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Proline oxidation fuels mitochondrial respiration during dark-induced leaf senescence in Arabidopsis thaliana

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Abstract

Leaf senescence is a form of developmentally programmed cell death that allows the remobilization of nutrients and cellular materials from leaves to sink tissues and organs. Among the catabolic reactions that occur upon senescence, little is known about the role of proline catabolism. In this study, the involvement in dark-induced senescence of proline dehydrogenases (ProDHs), which catalyse the first and rate-limiting step of proline oxidation in mitochondria, was investigated using prodh single- and double-mutants with the help of biochemical, proteomic, and metabolomic approaches. The presence of ProDH2 in mitochondria was confirmed by mass spectrometry and immunogold labeling in dark-induced leaves of Arabidopsis. The prodh1 prodh2 mutant exhibited enhanced levels of most tricarboxylic acid cycle intermediates and free amino acids, demonstrating a role of ProDH in mitochondrial metabolism. We also found evidence of the involvement and the importance of ProDH in respiration, with proline as an alternative substrate, and in remobilization of proline during senescence to generate glutamate and energy that can then be exported to sink tissues and organs.

Keywords: Arabidopsis thaliana, dark-induced leaf senescence, mitochondria, primary metabolism, proline dehydrogenase, proline metabolism.

Introduction

Leaf senescence is the final stage of leaf development and is critical for converting cellular materials into exportable nutrients that can be transported to sink tissues and organs (Lim et al., 2007). Senescence starts with programmed dismantling of chloroplasts, which leads to a general coordinated remodelling of cellular metabolism. In order to fuel the energetic demand of cells after the loss of the photosynthetically active chloroplasts, mitochondria persist long after chloroplast degradation.
Mitochondria are involved in oxidative energy metabolism and release energy to allow the maintenance of primary metabolism prior to cell death. Mitochondria numbers have been shown to decrease by 30% during developmental leaf senescence, although the integrity of the remaining mitochondria is preserved (Keech et al., 2007; Chrobok et al., 2016). It has also been postulated that mitochondria acquire additional functions, such as selective catabolism of lipids and amino acids, that compensate the loss of energy supplied by chloroplasts and maintain cell viability until the completion of nutrient remobilization from the cell (Keech et al., 2007).

Glutamate, which is released from several mitochondrial catabolic pathways, is proposed to play a crucial role in nitrogen remobilization during leaf senescence (Chrobok et al., 2016). It is one of the main amino acids present in phloem sap in senescing Arabidopsis (Masclaux-Daubresse et al., 2010) and in poplar (Couturier et al., 2010). Glutamate may originate from proline catabolism and from conversion through arginine activity of arginine to ornithine, which in turn enters in the proline catabolic pathway (Funck et al., 2008). Taken together, although little is known about its role at this developmental stage, these findings suggest that proline catabolism could play a role in recycling proline and arginine during leaf senescence. Proline accumulation has been described in excised rice leaves during dark-induced senescence (Wang et al., 1982; Mondal et al., 1985), although a decrease in proline content has been reported during developmental leaf senescence in Arabidopsis (Chrobok et al., 2016).

Energy deprivation triggers dramatic reprogramming of transcription, which supports metabolic adjustment. Proline metabolism and its regulation have been the subject of numerous studies in plants (Szabados and Savouré, 2010; Sharma et al., 2011; Servet et al., 2012). Anabolism and catabolism are each composed of two steps and take place in different cellular compartments. In response to environmental stresses such as drought and salinity, proline is synthesized from glutamate in the cytosol and most probably in chloroplasts via the activities of bifunctional pyrroline-5-carboxylate synthetase (P5CS) and P5C reductase (P5CR) (Székely et al., 2008). Upon recovery from stress, proline is rapidly oxidized to glutamate in mitochondria via sequential action of proline dehydrogenase (ProDH) and P5C dehydrogenase (P5CDH). In Arabidopsis, proline oxidation occurs via two non-redundant ProDH isoforms (Funck et al., 2010). ProDH1 and ProDH2 share high sequence homology, with 75% identical amino acids (Servet et al., 2012). ProDH1 expression has been shown to be repressed during water stress but is induced by rehydration (Kiyosue et al., 1996; Verbruggen et al., 1996). In addition to its developmental regulation, ProDH1 expression appears to be triggered by hypo-osmolality and exogenously applied proline (Verbruggen et al., 1996; Nakashima et al., 1998). Expression of ProDH1 in response to proline or hypo-osmolality is controlled by a proline-responsive element (PRE) cis-acting element via direct binding of the S1-bZIP transcriptional activator to its promoter (Satoh et al., 2002, 2004; Weltevreden et al., 2006). In addition bZIP1, bZIP2, and bZIP53 have been shown to regulate ProDH1 transcript levels and proline content during dark-induced starvation (Dietrich et al., 2011; Pedrotti et al., 2018). Transcriptional analysis of ProDH1 indicates abundant expression in flowers, particularly in pollen and stigma (Funck et al., 2010). In contrast with ProDH1, low expression is generally observed for ProDH2 with the exception of vascular tissues and the abscission zones of floral organs and senescent leaves, where high expression levels have been recorded (Funck et al., 2010; Faës et al., 2015).

