.6H<sub>2</sub>O
```

Dissolve in ultrapure H<sub>2</sub>O; filter through 0.22 μm membrane for long shelf life. δ 4 °C (storage)

```
[M] 0.5 Molarity (M) NaHCO3
```

Dissolve in ultrapure H<sub>2</sub>O; filter through 0.22 μm membrane for long shelf life. δ 4 °C (storage)

```
[M] 1 Molarity (M) Phosphocreatine
```

Dissolve in ultrapure H<sub>2</sub>O; aliquot; § -80 °C (storage)

```
[M]80 Milimolar (mM) ATP
```

Add powder to a tube, add some ultrapure H<sub>2</sub>O to dissolve, add [M]4 Molarity (M) NaOH (~10 μL/mL) to approximately pH7; add ultrapure H<sub>2</sub>O to the final volume; aliquot. δ -80 °C (storage)

protocols.io
5
06/30/2020

Citation: Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva (06/30/2020). NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state â GAPDH-GlyPDH. https://dx.doi.org/10.17504/protocols.io.bgasjsee



To test the pH use universal pH paper, add  $\Box 1 \mu I$  of the solution and check colour.

#### [M]14 Milimolar (mM) NADH

■ Dissolve in [M] 100 Molarity (m) Bicine-NaOH pH8 ; aliquot. 8 -80 °C (storage)



Protect from light as it is light sensitive.

Please, check notes in section 2.1 to a more efficient way of aliquoting NADH.

#### [M]1.8 KU/ml PCK

Dissolve all solid by adding [M]100 Molarity (m) Bicine-NaOH | pH8 | to the container; aliquot. § -80 °C (storage)

#### [M] 5 KU/ml GAPDH

Dissolve all solid by adding [M]100 Molarity (m) Bicine-NaOH pH8 to the container; aliquot. & -80 °C (storage)

#### [M] 17.9 KU/ml PGK

- Store at 8 4 °C when in ammonium sulfate suspension.
- Before use, it is necessary to buffer exchange as ammonium sulfate can affect the assay.
- Pipette a portion, e.g.,100 μL.
- Proceed to a buffer exchange using Amicon molecular weight cut-off of 10000 (Sigma-Aldrich); substitute ammonium sulfate for [M]100 Molarity (m) Bicine-NaOH pH8 in the same volume centrifuged (in this example 100 μL).
- Assay for TSP to know the concentration; aliquot. § -80 °C (storage)

# [M]1.8 KU/ml GlyPDH+ [M]180 KU/ml TPI (GlyPDH/TPI)

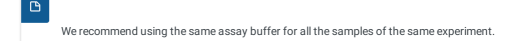
- Store at § 4 °C when in ammonium sulfate suspension.
- Before use, it is necessary to buffer exchange as ammonium sulfate can affect the assay.
- Pipette a portion, e.g.,100 μL.
- Proceed to a buffer exchange using Amicon molecular weight cut-off of 10000 (Sigma-Aldrich); substitute ammonium sulfate for [M]100 Molarity (m) Bicine-NaOH pH8 in the same volume centrifuged (in this example 100 μL).
- Assay for TSP to know the concentration; aliquot. § -80 °C (storage)

# 2.1 Complete assay buffer for Rubisco activity

- The basic assay buffer (table below) can be prepared the day before the assays and kept at & 4 °C , or prepared in advance (e.g. at the start of an experiment), snap-frozen in aliquots and kept at & -80 °C .
- Each assay buffer aliquot should only be thawed once, as repeated freeze thawing can result in degradation of the coupling enzymes; thus, it is important to aliquot adequate volumes for use in a day.
- NADH is prepared separately, snap-frozen in aliquots and kept at & -80 °C , and added to the assay buffer just before the assays.

Final concentration component	Stock	Volume in 200 µL assay (µL)
100 mM Bicine-NaOH pH 8.2	1 M	20
20 mM MgCl <sub>2</sub>	1 M	4
10 mM NaHCO₃	0.5 M	4
5 mM DTT	1 M	1
5 mM Phosphocreatine	1 M	1
1 mM ATP	80 mM	2.5
25 U mL <sup>-1</sup> PCK	1.8 KU mL <sup>-1</sup>	2.8
25 U mL <sup>-1</sup> GAPDH	5 KU mL <sup>-1</sup>	1
25 U mL <sup>-1</sup> PGK*	17.9 KU mL <sup>-1</sup>	0.28
20 / 200 U mL <sup>-1</sup> GlyPDH/TPI*	1.80 / 180 KU mL <sup>-1</sup>	2.2
Total		38.8
To be prepared separatel	y and added just before the	assays
0.4 mM NADH	14 mM	5.71
Total for each well		44.5





Stock solutions and the assay buffer should thaw § On ice. The assay buffer should be kept in a tube wrapped in aluminium foil § On ice during the assays, as NADH is light sensitive.

