SHORT COMMUNICATION

Metabolic interplay between cytosolic phosphoenolpyruvate carboxylase and mitochondrial alternative oxidase in thermogenic skunk cabbage, *Symplocarpus renifolius*

Md. Abu Sayeda, Yui Umekawa, and Kikukatsu Itoa,b

aUnited Graduate School of Agricultural Science, Iwate University, Ueda, Morioka, Iwate, Japan; bCryobiofrontier Research Center, Faculty of Agriculture, Iwate University, Ueda, Morioka, Iwate, Japan

**ABSTRACT**

Skunk cabbage (*Symplocarpus renifolius*) blooms in early spring and its inflorescence, referred to as the spadix, can produce enough heat to melt snow. Here, we investigated glycolytic carbon flow at the PEP branch-point in thermogenic spadices. Our analyses revealed that petals and pistils in thermogenic flores showed higher expression of *SrPEPC* and *SrAOX* transcripts than those of *SrPK*, *SrPCK*, and *SrPEPtase*. Moreover, enzymatic analyses showed high activities of PEPC in the extracts from thermogenic flores. Finally, mitochondria from thermogenic florses showed low respiratory activities when pyruvate was used as a substrate, although a significant malate-mediated cyanide-insensitive respiration was observed. Collectively, these results suggest that PEP metabolism, primarily catabolized by PEPC, plays a critical role in thermogenesis in *S. renifolius*.

**Abbreviations:** AOX, alternative oxidase; BSA, bovine serum albumin; EDTA, ethylenediaminetetraacetic acid; EF1α, elongation factor 1α; MOPS, 3-morpholinopropanesulfonic acid; MPCs, mitochondrial pyruvate carriers; NADH, nicotinamide adenine dinucleotide; PCR, polymerase chain reaction; PEP, phosphoenolpyruvate; PEPC, phosphoenolpyruvate carboxylase; PEPCK, phosphoenolpyruvate carboxykinase; PEPtrase, PEP phosphatase; PK, pyruvate kinase; qRT-PCR, quantitative real-time PCR; TPP, thiamine pyrophosphate

**Introduction**

Glycolysis is a central pathway of carbohydrate metabolism among living organisms from bacteria to plants and humans.1,2,3 In the glycolytic pathway, phosphoenolpyruvate (PEP) is an important intermediate, as it occupies the highest position on the thermodynamic scale of known phosphorylated metabolites.4 In animal cells, PEP is predominantly catalyzed by pyruvate kinase (PK; EC 2.7.1.40) with the concomitant phosphorylation of ADP to ATP.3 However, in addition to PK, PEP carboxylase (PEPC; EC 4.1.1.31) and PEP phosphatase (PEPtrase; EC 3.1.3.60) play roles in the catabolism of PEP and overall regulation of mitochondrial respiration in plant cells.2,5 In contrast, PEP carboxykinase (PEPCK; EC 4.1.1.49) is an enzyme involved in gluconeogenesis.5

Although thermogenesis is uncommon in plants, the glycolytic pathway is important in thermogenic plants such as skunk cabbage (*Symplocarpus renifolius*), which utilize carbohydrates as a major respiratory substrate for their metabolic heat-production.6 In the case of *S. renifolius*, organ-specific thermogenesis occurs in the inflorescence known as the spadix, and maintains a temperature of approximately 23°C during flowering, even when ambient temperatures drop below freezing.7,8,9 Because the mitochondrial cyanide-insensitive alternative oxidase (AOX) allows for a dramatic decrease in free energy between ubiquinol and oxygen,10,11 *SrAOX* identified in *S. renifolius* is likely to play a role in thermogenesis in this plant.7,12,13 In our previous study, sustained thermogenesis in spadices was associated with the import of carbohydrates including sucrose, glucose, and fructose from roots.14 Although these results suggest that glycolysis and subsequent AOX-mediated mitochondrial respiration plays a crucial role in organ-specific thermogenesis in *S. renifolius*, the mechanisms of carbohydrate metabolism in this plant remain poorly understood.