ProDH1 subcellular localization has unambiguously been shown to be mitochondrial (Mani et al., 2002; Schertl and Braun, 2014; Cabassa-Hourton et al., 2016). In response to proline treatment, ProDH1 has been shown to be part of low molecular mass complexes in mitochondria and to play an essential role in mitochondrial proline oxidation, which may provide reducing power, energy, and nitrogen sources for growth (Cabassa-Hourton et al., 2016). The function of ProDH2 was addressed later because its gene had for a long time been considered to be a pseudo gene. The use of ProDH2–GFP fusion proteins suggests either mitochondrial or plastid localization (Van Aken et al., 2009; Funck et al., 2010). Arabidopsis ProDH2 has been shown to complement a yeast Δput1 mutant and, when overexpressed in a GFP-tagged form, to rescue an Arabidopsis prodh1 mutant using proline as sole source of nitrogen (Funck et al., 2010), demonstrating that ProDH2 as well as ProDH1 can mediate proline oxidation. However, in Arabidopsis supplied with an excess of proline and other nitrogen sources, ProDH2 is neither able to overcome the proline sensitivity of the prodh1 mutant (Funck et al., 2010; Cabassa-Hourton et al., 2016) nor to restore the lack of ProDH1 in mitochondrial proline oxidation (Cabassa-Hourton et al., 2016). Therefore the two ProDHs appear to have distinct and non-redundant physiological functions in proline oxidation, at least in Arabidopsis (Funck et al., 2010; Rizzi et al., 2017; Cabassa-Hourton et al., 2016).

All of these data suggest that remobilization of proline and its oxidized derivatives could participate in the senescence process; however, the role of ProDHs remains to be demonstrated. Therefore, in this study we investigated the contribution of ProDHs during dark-induced leaf senescence in Arabidopsis. Our analyses of the prodh1 prodh2 double–mutant in comparison to the wild-type revealed an important function of ProDHs in proline oxidation and its contribution to respiration during senescence. Distinct metabolome patterns demonstrated a link between the tricarboxylic acid cycle and ProDH, demonstrating a key role of proline oxidation for providing energy and glutamate during senescence.

Materials and methods

Plant growth and senescence treatment

All the Arabidopsis italiana lines tested were in the Columbia-0 (Col-0) background. The prodh1–4 (SALK_119334) and prodh2–2 (GABI_328G05) insertion mutants were obtained from the Salk Institute, La Jolla, CA, USA (Alonso and Stepanova, 2003) and from the Center for Biotechnology, Universität Bielefeld, Germany (Kleinboelting et al., 2012), respectively, and genetically characterized (Cabassa-Hourton et al., 2016). A double-mutant prodh1–4 prodh2–2 (hereafter called prodh1 prodh2) was obtained by crossing prodh1–4 and prodh2–2 as described by Cabassa-Hourton et al. (2016). Seedlings were sown and grown in soil under a 16/8-h light/dark cycle at 80–100 μmol photons m−2 s−1 at 21 °C for 4 weeks.

Senescence experiments were carried out on excised leaves that were left to age in the dark. Leaves were collected from 4-week-old plants and placed on wet Whatman® paper in Petri dishes. The dishes were then
wrapped in aluminum foil and kept in the growth chamber until further analysis. For mitochondrial respiration, ProDH activity, transcript and metabolomic analyses, leaves 7, 8, and 9 from the base of the plant were harvested from 4-week-old plants and were processed as indicated above to trigger dark-induced senescence.