#### **3 SOLUTIONS TO PREPARE JUST BEFORE USE**

• Prepared with reagents/solutions described in step 1.

## 3.1 Complete extraction buffer

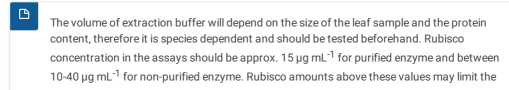
1x Basic extraction buffer (from step 1.1)

[M] 10 Milimolar (mM) DTT (from step 1.2)

[M] 1 Milimolar (mM) PMSF (from step 1.3)

[M] 1 % (v/v) Plant protease inhibitor cocktail (from step 1.4)

 Prepare the volume considering the number of extractions to be performed throughout the day plus two extras (to have a little excess). Mix all together.
 On ice



protocols.io
7
06/30/2020



To test if the assay is giving reliable results (i.e, none of the chemicals are limiting the reactions) it is important to always perform a test when the plant species and/or growth conditions change. Perform the assay with different extract concentration (e.g. 1/2 the amount, 1/5 of the amount, etc) and check if the activity expressed per protein content (TSP or Rubisco) is mantained.

## 3.2 Complete assay buffer

- Thaw & On ice the assay buffer and NADH aliquots (prepared according to step 2.1) to be used
  in the day.
- Mix the correspondent volume of both solutions together. Keep § On ice, wrapped in aluminium



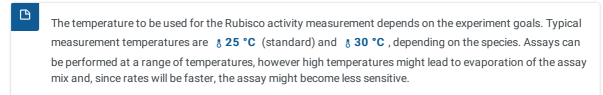
Example of how to prepare the complete assay buffer: e.g. 20 samples in a day

- 20 samples x (1 Blank + 3xTotal + 3xInitial activity assays) = 140 wells
- Assay buffer without NADH per well = 38.8, for 140 wells = 38.8 x 140 = 5432μL
- NADH per well = 5.71, for 140 wells = 5.71 x 140 =  $799.4 \mu L$
- Mix both together (5432 + 799.4 μL)

#### PROCEDURE

## 4 START

- Thaw the frozen solutions that will be used in the day.
- Prepare CO<sub>2</sub>-free ultrapure H<sub>2</sub>O by bubbling with nitrogen (5 min/100 mL).
- Turn on the microplate reader and set up for the temperature that Rubisco activity will be performed, select kinetic protocol at 340 nm.



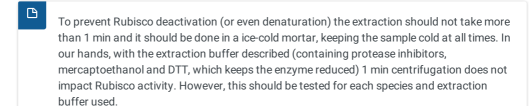
- Turn on the centrifuge and set to § 4 °C.
- Collect samples from § -80 °C into liquid nitrogen container.
- Prepare the complete extraction buffer (from step 3.1) and the complete assay buffer (from step 3.2) and keep it
   On ice

#### **5 EXTRACTIONS & RUBISCO ASSAYS**

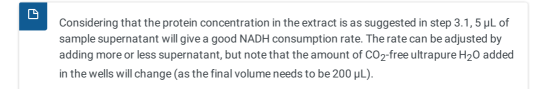
- 5.1 Before starting the extraction, pipette to the 96-well microtiter plate 150.5 μl CO<sub>2</sub>-free ultrapure H<sub>2</sub>O for the blank (singlet) and 144.5 μl for the samples into each well (6 wells, i.e., triplicates for initial activities and triplicates for total activities), followed by 44.5 μl of complete assay buffer (from step 3.2).
  - Gently mix components by pipetting up and down 5 times whilst stirring. Add a6 μl of 20 mM RuBP (from step 1.5) to the wells for measuring initial Rubisco activity (see table below). Cover to protect from light. Proceed to extraction.

# 5.2 Extraction

- Add the complete extraction buffer (from step 3.1) to an ice-cold mortar.
- Take a sample from the liquid nitrogen container and add to the mortar.
- Grind the sample thoroughly for **© 00:00:30** to maximum of **© 00:01:00**.
- Collect the homogenate into an ice-cold 1.5 mL microtube and centrifuge
   14000 x g, 4°C 00:01:00 .



- When centrifugation stops, take the extract supernatant into another ice-cold 1.5 mL microtube.
- Proceed to the Rubisco assays straight away.
- Add ■5 µl of sample supernatant to the wells for total activity first, followed by those for initial activity, mixing well by pipetting up and down 10 times whilst stirring. Place microplate in the reader and start monitoring the change in absorbance at 340 nm immediately. The addition of the extract initiates the reaction for the initial activity assays, which is measured while incubating Rubisco with CO<sub>2</sub> and MgCl<sub>2</sub> in the absence of RuBP (total activity) for ⑤00:03:00 at § 30 °C to enable carbamylation of the enzyme. The absorbance value should start decreasing in the wells for the initial activity assay (containing RuBP).