The purpose of the present study was to investigate glycolytic carbon flow at the PEP branch-point in thermogenic spadices of *S. renifolius*. Our data showed that co-expression of PEPC and AOX is central for metabolic heat-production in this plant.

**Results**

**Gene expression analyses for PK, PEPtrase, PEPC, and PEPCK in thermogenic and non-thermogenic tissues**

Tissue-specific expression patterns of *SrPK*, *SrPEPtrase*, *SrPEPC*, and *SrPepck* transcripts were determined by qRT-PCR using...
that expression levels of the SrPEPC and SrAOX transcripts were significantly higher in florets than those in the non-thermogenic spathe, leaf, and pith tissues (Fig. 1b). In contrast, the expression levels of SrPK, SrPEPtase, and SrPEPCK were nearly undetectable in the spathe, leaf, floret, and pith tissues (Fig. 1b).

Next, to examine gene expression in more detail, we focused on 3 tissues, the petal, stamen, and pistil, each of which comprises the florets of the spadix (Fig. 1a). SrPEPC transcripts were again co-expressed with SrAOX transcripts in petals and pistils, whereas SrPK, SrPEPtase, and SrPEPCK transcripts were expressed at low levels in all tissues examined (Fig. 1c). Expression levels of SrPK, SrPEPtase, SrPEPC, SrPEPCK, and SrAOX were also determined at 2 thermogenic stages of plants collected on the same day, and showed considerably higher expression of SrPEPC and SrAOX transcripts in the floret, petal, and pistil tissues (Figs. S1 and S2).

**Enzyme assays for PK, PEPtase, PEPC, and PEPC**

To determine whether similar high enzymatic activities of PEPC are found in thermogenic florets, we determined the PK, PEPtase, PEPC, and PEPC activities in cytosolic fractions from thermogenic florets. These analyses showed that the specific activity of PEPC was significantly higher than that of PK, PEPtase, and PEPC (Table 1). Another independent preparation from different thermogenic plants also showed similar results (Supplementary Table 1).

**Predicted amino acid sequence of SrPEPC**

Phylogenetic analysis of PEPC amino acid sequences from 15 plants, 3 algae, and 4 bacteria clearly characterized SrPEPC as a C3-plant-type PEPC (Fig. 2a). Moreover, amino acid sequences of SrPEPC and the homolog NnPEPC from the thermogenic plant species *Nelumbo nucifera* were closely related on the phylogenetic tree (Fig. 2a).

Multiple sequence alignments of the deduced PEPC amino acid sequence with other protein sequences from C3- and C4-plant PEPCs showed that SrPEPC possesses conserved alanine and arginine residues, characteristic of C3-plant-type PEPCs (Fig. 2b). Moreover, a serine residue with a potential phosphorylation site was highly conserved across C3- and C4-types of plant PEPCs, including that from *S. renifolius*.

![Figure 1. Levels of SrPK, SrPEPtase, SrPEPC, SrPEPCK, and SrAOX transcripts in various tissues during thermogenesis in *S. renifolius*. (a) Infrared thermal imaging of longitudinal section of the spadix. The positions of florets, pith, and spathe are shown. Each floret composed of the stamen, pistil, and petal is also illustrated. Expression profiles of SrPK, SrPEPtase, SrPEPC, SrPEPCK, and SrAOX transcripts in the (b) spathe, leaf, floret, and pith and (c) petal, stamen, and pistil. SrEF1α transcripts were used as a normalization control. Experiments were performed in triplicate for each sample. Data are expressed as the mean ± standard deviation. Values with different letters in the graph indicate that they are statistically significantly different (n = 3; P < 0.05). AOX, alternative oxidase; PK, pyruvate kinase; PEP, phosphoenolpyruvate; PEPtase, PEP phosphatase; PEPC, PEP carboxylase; PEPC, PEP carboxykinase.](image)

| Enzyme | Specific activity (nmol min⁻¹ mg⁻¹ protein) |
|--------|------------------------------------------|
| PK     | 9.5 ± 2.1*                                |
| PEPtase| 10.2 ± 4.3*                               |
| PEPC   | 107.5 ± 13.7*                             |
| PEPC   | 14.8 ± 5.7*                               |

Value for PEPC is depicted as decarboxylation activity. Assays were performed in triplicate for each sample. Data are expressed as mean ± standard deviations. Values with different letters indicate that they are statistically significantly different (n = 3; P < 0.05). PK, pyruvate kinase; PEP, phosphoenolpyruvate; PEPtase, PEP phosphatase; PEPC, PEP carboxylase; PEPC, PEP carboxykinase.

