C4 photosynthesis with Kranz anatomy evolved in the Oryza coarctata Roxb.

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Abstract

The C4 cycle is a complex biochemical pathway that has been evolved in plants to deal with the

adverse environmental conditions. Mostly C4 plants grow in arid, water-logged area or poor

nutrient habitats. Wild species, *Oryza coarctata* (genome type KKLL; chromosome number (4x)

=48, genome size 665 Mb) belongs to the genus of *Oryza* which thrives well under high saline as

well as submerged conditions. Here, we report for the first time that O. coarctata is a C4 plant by

observing the increased biomass growth, morphological features such as vein density, anatomical

features including ultrastuctural characteristics as well as expression patterns of C4 related

genes. Leaves of O. coarctata have higher vein density and possess Kranz anatomy. The

ultrastructural observation showed chloroplast dimorphism i.e. presence of agranal chloroplasts

in bundle sheath cells whereas, mesophyll cells contain granal chloroplasts. The cell walls of

bundle sheath cells contain tangential suberin lamella. The transcript level of C4 specific genes

such as phosphoenolpyruvate carboxylase, pyruvate orthophosphate dikinase, NADP-dependent

malic enzyme and malate dehydrogenase was higher in leaves of O. coarctata compare to high

yielding rice cultivar (IR-29). These anatomical, ultra structural as well as molecular changes in

O. coarctata for C4 photosynthesis adaptation might be might be due to its survival in wide

diverse condition from aquatic to saline submerged condition. Being in the genus of *Oryza*, this

plant could be potential donor for production of C4 rice in future through conventional breeding,

as successful cross with rice has already been reported.

Introduction

Most of the plant species rely on RuBsiCO (Ribulose-1,5-bisphosphate carboxylase) enzyme for

carboxylation of atmospheric carbon dioxide to produce phosphoglyceric acid (Calvin et al.,

1949). The adverse environmental conditions allow the evolution of another biochemical

mechanism to fix carbon dioxide by phosphoenolpyruvate carboxylase (PEPC) to produce

oxaloacetic acid (Ehleringer et al., 1997). Beside this biochemical evolution, there are two major

anatomical characteristics that had been evolved in C4 plants. They are i) presence of special

anatomy of leaf i.e. Kranz (ring/ wreath-like) anatomy where bundle sheath cells surround

vascular bundles (Dengler and Nelson, 1999) and ii) chloroplast dimorphism i.e. mesophyll

chloroplasts contain well developed grana and bundle sheath chloroplasts are either agranal or

possess unstacked grana (thylakoids) (Laetsch, 1974). The C4 plants evolved around 20 to 30

million years ago (Kellogg, 1999). So, far C4 photosynthesis had been found in 21 taxonomical

groups i.e. 18 dicots (2,994 species) and 3 monocots (11,900 species) (Sage at al., 2001).

However, it has been suggested that C4 plants are more like to be tolerant to different abiotic

stresses (Bromham and Bennett, 2014).

Oryza genus comprises of 24 species out of which two species are edible and rests are grown as

wild species (Sanchez et al., 2013). The wild plant species are excellent source of stress tolerance

genes (Mondal et al., 2018). However, rice is a C3 plant therefore several attempts have been

made to transfer C4 genes from maize into rice for increasing yield as well as better tolerance of

stress (Ku et al., 1999; Takeuchi et al., 2000). O. coarctata is a wild halophyte found in the

coastal regions. It had ability to tolerate high salt stress condition (Electrical conductivity-E.Ce

20-40dSm⁻¹) and submergence condition for 12h (Sengupta and Mujumder, 2010; Chowrasia et

al., 2019; 2018). Recently the genome sequence of this wild species of rice has been decoded in

our laboratory (Mondal et al., 2018). Several salt stress responsive genes from this plant had

been characterized (Sengupta et al., 2009; Kizhakkedath et al., 2015). Moreover, attempts were

made for introgression of salt stress responsive genes from O. coarctata into rice cultivar (Jena,

1994; Islam et al., 2017) through conventional crossing method. Although, O. coarctata is an

obligate halophytic grass but no attempt was made to study its mode of photosynthesis.

Moreover, it is well-known that stress tolerant plants found in arid and saline habitats are mostly

C4 (Bromham and Bennett, 2014). Therefore, present study deals with biomass growth,

anatomical and ultrastructural features as well as expression pattern of C4 related genes in the

leaves of *O.coarctata* to understand the mode of photosynthesis.