**Immunocytochemical studies**

For electron microscopy, 2×5-mm pieces of leaves were cut and placed in a fixation solution containing 3% (v/v) formaldehyde and 0.5% (v/v) glutaraldehyde in 1× PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, pH 7.4). The samples were then subjected to a low vacuum for 20 min, and washed twice in PBS and once in distilled water for 20 min each. A dehydration procedure was then performed with 25% and 50% ethanol successively for 20 min each under gentle shaking. The samples were then stored in 70% ethanol overnight at 4 °C. Dehydration was achieved by successive incubation in 80% and 90% ethanol for 1 h each. Embedding was started in a mix of 25% London Resin White (LR) in 90% ethanol for 1 h under gentle shaking, followed by incubation in 50% LR in 90% ethanol overnight, 75% LR in 90% ethanol for 2 h, and finally in 100% LR for 3 h. The embedded samples were then placed in gelatine capsules at 55 °C for 48 h. Ultra-thin sections were then cut with a diamond knife to a thickness of 70 nm. Sections were collected on 200 mesh Formvar-coated nickel grids.

For immunological detection of ProDH, the grids were firstly incubated in goat serum 5% (v/v) in T1 buffer [0.05 M Tris-HCl, 2% (w/v) NaCl, 0.1% (w/v) BSA grade V, and 0.05% (v/v) Tween-20, pH 7.4] for 1 h. Antibodies raised against the ProDH1 protein were diluted 50-fold in T1 buffer and applied to the grids overnight at 4 °C. The grids were then washed five times with T1 buffer and three times with T2 buffer [0.02 M Tris-HCl, 2.5% (w/v) NaCl, 0.1% (w/v) BSA grade V, and 0.05% Tween-20 (v/v), pH 8.0] for 5 min each. The grids were incubated with a 10-nm colloidal gold–goat-anti-rabbit IgG whole molecule complex in T2 buffer for 2 h at room temperature and were then washed five times with T2 and five times with ultra-pure water for 5 min each. The grids were finally contrasted with 5% uranyl acetate in water for 5 min, washed three times with ultra-pure water and then with lead citrate tri-hydrate in water for 30 s, and washed 10 times with ultra-pure water. Observations were made using a Zeiss 912 TEM with Omega filter.

**RNA extraction and droplet digital PCR analysis**

RNAs were extracted from three leaves at the same developmental stage (~100 mg of plant tissue) that were ground in liquid nitrogen using a mixer mill (MM301; Retsch, Haan, Germany) according to Ben Rejeb et al. (2015). RNA quality and concentration were determined using a NP80 Nanophotometer® (Implen). A mixture of cDNAs corresponding to four copies per cDNA. The absolute expression quantitation of the formed using the Quantasoft™ software (Bio-Rad) to calculate the number of their corresponding wells of the DG8™ cartridge in order to be emul-sioned into 20 000 000 encapsulated nanodroplets. Data analysis was performed using the Quantasoft™ software (Bio-Rad) to calculate the number of copies per cDNA. The absolute expression quantitation of the ProDH genes was normalized using adenine phosphoribosyltransferase 1 (APRT1, At1g27450) and the ubiquinol–Cyt c reductase iron–sulfur subunit (AT5G, At5g13440) as housekeeping genes (Ben Rejeb et al., 2015). All the primers used for ddPCR are given in Supplementary Table S1 at JXB online.

**Protein extraction and western blotting**

Leaves (100–150 mg) were ground in a fine powder in liquid nitrogen. The frozen powder was then suspended into 10 ml of extraction buffer containing 8 M urea, 5% (w/v) SDS, 40 mM Tris-HCl pH 6.8, 0.1 mM EDTA, and 0.4 mg ml−1 bromophenol blue, supplemented with 1% protease inhibitor cocktail (Merck). Samples were shaken three times for 5 min with 5 min intervals on ice. The samples were then centrifuged at 18 000 g for 10 min at 10 °C and the supernatants (30 μl) were loaded and separated onto a 12% polyacrylamide gel. The proteins were then transferred onto nitrocellulose membranes (Amersham Hybond-ECL® from GE Healthcare) and were immunoblotted using purified antibodies directed against either Arabidopsis recombinant ProDH1, ProDH2, PSCDH, or PSCS proteins. Antibody binding was detected using a secondary antibody conjugated with a horseradish peroxidase and the ECL prime system from GE Healthcare using the G:BOX® system (Syngene).