**Table 1. Enzyme activities of PK, PEPtase, PEPC, and PEPC in cytosolic fraction of thermogenic florets of *S. renifolius*.**

RNAs from the spathe, leaf, floret, and pith tissues collected from *S. renifolius* plants during thermogenesis (Fig. 1). In these experiments, temperatures of ambient air and thermogenic spadix were 12.2°C and 22.3°C, respectively. Our previous study showed that florets are thermogenic, while the spathe, leaf, and pith are non-thermogenic. Thermogenic florets contain the stamen, pistil, and petal (Fig. 1a). SrAOX mRNA expression was used as a thermogenic tissue-specific control to reflect the thermogenic status of the samples. Our data clearly showed
Substrate-dependent oxygen uptake

The present results indicate that pyruvate, which is formed from PEP by PK and/or PEPtase, is not a major end-product of the glycolytic pathway in thermogenic cells of *S. renifolius*. Thus, we next wished to clarify whether purified mitochondria from thermogenic florets oxidize pyruvate as a substrate for respiration. In this analysis, we used the same experimental conditions at pH 7.2 as previously reported for pyruvate-mediated respiration in the mitochondria from thermogenic appendices of *Arum maculatum* (Fig. 3a and b). In these experiments, we used the cinnamate derivative UK5099 to inhibit mitochondrial pyruvate transport and determined the effects on respiratory rates (Fig. 3b). Our results showed that pyruvate does not act as a substrate for mitochondrial respiration and that UK5099 does not affect respiration. However, subsequent addition of NADH led to significantly increased respiration rates both in absence and in presence of cyanide, indicating cyanide-insensitive mitochondrial AOX activities (Fig. 3a and b).

To confirm that our purified mitochondria from thermogenic florets do not utilize externally added pyruvate as a substrate for respiration, we performed experiments at pH 6.8, 7.2, and 7.6 in presence of the pyruvate dehydrogenase complex cofactors NAD⁺ and thiamine pyrophosphate (TPP). These experiments indicated that pyruvate was not used as a substrate.

Figure 2. Phylogenetic tree and amino acid alignments of PEPC proteins. (a) Phylogenetic tree of deduced amino acid sequences of PEPC from *S. renifolius*, other C3- and C4-plants, and bacterial and algal type organisms. (b) Positions of HCO₂⁻-binding loop (red arrow) and inhibitor binding site (blue arrow). (c) Conserved potential phosphorylation sites (black arrow) at the N-terminal region of PEPC. PEPC, phosphoenolpyruvate carboxylase.
respiratory substrate (Fig. 3c–e). However, addition of exogenous malate led to increased respiration rates regardless of whether cyanide was present, with the highest respiration rate observed at pH 7.2 (Fig. 3c–e).

Discussion

PEP is an energy-rich compound and its metabolism in replacement of phosphate group plays an important role in glycolysis. Therefore, analysis of the metabolic partitioning of PEP will provide insight into the glycolytic end products and energy substrates used in subsequent mitochondrial respiration.

Herein, we showed that transcripts encoding the C3-type of PEPC protein SrPEPC were specifically and highly co-expressed with those encoding SrAOX in the petals and pistils of thermogenic florets, whereas the expression levels of SrPK, SrPEPtase, and SrPEPCK transcripts were extremely low in all tissues examined (Fig. 1). Moreover, enzymatic activities of PEPC in thermogenic tissues were higher than those of PK, PEPtase, and PEPCK (Table 1), indicating that PEP is predominantly catabolized by PEPC in thermogenic tissues of *S. renifolius*. Although extremely
high enzymatic activities of PEPC have been reported in other thermogenic plants including A. maculatum,20 this is the first study to show tissue-specific co-expression of PEPC and AOX in quantitative gene expression analyses of thermogenic plants.