Methods

Plant materials

The plantlets of O. coarctata was collected from coastal region of Sunderban, West Bengal and

grown in net house of NIPB (National Research Centre for Plant Biotechnology), New Delhi

under control condition in soil under 16h daylight and 8h dark, an irradiance of 56 µmol m⁻² s⁻¹

temperatures of 30°C, and a relative humidity of 60 -70 %. The rice seeds of IR-29 were sown in

soil and maintained under same condition. All sampling was done at morning between 10-11 am

3

from two months old seedling.

Anatomical study of leaf

The young leaves of two month old plants were collected from both IR-29 and O. coarctata for

anatomical studies. Fine sections of leaf were cut with the help of a sharp razor to observe under

light microscope (Nikon Digital Microscopes Eclipse 80i) at magnification of 40X and 100X.

Three biological replicates were taken for analysis of vein density.

For ultrastructural observation, the young leaves of O. coarctata were sectioned to 1 mm² size

and fixed in a fixative (2% glutaraldehyde and 2% paraformaldehyde in 0.1M phosphate buffer

pH-7) for 24h (Giuliani et al., 2013). The leaves sections were then treated with 1% (w/v)

osmium tetroxide followed by standard serial dehydration of acetone for 15 min each and finally

embedded in Spurr's epoxy resin. The ultra thin leaf blocks were made by employing microtome

(Leica, Model: RM2235, Germany) and stained with 4% (w/v) uranyl acetate and 2% (w/v) lead

citrate. For observation of ultrathin blocks, transmission electron microscope (JEOL, Model:

JEM-2100F, Japan) equipped with a Mega View III Digital Camera and Image J Imaging System

was employed.

qReal Time-PCR (qRT-PCR) analysis of C₄ specific genes

To study the expression patterns of C4 specific genes such as carbonic anhydrase (CA), PEPC,

PPDK (pyruvate orthophosphate dikinase), NADP-dependent malic enzyme (NADP-ME) and

malate dehydrogenase (MDH), were selected from previous reports (Supplementary Table 1).

The young leaves of O. coarctata were collected from two months old plants for RNA extracted

by Trizol reagent as per the manufacturer's protocol (Invitrogen, Carlsbad, USA). The integrity

of RNA was checked on formaldehyde- agarose gel (2% w/v). Around 1µg RNA was taken for

cDNA preparation by employing PrimeScript 1st strand cDNA Synthesis Kit (Clonetech Takara

Bio, USA) (Chowrasia et al., 2019). Detail of primers has been given in Supplementary Table 1.

The relative expression level of C4 genes were studied by using qPCR (Light Cycler 480 II,

Roche Molecular Diagnostics, USA) (Chowrasia et al., 2019). The expression level of C4 genes

were normalized with eukaryotic initiation factor 4- α (eIF4- α) as internal control by using 2

^{ΔΔCq} (Cq=threshold cycle) formula (Schmittgen and Livak 2008). Three biological replicates

were taken for expression level analysis.

Statistical analysis

All the differences in mean values of observed characteristics in both rice and O. coarctata were

calculated by Student *t*-test at significance of P = 0.05 and 0.001 (Chowrasia et al., 2019).

Result and discussion

Plants evolved several physiological and biochemical features to encounters different stresses

condition. One such evolution in plants is change of mode of photosynthesis i.e. origin of C4 and

CAM (Crassulacean Acid Metabolism) (Ehleringer and Monson, 1993). C4 mode of

photosynthetic plants are found to be grown in adverse environmental condition with compair to

C3 mode of photosynthetic plants (Sage et al., 1999). The growth and development of C4 plants

were less affected by these harsh environmental condition because of higher rate of

photosynthesis, efficient carbon dioxide fixation, lower rate of photorespiration and transpiration

to check water loss (Bromham and Bennett, 2014). Rice is a C3 plant (Covshoff et al., 2016) and

being in the same genus, we selected a high yielding but highly salt stress sensitive cultivar IR-

29 for comparative study. Further, biomass of *O. coarctata* was found to be more than IR-29

(Fig. 1 A) which might be due to higher rate of photosynthesis in *O. coarctata* compare to IR-29.

O. coarctata is a perennial grass which predominantly propagates vegetatively because seeds are

recalcitrant in nature (Jagtap et al., 2006). The leaves of this wild species of rice are thicker, waxy and leathery (Figure 1B) to prevent transpiration loss.