**Identification of ProDH2 by MS analysis**

Preparation of samples and identification of proteins by MS was performed as described by Fromm et al. (2016). Briefly, 50 μg of proteins from crude mitochondria isolated from leaves of prold2 seedlings were loaded onto polyacylamide gels (staining gel 10% acrylamide, separating gel 14% acrylamide). Gel runs were stopped when the bromophenol blue dye front reached the end of the staining gel, and were subsequently stained with Coomassie Brilliant Blue G250. Gel pieces containing the proteins that concentrated at the stacking gel–separation gel interface were subjected to in-gel digestion followed by peptide extraction. MS analyses were performed using a Dionex Ultimate 3000 (ThermoFisher Scientific) equipped with a 50-cm reverse-phase analytical column (C18, ID 75 μm, particle diameter 3 μm, 100A pore size) in line to a 2-cm
reverse-phase pre-column (C18, ID 100 μm, particle diameter 5 μm, 100Å pore size). Peptides were eluted using a 1-h non-linear acetonitrile gradient and submitted to tandem MS employing a top-10 duty cycle. MS/MS data were subsequently evaluated using the Mascot (Matrix Science, London, UK) search engine querying against an in-house Arabidopsis (TAIR10) protein database.

Metabolic profiling using GC-MS
Sample preparation, analysis, and data processing were performed as previously described by Fiehn (2006). Ground, frozen samples (100 mg) from harvested leaves (7, 8, and 9 from the base of the plant) exposed to 5 d of either dark or light conditions were resuspended in 1 ml of frozen (−20 °C) water:acetonitrile:isopropanol (2:3:3) containing ribitol at 4 μg ml−1 and extracted by shaking for 10 min at 4 °C. Insoluble material was removed by centrifugation. Three blank tubes were subjected to the same steps as the samples. A quality control was made by pooling an equal volume from each light/dark condition. A sample of 30 μl was collected from each tube and dried overnight at 35 °C in a vacuum centrifuge.

Derivatization was performed according to Clément et al. (2018), 4 h after the end of which the whole sample series was first injected in splitless mode and then in split mode (1/30). Four different standard mixes were injected at the beginning and the end of the analysis as well as three independent derivatizations of the quality control (beginning, middle, and end) for monitoring the derivatization stability. Samples were randomized. The instrument was an Agilent 7890A gas chromatograph coupled to an Agilent 5977B mass spectrometer. The column was a Rxi-5SilMS from Restek (30 m with 10-μm integraguard column). The liner (Restek # 20994) was changed before analysis. The oven temperature ramp was 70 °C for 7 min then 10 °C min−1 to 330 °C for 5 min (run length 38 min). The helium constant flow was 0.7 ml min−1. The following temperatures were used: injector, 250 °C; transfer line, 290 °C; source, 250 °C; and quadrupole, 150 °C. Five scans per second were acquired spanning a 50–600 Da range. The instrument was tuned with perfluorotributylamine (PFTBA) with the 69 m/z and 219 m/z of equal intensities. The split mode conditions were 70 °C for 2 min, then 30 °C min−1 to 330 °C for 5 min. The helium constant flow was 1 ml min−1.

The data were processed as follows. The raw Agilent data files were converted into NetCDF format and analysed with AMDIS (http://chemdata.nist.gov/mass-spc/amdis/). A home retention indices/mass spectra library built from the NIST, Golm (http://gmd.mpimp-golm.mpg.de/), and Fiehn databases and standard compounds was used for metabolic identification. Peak areas were also determined with the Targetlynx software (Waters) after conversion of the NetCDF files into masslynx format. AMDIS, Target Lynx in splitless, and split 30 mode data were compiled in one single Excel file for comparisons. After subtraction of the blank mean, peak areas were normalized to ribitol and expressed on a fresh weight basis. Only metabolites showing repeatable and significant differences (based on t-tests) with respect to light and dark conditions are presented in heat maps. A complete list of all the metabolites that were detected can be found in Supplementary Data Set S1.

Statistical analysis
Two-way ANOVAs were performed for the qPCR analysis. The effects of senescence on chlorophyll and proline contents in wild-type and mutants, including time effects, as well as across days was assessed by ANOVA. Tukey tests were performed for comparisons of ProDH activities.