- Pause the reading in order to add ☐6 µl of 20 mM RuBP (from step 1.5) to the wells for measurement of total Rubisco activity ⊙ 00:03:00 after addition of sample supernatant.
- Place the microplate in the reader and continue monitoring the change in absorbance.
- The reading can be stopped once the reaction reaches a plateau.



The Initial activity assays start with extract addition, while total activity assays start with addition of RuBP after 3 min of extract incubation with CO<sub>2</sub> and Mg<sup>2+</sup> to allow for Rubisco carbamylation.

protocols.io 9 06/30/2020



This protocol can be adapted for measuring Rubisco activity with purified enzyme. In this case, Rubisco is frequently pre-activated and initial activity assays are performed.

Below is a pipetting scheme for the microplate assay

Solution (in pipetting order)		Volume to add (µL)		
	Blank	Initial activity	Total activity	
	To be added before	the extraction		
CO <sub>2</sub> -free H <sub>2</sub> O	150.5	144.5	144.5	
Complete assay buffer	44.5	44.5	44.5	
20 mM RuBP	0	6	0	
Т	o be proceeded aft	er the extraction		
Leaf extract	5	5	5	
Start measuring absorbance at 340 Pause readii		g the plate at the desired t art reaction for total activit		
20 mM RuBP	0	0	6	
Continue measuri	ing absorbance at 3	40 nm until a plateau is rea	ached.	





It is important to ensure that air bubbles are not introduced in the wells during the pipetting steps, as these will interfere with the absorbance measurements.

#### CALCULATIONS

- The activity of Rubisco is inferred from the consumption of RuBP (μmol s<sup>-1</sup>) measured by absorbance change per second at 340 nm due to NADH oxidation, using an extinction coefficient of 6220 M<sup>-1</sup>cm<sup>-1</sup> or 6.22 μmol<sup>-1</sup> mL cm<sup>-1</sup>.
  - The carboxylation of one molecule of RuBP results in two molecules of 3-PGA, thus requiring four NADH in the subsequent coupling reactions. The rate of RuBP consumption (μmol s<sup>-1</sup>) in the assay volume, is therefore calculated by:

$$RuBP consumption = (Slope imes Volume)/(6.22 imes 4 imes Pathlength)$$

where the *Slope* represents the change in absorbance per second in the linear part of the absorbance trace change, *Volume* is the final volume per well in mL (0.2), 6.22 is the extinction coefficient of NADH in  $\mu$ mol<sup>-1</sup> mL cm<sup>-1</sup> and the factor 4 is used to account for the four molecules of NADH which are oxidized per molecule of RuBP. The *Pathlength* of the assay mix contained in each well is measured in cm.

■ Rubisco initial (VI) and total (VI) activity expressed on a leaf area basis (µmol m<sup>-2</sup> s<sup>-1</sup>) is then calculated by:

$$V_iorV_t = (RuBPconsumption imes Extraction)/(Leafarea imes Aliquot)$$

protocols.io
10
06/30/2020

Citation: Cristina Rodrigues Gabriel Sales, Anabela Silva, Elizabete Carmo-Silva (06/30/2020). NADH-linked microtiter plate-based assay for measuring Rubisco activity & activation state â GAPDH-GlyPDH. https://dx.doi.org/10.17504/protocols.io.bgasjsee

where the Extraction is the volume of buffer in mL used for leaf extraction, leaf area is in m<sup>2</sup>, and Aliquot is the volume of leaf extract supernatant used in the assay in mL.

• Rubisco activity can also be expressed on a Rubisco or total soluble protein (TSP) content basis (μmol min<sup>-1</sup>mg<sup>-1</sup>):

$$V_iorV_t = (RuBPconsumption \times 60)/(Protein \times Aliquot)$$

where 60 is to convert seconds to minutes, *Protein* is the Rubisco or TSP content in mg mL<sup>-1</sup>, and *Aliquot* is the volume of leaf extract supernatant used in the assay in mL.

• From the Rubisco activity calculations above for initial (*Vi*) and total activity (*Vt*), the Rubisco activation state (AS, %) can be calculated:

$$AS = 100 \times V_i/V_t$$

Measured absorbance values in a microtiter plate need to be normalized to a 1 cm pathlength, which would be found in a typical cuvette used in spectrophotometers. Measurements are corrected using Lambert-Beer's Law and considering both the volume in each well and the specific well dimensions for each type of microtiter plate. Modern microtiter plate readers frequently include a pathlength correction option, but this feature normally does not consider the properties of the solution. It is important to use the respective assay mix in determining the pathlength correction factor as the meniscus will affect the pathlength and absorbance reading in the microtiter plate. The pathlength can be determined according to Lampinen et al., (2012). Please, check the SI information in the publication linked to this protocol for more details.