Further, we studied the anatomical feathers of photosynthetically active two months old leaves of O. coarctata and IR-29. The transverse section of O. coarctata clearly shows Kranz anatomy i.e. wreath or ring like larger bundle sheath cells around the vascular bundle (Figure 1C). The Kranz anatomy is considered to be an exclusive characteristic of C4 plants (Lundgren et al., 2014). The vascular bundle of O. coarctata leaves is surrounded by the large bundle sheath cells which is a typical characteristics of C4 plant (Figure 1C) whereas, IR-29 being C3 plant showed anomalous non-Kranz anatomy (Weerasooriya et al., 2018). Like other C4 plants, bundle sheath cells were in turn surrounded by mesophyll cells. Each pair of vascular bundle is thus separated by two layers of bundle sheath and two layers of mesophyll cells (Figure 1C). The bundle sheath cells of O. coarctata are about 1.4 times larger than rice (Table 1). The leaf of O. coarctata was found to have many furrows and ridges, each ridge possess single vascular bundle. Therefore, the number of vein was also found to be higher in O. coarctata compare to IR-29 (Figure 2 A, B). For further detail study of veins, the leaves of both plant species were sectioned longitudinally. Both transverse (Figure 1C) and longitudinal (Figure 2C) sections of O. coarctata leaf blades showed large number of veins in compare to rice (Figure 2 B, D). In addition, the vein density of leaves was calculated and it was found that the distance between large transverse veins (TV), small longitudinal veins (SLV) and longitudinal veins (LLV) were 1.7, 1.56 and 1.86 times shorter in length in O. coarctata compared to rice (Table 1). This result depicts that large number of veins in leaves of O. coarctata might increases supply of water in leaves to maintain relative water content under saline condition. This increment of vein density in O. coarctata is one of the characteristic feathers of C4 plant (Kumar and Kellogg, 2019; Sage et al., 2004).

The chloroplast dimorphism as well as peripheral localization of chloroplasts and mitrochondria are important characteristics of the C4 plant (Laetsch, 1974). To investigate chloroplast dimorphism and organelles arrangement in leaves O. coarctata, transmission electron microscopy was employed. It has been found that O. coarctata leaves exhibits typical arundinelloid type C4, where chloroplasts were present centrifugally in both mesophyll and bundle sheath cells. The mitochondrial number was higher as observed under TEM photograph in mesophyll cells compare to bundle sheath cells (Figure 3) which suggests slow rate of photorespiration in bundle sheath cells, an important adaptation of C4 plant (Kennedy et al., 1976). The outlines of bundle sheath and mesophyll cell wall were uneven and intercellar space was absent like other C4 plants (Dengler and Nelson, 1999) (Figure 3, 4). The mesophyll chloroplast possess granal stacking whereas, bundle sheath chloroplast lacks grana or contain rudimentary thylakoids (Figure 4, 5). The size of bundle sheath chloroplast was larger than mesophyll cells (Figure 5) which is one of the important criteria to depict that the C4 mode of photosynthesis of O. coarctata. The cell wall of bundle sheath cell was thicker than other mesophyll cells as shown in Figure 5 and suberin lamella was present on the cell wall of bundle sheath cell (Figure 6) which further indicates that O. coarctata possess C4 type of photosynthesis (Dengler and Nelson, 1999). Collectively, these ultrastructure characteristics of O. coarctata leaves revealed that it has adapted C4 mode of photosynthesis to deal with its adverse habitat (Muhaidat et al., 2011; Sage et al., 2004).

In addition, we had also studied the relative expression level of C4 specific genes by RT-qPCR. Five C4 specific genes were selected based upon transcript abundance in C4 plants such as *Echinochloa glabrescens* (Covshoff et al., 2016) and maize (Xu et al., 2016) (Supplementary Table 1). The relative expression level of five C4 related genes i.e. *CA*, *PEPC*, *PPDK*, *NADP*-

ME and MDH were compared in both leaves of Oryza species. The expression level of CA gene which is involved in hydration of carbon dioxide was found to be low in O. coarctata which corroborating with the findings of C4 plants (Xu et al., 2016). The transcript level of primary carboxylation gene of C4 cycle was significantly higher in O. coarctata compare to IR-29. Similarly, expression level of other important C4 related genes, NADP-ME and MDH were also found to be higher in O. coarctata. These two genes are involved in reduction of oxaloacetic acid into malate (Hatch and Slack, 1968). PPDK gene expression was also found to be high in O. coarctata compare to IR-29 rice variety, which involved in synthesis of phosphoenolpyruvate in chloroplast of mesophyll (Figure 7). Therefore, these above features depict that O. coarctata has C4 mode of photosynthesis.