Results

Knockout Arabidopsis prodh mutants accumulate proline during dark-induced senescence

Dark-induced senescence (DIS) was triggered by cutting leaves of 4–week-old Arabidopsis Col-0 (wild-type, WT), prodh1, prodh2, and prodh1 prodh2 plants at the same developmental stage and placing them in the dark for 6 d. The progression of DIS was followed by measuring chlorophyll degradation over time. As shown in Fig. 1A, chlorophyll contents progressively decreased during DIS in all genotypes to less than 0.2 mg chlorophyll g−1 FW at Day 6 (contents could not be determined at 7 d due to degradation of the leaf tissue). The kinetics of chlorophyll degradation were similar to the WT in the both prodh single- and double- mutants. Chrobok et al. (2016) determined five time periods of senescence from T0, which corresponds to leaf expansion, to T4, which corresponds to the mature leaf at the late stage of senescence. Based on this reference, the chlorophyll contents indicated that Day 3 and Day 4 in our study were equivalent to Chrobok’s T2 period, and Day 5 and Day 6 were equivalent to the T3 period. We also monitored the proline content, and no significant variations were observed in the WT and the single-mutants up to Day 6 of DIS (Fig. 1B). However, significant increases in proline accumulation were measured in the prodh1 prodh2 double-mutant compared to the WT at Day 4 and Day 5, suggesting a putative role of ProDH in the oxidation of proline during senescence.

Expression of ProDH during DIS

ProDH expression was investigated in dark-induced senescent leaves in a time-course experiment. Leaves from 4-week-old Arabidopsis WT and prodh1 prodh2 plants were collected during senescence. Transcript analyses using the highly sensitive droplet digital PCR (ddPCR) method revealed very low transcript levels of both ProDH1 and ProDH2 in the WT at Day 0 (Fig. 2A), although the ProDH1 transcript level was 5-fold higher than ProDH2. During the 4 d of the senescence experiment, a consistent increase of ProDH1 transcripts (575-fold induction between 0–4 d), and to a lesser extent ProDH2, occurred in the WT (at Day 5, ProDH2 transcript levels could not be determined due to high degradation of RNAs). At Day 4, the ProDH1 transcript level was 6.5-fold higher than that of ProDH2. These results prompted us to monitor ProDH protein accumulation in a time-course experiment using an antibody directed against ProDH1 recombinant protein (Cabassa-Hourton et al., 2016). In the WT total protein extracts, no ProDH signal was detected at Day 0 (Fig. 2B). A faint signal appeared at Day 3, increased at Day 4 and Day 5, and reached a maximum at the end of the senescence period. A similar progression of ProDH accumulation was observed in the prodh2 mutant, but no ProDH signal could be detected in either the prodh1 or prodh1 prodh2 mutants. These results suggested that at least the ProDH1 isoform accumulated during DIS. ProDH2 could not be detected in the total protein extracts but an increase of ProDH2 transcripts was observed. As the antibody directed against ProDH1 could detect recombinant ProDH2 but with a lower sensitivity than recombinant ProDH1 (Cabassa-Hourton et al., 2016), the lack of detection of ProDH2 in the total protein extracts could have been due to the fact that ProDH2 is a very low abundant enzyme, as postulated by Cabassa-Hourton et al. (2016).

ProDH2 was therefore investigated by western blotting using a new antibody directed against ProDH2 recombinant protein. Leaves from 4-week-old plants were sampled after 5 d of DIS and crude mitochondria were isolated (Cabassa-Hourton et al., 2016). Western blots showed different signal patterns with antibodies directed against either ProDH1 or ProDH2 (Fig. 3). The anti-ProDH1 antibody showed a signal at an apparent molecular
mass of ~60 kDa that was not found in the prodh1 and prodh1 prodh2 mutants. In contrast, the use of the new antibody directed against ProDH2 recombinant protein resulted in the observation of both ProDH isoforms (Fig. 3). ProDH2 was only detected in the WT and prodh1 mutant with the anti-ProDH2 antibody. However, this antibody also detected ProDH1 in the WT and prodh2 mutant but to a lesser extent than with the antibody raised against ProDH1. A non-specific signal was observed at a higher apparent molecular mass for all the genotypes.

In order to further confirm the nature of the signal detected by the anti-ProDH2 antibody, a MS analysis was performed using crude mitochondria isolated from dark-induced senescent leaves of the prodh1 mutant. This resulted in the identification of 13 unique peptides exactly matching the ProDH2 protein, which covered 45% of the protein sequence with a MASCOT protein score (sum of the peptide score) of 633.16 (Fig. 4). These results confirmed the presence of the ProDH2 isoform in crude mitochondria of leaves subjected to DIS.