Because the AOX-mediated energy-dissipative respiration pathway contributes significantly to cellular thermogenesis in plants, co-expression of SrAOX and SrPEPC detected in the present study may be critical for metabolic cross-talk between the cytosol and AOX-expressing mitochondria in thermogenic cells (Fig. 4). In thermogenic cells, PEP was primarily catabolized by PEPC to produce oxaloacetic acid, which is used directly as a mitochondrial respiration substrate or is converted to malate by malate dehydrogenase21,22 for use as a respiratory substrate (Fig. 3c–e). In either case, such PEPC-mediated metabolism may contribute significantly to continuous carbon flow in furnishing C4-dicarboxylic acids that maintains increased respiration for thermogenesis in S. renifolius. These data are consistent with those of previous report of constitutive PEPC-overexpressing transgenic plants, in which carbon flow was redirected from soluble sugars to organic acids.23 More importantly, because PEPC catalyzes the addition of HCO$_3^-$ to PEP,4 excess CO$_2$ that is liberated with increased mitochondrial respiration in thermogenic cells may be catabolized by complex I-integrated mitochondrial γ-carboxylic anhydrases to form HCO$_3^-$,24,25 which is subsequently converted to oxaloacetic acid by PEPC. Previously, it was shown that PEPC is highly expressed and participates in the recycling of respired CO$_2$ in the spikelets of C3-type plants.26 Similarly, fruiting plants possess a system known as fruit photosynthesis,27 in which CO$_2$ from mitochondrial respiration is refixed by PEPC. These data indicate that thermogenic plants express higher levels of PEPC and AOX enzymes in their non-photosynthetic organs, such as in the spadices of S. renifolius which developed specialized metabolisms not only for recycling of excess CO$_2$ similar to that seen in other C3 plants but also for energy-dissipating AOX-mediated mitochondrial respiration during evolution. Accordingly, integration of PEPC-mediated CO$_2$ assimilation and AOX-mediated mitochondrial respiration probably act as substantial carbon resources in thermogenic plants.

In the present study, mitochondria purified from thermogenic florets of S. renifolius did not oxidize exogenous pyruvate as a respiratory substrate in the presence or absence of cofactors (Fig. 3). In contrast, mitochondria from thermogenic spadices of A. maculatum were previously shown to oxidize exogenous pyruvate in the absence of exogenous cofactors.18 In addition, rapid oxidation of pyruvate by isolated A. maculatum mitochondria was sensitive to UK5099, suggesting the presence of mitochondrial pyruvate carriers, as shown recently in Arabidopsis.28 Hence, expression of mitochondrial pyruvate carriers in thermogenic florets of S. renifolius may be lower than in those of A. maculatum. Because pyruvate has been identified as an allosteric activator of S. renifolius AOX,12 AOX activities in this plant may be post-translationally regulated by intra-mitochondrially produced pyruvate via malic enzyme29,30 that uses malate from the PEPC-mediated metabolic pathway. It should be noted here that NAD-dependent malic enzyme α- and/or β-subunits have been identified in mitochondria from thermogenic florets of S. renifolius.13

In conclusion, we report that SrPEPC is abundantly co-expressed with SrAOX in thermogenic florets of S. renifolius. These results further suggest that PEPC plays a role in metabolic heat-production in furnishing C4-dicarboxylic acids to AOX-expressing mitochondria in other thermogenic plants.

**Materials and methods**

**Plant materials**

All plant materials were sampled from wild S. renifolius grown outdoors. For thermal imaging, S. renifolius plants that were transplanted from Hakuba (Nagano prefecture, Japan) to Iwate University campus (Iwate prefecture, Japan) in April 2005 were used. For preparation of total RNAs, fresh spathe, leaf, floret and pith were collected from S. renifolius at Fujine (Iwate prefecture, Japan) on April 3, 2012 and at Omori (Akita prefecture, Japan) on April 1, 2014. Mitochondria were purified from spadices of S. renifolius sampled at Omori on March 27, 2015. For enzyme assays, florets were collected from thermogenic spadices of S. renifolius that were sampled at Omori on April 20, 2016.
Thermal imaging and temperature measurements

Thermal images were obtained using an infrared thermal camera as described previously. Temperatures of the spadices were measured using an automatic recording thermometer (TR-52; T & D, Nagano, Japan).