It was mainly found that C4 plants usually grown in warm temperate zone to tropical zone which consist of grassland, salt marshes (Sage et al., 1999). The high irradiance, low level of atmospheric carbon dioxide, high temperature, salinity, low nitrogen availability and drought promote C4 mode of photosynthesis (Osmond et al., 1982). Like other halophytic C4 plants, *O. coarctata* is shade loving and obligate halophyte where sodium is necessary for its growth (Sanchez et al., 2013). Sodium ions play an important role in pyruvate transportation through sodium:proton antiporter in plastid where it metabolized into carbon dioxide thus increasing its level for photosynthesis (Furumoto et al., 2011). Therefore, in order to scope such adverse condition (higher salt concentration E.Ce: 20-40 dSm⁻¹) *O. coarctata* adapted efficient C4 mode of photosynthesis by expressing higher level of C4 specific genes and adapted special anatomical features to fix carbon dioxide.

Therefore, O. coarctata being halophyte has adapted C4 mode of photosynthesis mechanism

among Oryza genus. Thus, introgression of C4 photosynthetic genes into rice by genetic

engineering or by conventional breeding method will make C4 rice for higher yield.

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Legends

Figure 1. Morpho-anatomical study of *O. coarctata*. (A) Comparative observation of IR-29 (*O.*

sativa) and O. coarctata biomass, (B) Vegetatively propagated seedling of O. coarctata and (C)

Transverse section of young leaf of O. coarctata under light microscope (40X magnification).

Scale bar=50µm. The means value of fresh weight (mg) were compared with t-test (** P ≤

0.001).

Figure 2. Images of leaf veins as observed under light microscope, (A) O. coarctata and (B) O.

sativa (IR-29). (C) Longitudinal section of leaf blade of O. coarctata and (D) Rice. LLV-large

longitudinal vein; SLV- small longitudinal vein and TV- transverse vein are indicated. Scale bar

 $=10\mu m$.

Figure 3. O. coarctata bundle sheath cell containing chloroplast and mitochondria. C-chloroplast,

M- mesophyll, BS- bundle sheath and Mi- mitochondria. Scale bar = 500nm.

Figure 4. O. coarctata leaves exhibit chloroplast dimorphism i.e. mesophyll chloroplast exhibit

extensive granal stacking (black arrow) and bundle sheath show no granal stacking under

transmission electron microscope. Scale bar = 500nm.

Figure 5 Ultrastructure of O. coarctata leaves illustrating centrifugal arrangement of the

chloroplast in bundle sheath and mesophyll cells. The chloroplast of bundle sheath is larger than

mesophyll chloroplast. The bundle sheath cell wall is thicker (black arrow). Scale bar = 500nm.

Figure 6. The bundle sheath cell wall showing suberin lamella (white arrow). Scale bar =

500nm. SL- suberin lamella.

Figure 7. The relative expression levels of five C4 genes in rice and O. coarctata as normalized

with eukaryotic initiation factor 4-α. The five C4 genes are carbonic anhydrase (CA),

phosphoenolpyruvate carboxylase (PEPC), malate dehydrogenase (MDH), NADP-dependent

malic enzyme (NADP-ME) and pyruvate orthophosphate dikinase (PPDK). The means value of

relative expression levels were with compared *t*-test (* $P \le 0.05$, ** $P \le 0.001$).

Table1. Detail study of leaf vein density. The value with same lower case are not significantly

different in Student *t*-test at significance of P = 0.05 and 0.001. (P < 0.001).

Supplementary Table1. Gene specific primers of the selected C4 genes used in qRT-PCR.

Table1. Detail study of leaf vein density. The value with same lower case are not significantly different in Student t-test at significance of P = 0.05 and 0.001. (P < 0.001).