The mitochondrial localization of both ProDHs was confirmed by preparation of ultra-thin leaf sections that were immunogold-labelled with the anti-ProDH1 antibody and observed using TEM. The gold labelling showed a mitochondrial localization of ProDH in the WT at 5 d of DIS (Fig. 5). No signal was detected in the chloroplasts. Additional experiments were performed with the prodh single- and double-mutants in order to assess the identity of the isoform. Signals were detected in mitochondria of the prodh1 (Fig. 5B) and prodh2 (Fig. 5C) mutants, which indicated the presence of both ProDH isoforms in mitochondria during DIS. No signal was detected in the prodh1 prodh2 double-mutant (Fig. 5D). Taken together, these results demonstrated that both the ProDHs were expressed and accumulated in mitochondria during DIS.

P5CDH but not P5CS accumulates during DIS

We also investigated the effect of dark-induced leaf senescence on P5CS and P5CDH, markers of proline biosynthesis and proline catabolism, respectively, by western blotting using antibodies raised against P5CS1 and P5CDH recombinant enzymes (Fig. 6). In contrast to ProDH, P5CS and P5CDH were already...
present at Day 0. The level of P5CS protein was not really affected during the time-course experiment in the WT and the prodh1 prodh2 double-mutant. On the other hand, the P5CDH content slightly increased during DIS. These results showed that DIS enhanced proline catabolism but not proline biosynthesis.

**DIS triggers ProDH activity and proline respiration**

The enzymatic activity of ProDH was investigated in crude mitochondria extracts during DIS. After 5 d of DIS, ProDH activity increased by 5-fold in the WT (Fig. 7), but no activity could be measured in the prodh1 prodh2 double-mutant whatever the growth conditions, as has previously been reported by Cabassa-Hourton et al. (2016). Taken together, these results indicated that ProDH was functional during DIS.

Oxygen consumption by crude mitochondria extracts was investigated in the WT and the prodh1 prodh2 double-mutant. Complex I capacity through the cytochrome pathway increased by 2.5-fold after 5 d of DIS in both the WT and the double-mutant (Fig. 8A). Complex II capacity increased in the same range for the WT but to a lower extent for the double-mutant (Fig. 8B). The potential contribution of the AOX pathway was determined by completely inhibiting the CP with KCN and by activating the AOX pathway with DTT in the presence of excess of pyruvate. The capacity for electron flux through the AOX pathway increased by 2-fold in the WT during DIS when succinate was used as the substrate (Fig. 8C). In the WT, proline respiration increased through both the CP and AOX pathway by 14- and 3-fold, respectively, after 5 d of DIS (Fig. 8D). Interestingly, the potential contribution of the AOX pathway upon proline respiration was higher than the CP at Day 0 while the opposite was observed after 5 d of DIS, when 65% of the electron flux passed through the CP. No proline respiration was measured in the double-mutant (data not shown), as was previously observed by Cabassa-Hourton et al. (2016).

Taken together, these results indicated that there was a higher proline oxidation rate during DIS, making it a potential respiratory substrate, together with a shift of the major electron pathway for proline respiration from AOX to cytochrome.

**Metabolic changes in detached leaves subjected to DIS**

To gain more insights into ProDH function during senescence, a metabolic profiling analysis was carried out in detached leaves of the WT and prodh1 prodh2 double-mutant exposed for 5 d to either light or dark conditions. A GC-MS untargeted approach identified 63 metabolites, of which 32 were unknown, that were differentially accumulated in the double-mutant compared with the WT (Fig. 9). Clearly opposite behavior was generally observed mainly for differentially accumulated metabolites of the primary metabolism in the two genotypes exposed to light or dark conditions. Under light conditions, the identified metabolites were generally accumulated to a higher level in the WT than in the double-mutant, which was in contrast to DIS where the opposite response was observed. In the
Involvement of ProDHs in dark-induced leaf senescence

The TCA cycle, levels of citrate, aconitate, 2-oxoglutarate, succinate, and malate were significantly higher in prodh1 prodh2 than in the WT under DIS. This was also the case with some amino acids such as proline, hydroxyproline, lysine, acetyl-serine, tyrosine, GABA, aminoadipate, and homocysteine, with nucleobases such as adenine and uracil, and with sugars such as ribose, xylose, xylulose, erythritol, and fructose. Interestingly, proline and its derivatives such as hydroxyproline and GABA followed the same trend. Only glycerate and glucose were present at lower levels in the prodh1 prodh2 double-mutant compared to the WT in dark conditions. In contrast, the glutamate level was not significantly changed between light and dark conditions in the WT and the double-mutant.