Total RNA extraction, cDNA amplification, and isolation of full-length cDNAs encoding SrPK, SrPEPtase, SrPEPC, and SrPEPCK

Total RNAs were extracted using either an RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) or a FastPure® RNA Kit (Takara Bio, Shiga, Japan). First-strand cDNA synthesis was performed using PrimerScript™ II 1st strand cDNA Synthesis Kit (Takara Bio) with oligo-dT primers provided by the manufacturer. Procedures for cDNA cloning are described in the Supplementary information. Briefly, partial fragments of targeted genes for SrPK, SrPEPtase, SrPEPC, and SrPEPCK were first amplified using PCR with Takara Ex Taq® (Takara Bio) and the primers are listed in Supplementary Table 2. Gene-specific primers were then designed to perform 5'- and 3'-rapid amplification of cDNA ends using SMARTer™ RACE cDNA Amplification Kit (Clontech, Mountain View, CA, USA). To isolate full-length cDNAs, final PCR amplifications were performed with the KOD -Plus- (Toyobo, Osaka, Japan) with the primers shown in Supplementary Table 2. The obtained fragments were cloned into the T-Vector (pMD19, Takara Bio) and sequenced in both directions. Full-length sequences were determined by isolating at least 2 independent clones with identical sequences. DNA sequences were analyzed using GENETYX software (Genetyx, Tokyo, Japan). Complete cDNAs encoding SrPEPC, SrPK, SrPEPCK, and SrPEPtase were deposited in the DNA Data Bank of Japan with Accession numbers LC155943, LC155944, LC155945, and LC155946, respectively.

Expression analyses of genes encoding SrPK, SrPEPtase, SrPEPC, and SrPEPCK

Real-time qPCR was performed using a Thermal Cycler Dice (TP800; Takara Bio) instrument as described previously. Gene-specific primers (Supplementary Table 2) were designed from identified cDNAs and the housekeeping gene EF1α was used as a normalization control.

Phylogenetic tree analyses of PEPC proteins and sequence alignments

A phylogenetic tree was constructed using the neighbor-joining method with ClustalW for 22 PEPC proteins gathered from GeneBank. A bootstrap consensus tree was inferred from 1000 replicates to represent the evolutionary history of the present taxa. Branches corresponding to partitions that were reproduced in less than 50% of bootstrap replicates were collapsed. Multiple sequence alignments were performed using GENETYX software.

Enzyme assays

Enzyme activities of PK, PEPtase, PEPC, and PECK were determined in thermogenic florets after extraction in ice-cold extraction buffer containing 0.3 M mannitol, 20 mM MOPS (pH 7.5), 2 mM EDTA, 2 mM pyruvate, 7 mM cysteine, and 0.2% BSA. Extracts were filtered through 8 layers of Miracloth (EMD Millipore, Billerica, MA, USA). Filtrates were collected in 50-mL tubes and centrifuged at 120 g for 10 min at 4°C. Supernatants were collected and centrifuged again at 12,000 g for 10 min at 4°C, and stored at −80°C until enzymatic analyses. Enzyme assays of PK and PE Ptase and assays of PEPC and PE CK for decarboxylation were conducted as previously described with a double beam spectrophotometer (Bio-spec-1600, Shimadzu, Kyoto, Japan) at 25°C.

Isolation of intact mitochondria and respiration analyses

Mitochondria were isolated from S. renifolius spadices as described previously. Oxygen uptake by mitochondria was then measured according to our previous reports at 25°C.

Determination of protein concentrations

Protein concentrations of isolated mitochondria and crude extracts were determined as described previously.