Plant	Distance between	Distance	Distance	Area of Bundle
	large longitudinal	between small	between	shealth (µm²)
	veins (µm)	longitudinal	transverse veins	
		veins (µm)	(µm)	
O. sativa	68.05 ^a	14.01 ^a	12.01 ^a	37.25 ^a
O. coarctata	40 ^d	8.94 ^b	6.60 ^b	52.97 ^c

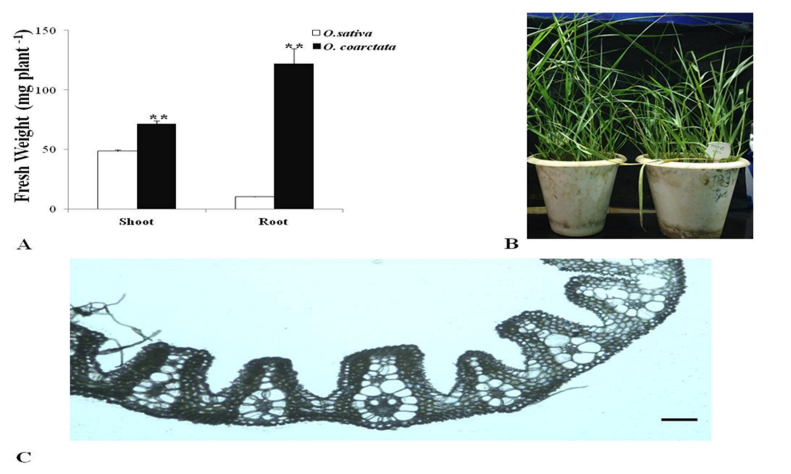


Figure 1. Morpho-anatomical study of O. coarctata. (A) Comparative observation of IR-29 (O. sativa) and O. coarctata biomass, (B) Vegetatively propagated seedling of O. coarctata and (C) Transverse section of young leaf of O. coarctata under light microscope (40X magnification). Scale bar= $50\mu m$. The means value of fresh weight (mg) were compared with t-test (** $P \le 0.001$).

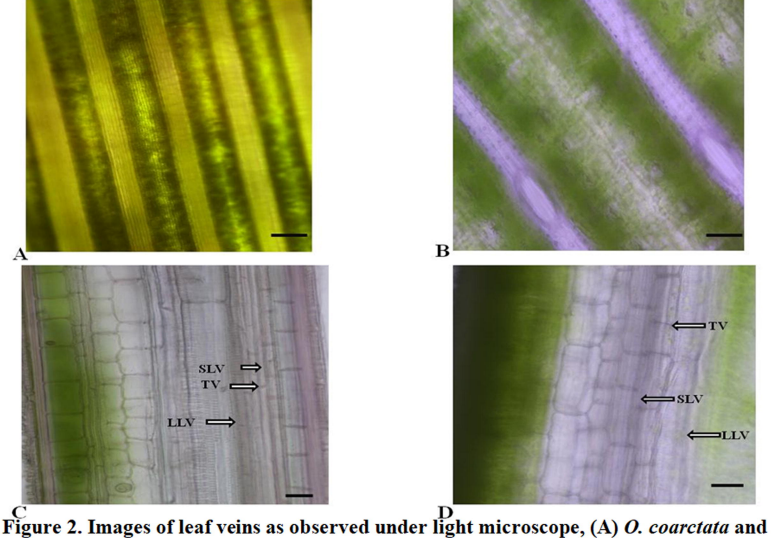
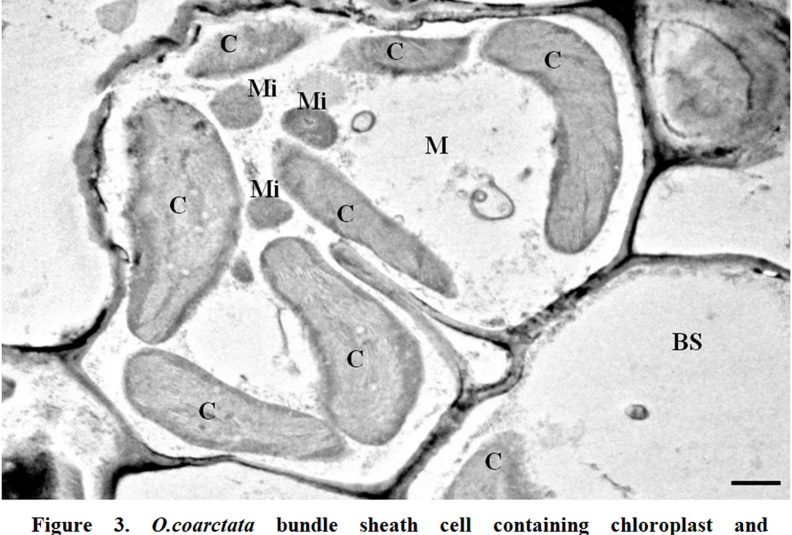


Figure 2. Images of leaf veins as observed under light microscope, (A) O. coarctata and (B) O. sativa (IR-29). (C) Longitudinal section of leaf blade of O. coarctata and (D) Rice. LLV-large longitudinal vein; SLV- small longitudinal vein and TV- transverse vein are indicated. Scale bar =10 μ m.



mitochondria. C-chloroplast, M- mesophyll, BS- bundle sheath and Mi-mitochondria. Scale bar = 500nm.