**Discussion**

The remobilization of metabolites during stress and senescence plays an important role in plant adaptation to the environment. Here, we hypothesized a requirement for ProDHs for alternative respiration and for the remobilization of proline during stress and senescence. This was supported by the higher levels of proline and its derivatives in the prodh1 prodh2 double-mutant compared to the WT under DIS. The remobilization of metabolites during stress and senescence could be facilitated by ProDHs, which are involved in the remobilization of proline, a key amino acid in stress and senescence. The identification of the ProDH2 isoform in Arabidopsis mitochondria by mass spectrometry using crude mitochondrial extracts from leaves of prodh1 plants after 5 d of dark-induced senescence further supports the role of ProDHs in stress and senescence.

**Fig. 4.** Identification of the ProDH2 isoform in Arabidopsis mitochondria by mass spectrometry using crude mitochondrial extracts from leaves of prodh1 plants after 5 d of dark-induced senescence. (A) Summary of MS results. (B) The ProDH2 protein sequence with the identified peptides marked in red.

**Fig. 5.** Immunolocalization using antibodies directed against ProDH during dark-induced senescence in leaves of Arabidopsis. The electron micrographs show sections of leaves after 5 d of dark-induced senescence for (A) Col-0 (wild-type, WT), (B) prodh1, (C) prodh2, and (D) the prodh1 prodh2 double-mutant, and show typical labelling within the mitochondria. Arrows indicate the presence of ProDH. No signal was detected in prodh1 prodh2.

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senescence, and this was demonstrated by using a combination of genetic, biochemical, proteomic, and immunolocalization approaches.

Increased transcript abundance of both ProDH1 and ProDH2 during DIS

We found that the transcript levels of both ProDH1 and ProDH2 steadily increased over 4 d in detached leaves incubated in the dark (Fig. 2), although much higher levels of ProDH1 than ProDH2 were observed (6.5-fold higher at 4 d). In the literature, data concerning ProDH expression during senescence are very mixed depending on the way it is induced. In developmental leaf senescence (DLS), Chrobok et al. (2016) observed an equal increase of both ProDH1 and ProDH2 transcript abundance in Col-0 leaves. van der Graaff et al. (2006) also observed an increase of ProDH1 transcript levels in DLS but to a lesser extent than ProDH2. Previous studies have also described accumulation of ProDH2 transcripts during senescence in Arabidopsis (Buchanan-Wollaston et al., 2005; Funck et al., 2010; Allu et al., 2014; Chrobok et al., 2016) and in Brassica napus (Faës et al., 2015). In our experiments, leaf senescence was induced by placing detached leaves in the dark. In these conditions, much higher amounts of ProDH1 transcripts than of ProDH2 were measured, as also observed by van der Graaff et al. (2006). More generally, ProDH transcript levels have been found to be higher in dark-induced senescence in detached leaves, in individually darkened leaves or darkened whole plants than during DLS (van der Graaff et al., 2006; Yu et al., 2016).

Both ProDH isoforms are located in the mitochondria

We found an increase in abundance of both ProDH transcripts but mRNA and protein levels are not always strictly correlated, as has been demonstrated for ProDH1 in response to water- and salt-stress conditions (Parré et al., 2007). Indeed, the protein concentration in a cell is determined by several factors such as transcription rate, transcript stability, and/or protein degradation. Van Aken et al. (2009) showed that variations in transcript levels of genes that putatively encode mitochondrial proteins are generally greater than those of proteins under stress-related conditions. Moreover, microarray analyses have shown that the half-life times of ProDH1 and ProDH2 transcripts are different, being estimated at 6.3 h and 2.2 h, respectively (Narsai et al., 2007). It was therefore essential to examine whether the observed changes in ProDH1 and ProDH2 transcript levels were reflected in the amounts of their corresponding protein. Accumulation of ProDH1 was clearly observed during DIS in total protein extracts (Fig. 2) and in crude mitochondrial fractions (Fig. 3), while ProDH2 was only found in crude mitochondria but not in total protein extracts (data not shown) when using antibodies raised against recombinant ProDH2. The absence of any signal in the total protein extracts when using antibodies raised against ProDH2 may probably be explained by a co-migration of ProDH2 with Rubisco, with the abundance of the latter masking ProDH2; however, the presence of ProDH2 in crude mitochondria was also confirmed by MS (Fig. 4). To our knowledge, this is the first time that ProDH2 has been identified using a proteomic approach (Klodmann et al., 2011; Rao et al., 2017).

ProDH1 has previously been shown to be clearly localized in mitochondria (Funck et al., 2010; Cabassa-Hourtou et al., 2016) while dual targeting to chloroplasts and mitochondria has been proposed for ProDH2 as a result of GFP fusion experiments (Van Aken et al., 2009). Our immuno-microscopy results clearly showed that ProDH2 was only detected in mitochondria during DIS (Fig. 5).