Statistical analysis

All data were compared using one-way factorial ANOVA (SigmaPlot 12, Systat Software, San Jose, CA, USA). Tukey’s honest significance posthoc tests were used to identify significantly different means. Significant differences between means were calculated at P = 0.05.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

Acknowledgments

The authors are grateful to members of our laboratory for technical assistance and helpful discussions, and to Dr. M. Kasai, Dr. Y. Saitoh, and Dr. M. Afzal Hossain for their valuable discussions. We would like to thank Dr. Y. Onda for sharing data of thermal imaging. This study is dedicated to Professor S. Kawai who passed away on March 10, 2016.

Funding

This work was supported in part by JSPS KAKENHI (Grant Numbers JP16H05064 and JP15KT0101).

References

1. Audesirk T, Audesirk G, Byers BE. Biology: Life on Earth, eighth ed., Upper Saddle River, NJ: Pearson; 2008; ISBN 13: 9780132380614
2. Plaxton WC. The organization and regulation of plant glycolysis. Annu Rev Plant Physiol Plant Mol Biol 1996; 47:185-214; PMID:15012287; http://dx.doi.org/10.1146/annurev.arplant.47.1.185
19. O'Leary B, Fark J, Plaxton WC. The remarkable diversity of plant PEPC (phosphoenolpyruvate carboxylase): recent insights into the physiological functions and post-translational controls of non-photo synthesizing PEPCs. Biochem J 2011; 436:15-34; PMID:21524275; http://dx.doi.org/10.1042/Bj20110078

20. Seymour RS, Blaylock AJ. Switching off the heater: influence of ambient temperature on thermoregulation by eastern skunk cabbage Symplocarpus foetidus. J Exp Bot 1999; 50:1525-32; http://dx.doi.org/10.1093/sxb/50.338.1525

21. Onda Y, Kato Y, Abe Y, Ito T, Morohashi M, Ito Y, Ichikawa M, Matsukawa K, Kakizaki Y, Koita H, Ito K. Functional coexpression of the mitochondrial alternative oxidase and uncoupling protein underlies thermoregulation in the thermogenic florets of skunk cabbage. Plant Physiol 2008; 146:636-45; PMID:18162588; http://dx.doi.org/10.1104/pp.107.113563

22. Seymour RS, Ito Y, Onda Y, Ito K. Effects of floral thermogenesis on pollen function in Asian skunk cabbage Symplocarpus renifolius. Bioll Lett 2009; 5:568-70; PMID:19364718; http://dx.doi.org/10.1098/rsbl.2009.0064

23. Knutson RM. Heat production and temperature regulation in eastern skunk cabbage. Science 1974; 186:746-7; PMID:4417289; http://dx.doi.org/10.1126/science.186.4165.746

24. Moore AL, Shiba T, Young L, Harada S, Kita K, Ito K. Unraveling the heater: new insights into the structure of the alternative oxidase. Annu Rev Plant Biol 2013; 64:637-63; PMID:23638828; http://dx.doi.org/10.1146/annurev-arplant-042811-105432

25. Moore AL, Bonner WD. Measurements of membrane potentials in plant mitochondria with the safranine method. Plant Physiol 1982; 70:1271-6; PMID:6662666; http://dx.doi.org/10.1104/pp.70.5.1271

26. Onda Y, Kato Y, Abe Y, Ito T, Inaba Y, Morohashi M, Ito Y, Ichikawa M, Matsukawa K, Otsuka M, Koita H, Ito K. Pyruvate-sensitive AOX exists as a non-covalently associated dimer in the homeothermic spadix of the skunk cabbage, Symplocarpus renifolius. FEBs Lett 2007; 58:5852-8; PMID:18060878; http://dx.doi.org/10.1016/j.febslet.2007.11.061

27. Kakizaki Y, Moore AL, Ito K. Different molecular bases underlie the mitochondrial respiratory activity in the homeothermic spadices of Symplocarpus renifolius and the transiently thermogenic appendices of Arum maculatum. Biochom J 2012; 445: 237-46; PMID:22512685; http://dx.doi.org/10.1042/Bj201101978