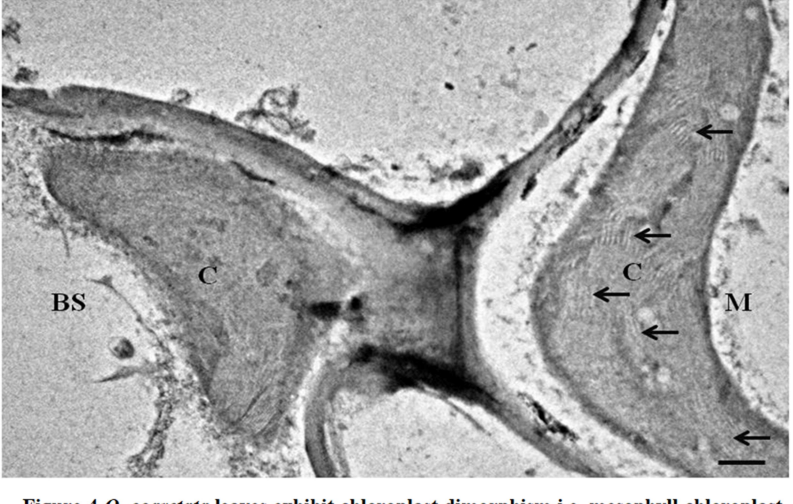


Figure 4 O. coarctata leaves exhibit chloroplast dimorphism i.e. mesophyll chloroplast exhibit extensive granal stacking (black arrow) and bundle sheath show no granal stacking under transmission electron microscope. Scale bar = 500nm.

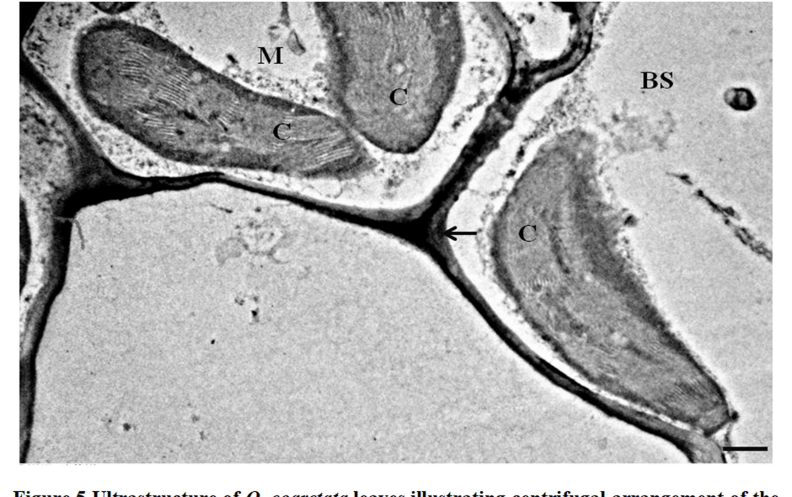


Figure 5 Ultrastructure of *O. coarctata* leaves illustrating centrifugal arrangement of the chloroplast in bundle sheath and mesophyll cells. The chloroplast of bundle sheath is larger than mesophyll chloroplast. The bundle sheath cell wall is thicker (black arrow). Scale bar = 500nm.



Figure 6. The bundle sheath cell wall showing suberin lamella (white arrow). Scale bar = 500nm. SL- suberin lamella.

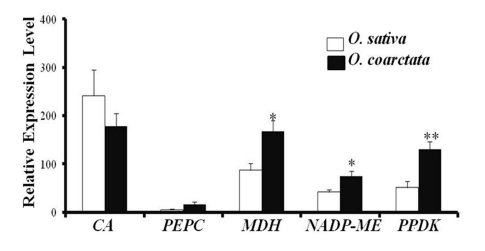


Figure 7. The relative expression levels of five C4 genes in rice and O. coarctata as normalized with eukaryotic initiation factor 4- α . The five C4 genes are carbonic anhydrase (CA), phosphoenolpyruvate carboxylase (PEPC), malate dehydrogenase (MDH), NADP-dependent malic enzyme (NADP-ME) and pyruvate orthophosphate dikinase (PPDK). The means value of relative expression levels were with compared t-test (* P < 0.05, ** P \le 0.001).