28. Onda Y, Ito K. Changes in the composition of xylem sap during development of the spadix of skunk cabbage (Symplocarpus foetidus). Biosci Biotechnol Biochem 2005; 69:1156-61; PMID:15973047; http://dx.doi.org/10.1271/bbb.69.1156

29. Ming R, Vanburen R, Liu Y, Yang M, Han Y, Li LT, Zhang Q, Kim MJ, Schatz MC, Campbell M, et al. Genome of the long-living sacred lotus (Nelumbo nucifera Gaertn.). Genome Biol 2013; 14; R41; PMID:23663246; http://dx.doi.org/10.1186/gb-2013-14-5-r41

30. Paulus JK, Schleper D, Groth G. Greater efficiency of photosynthetic carbon fixation due to single amino-acid substitution. Nat Commun 2013; 4; 1518; PMID:23243304; http://dx.doi.org/10.1038/ncomms2504

31. Proudflove MO, Beechey RB, Moore AL. Pyruvate transport by thermogenic-tissue mitochondria. Biochem J 1987; 247:441-447; PMID:3426546; http://dx.doi.org/10.1042/bj2470441

32. Proudflove MO, Moore AL. Metabolite fluxes across the inner membrane of plant mitochondria - inhibition by phthalic acid. Planta 1984; 160:407-14; PMID:24258667; http://dx.doi.org/10.1007/BF00492756

33. Patel MS, Nemeria NS, Furey W, Jordan F. The pyruvate dehydrogenase complexes: structure-based function and regulation. J Biol Chem 2014; 289:16615-23; PMID:24798336; http://dx.doi.org/10.1074/jbc.R114.563148

34. Rees T, Fuller WA, Green JH. Extremely high activities of phosphoenolpyruvate carboxylase in thermogenic tissues of Araceae. Planta 1981; 152:79-86; PMID:24303223; http://dx.doi.org/10.1007/BF00384989

35. Eron Y, Hata S, Izui K. Regulatory phosphorylation of plant phosphoenolpyruvate carboxylase: recent insights into the physiological function and regulation. J Biol Chem 2014; 289:16078-84; PMID:24824257; hydroxylation on phosphoenolpyruvate carboxylase, and the Krebs cycle to respiration and biosynthesis and to intracellular pH regulation during hypoxia in maize root tips observed by nuclear magnetic resonance imaging and gas chromatography-mass spectrometry. Plant Physiol 1998; 116:1073-81; PMID:9501140; http://dx.doi.org/10.1104/pp.116.3.1073

36. Ito K, Ogata T, Kakizaki Y, Elliott C, Albury MS, Moore AL. Identification of a gene for pyruvate-insensitive mitochondrial alternative oxidase expressed in the thermogenic appendices in Arum maculatum. Plant Physiol 2011; 157:1721-32; PMID:21988877; http://dx.doi.org/10.1104/pp.111.186932

37. Thompson JD, Higgins DG, Gibson TJ. CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nuclic Acids Res 1994; 22:4673-80; PMID:7984417; http://dx.doi.org/10.1093/nar/22.22.4673

38. Ireland RJ, De Luca V, Dennis DT. Characterization and kinetics of isoenzymes of pyruvate kinase from developing castor bean endosperm. Plant Physiol 1980; 65:1188-93; http://dx.doi.org/10.1104/pp.65.6.1188

39. Ueno Y, Hata S, Izui K. Regulatory phosphorylation of plant phosphoenolpyruvate carboxylase: role of a conserved basic residue upstream of the phosphorylation site. FEBS Lett 1997; 417:57-60; PMID:9395074; http://dx.doi.org/10.1016/S0014-5793(97)01254-4

40. Walker RP, Chen ZH, Acheson RM, Leegood RC. Effects of phosphorylation on phosphoenolpyruvate carboxykinase from the C4 plant Guinea grass. Plant Physiol 2002; 128:165-72; PMID:11788762; http://dx.doi.org/10.1104/pp.101632

41. Umezawa Y, Seymour RS, Ito K. The biochemical basis for thermoregulation in heat-producing flowers. Sci Rep 2016; 6:24830; PMID:27095582; http://dx.doi.org/10.1038/srep24830