ProDHs contribute to fueling respiration and the TCA cycle during senescence

We found a higher respiration capacity in complex I and II after 5 d of DIS, whatever the electron pathway (Fig. 8). In senescence induced by covering individual Arabidopsis leaves with aluminum foil, Keech et al. (2007) also reported higher levels of respiration at 6 d of senescence. An increase in respiration has also been shown in leaves of Populus tremuloides and Quercus rubra during autumnal senescence (Collier and
Fig. 8. Oxygen consumption in mitochondria isolated from Arabidopsis leaves from Col-0 (wild-type, WT) and the prodh1 prodh2 double-mutant (DM) in response to dark-induced senescence. Oxygen consumption was determined using a Clark-type electrode in crude mitochondria extracts from leaves 7, 8, and 9 from the base of the plant sampled at 0 d and 5 d of dark-induced senescence. The capacity of the cytochrome pathway (CP) was measured from (A) complex I, (B) complex II, and (D) proline. The capacity of the alternative oxidase pathway (AOX) was measured in WT mitochondria from (C) complex II and (D) proline. Data are means (±SE) of at least four biological replicates. Different letters indicate significant differences between means as determined using ANOVA (P<0.05).

Fig. 9. Primary metabolism pathways of metabolites measured in detached leaves of Arabidopsis under light or dark conditions compared between Col-0 (wild-type) and the prodh1 prodh2 double-mutant. Heat-maps are shown for changes in metabolites in pooled samples of leaves 7–9 from the base of the plant after 5 d under light (left square) or dark (right square) conditions: the blue-to-yellow scale shows the log2 ratio of the fold-change between the double-mutant and the wild-type. Data represents the mean values of four biological replicates for each time-point.
Interestingly, our work showed for the first time that senescence stimulated a striking increase in proline respiration (Fig. 8D). Cabassa-Hourton et al. (2016) demonstrated that treatment of Arabidopsis seedlings with proline triggers ProDH1 accumulation and activity. Concomitantly, proline respiration increases but to a significantly lesser extent than during senescence: it was two-fold higher in DIS than in proline-treated seedlings. To our knowledge, our data represents one of the highest rates of proline respiration that has been measured. Electron fluxes generated from proline respiration in proline-treated seedlings (Cabassa-Hourton et al., 2016), as well as in leaves before senescence (Day 0, this study), are mainly directed through the AOX pathway and do not efficiently contribute to ATP synthesis. However, upon DIS in our study, the electron flux from proline oxidation was preferentially directed through the CP, allowing a better coupling to ATP synthesis. Taken together, these observations suggest the importance of proline as an alternative respiratory substrate during senescence. Proline oxidation may contribute to the generation of ATP that is necessary for the maintenance of leaf viability while the remobilization of nutrients is carried out. It has previously been reported that the total oxidation of one proline molecule can generate reductants that can fuel the mitochondrial electron transfer chain and support the formation of 30 ATP molecules (Atkinson, 1977). DIS leads to intensive metabolite recycling through protein degradation and the generation of amino acids that are used as alternative substrates for respiration. More generally, energy-limited conditions require alternative pathways of respiration to provide the necessary respiratory function (Barros et al., 2017). Interestingly, proline has also been shown to be the main energy substrate to fuel flight in some insects (Gäde and Auerswald, 2002).

In addition to its role in respiratory electron flux, ProDH is more widely involved in mitochondrial metabolism. Our untargeted metabolomic approach based on GC-MS provided a more comprehensive view of the role of ProDH in primary metabolic pathways during senescence (Fig. 9). Interestingly, this analysis identified increased levels of citrate, aconitate, 2-oxoglutarate, succinate, malate, proline, and hydroxyproline in the prodh1 prodh2 double-mutant in detached leaves subjected to DIS when compared to the WT. These metabolites are linked to the TCA cycle and associated pathways. The increase of TCA intermediates in the double-mutant may indicate a lower availability of alternative substrates such as proline, which could no longer be oxidized to supply the carbon backbone in the form of glutamate or 2-oxoglutarate to sustain the TCA cycle. An increase in TCA intermediates has previously been observed in plants lacking D-2-hydroxyglutarate dehydrogenase or isovaleryl-CoA dehydrogenase, demonstrating the participation of lysine and branched amino acids as alternative substrates for respiration during senescence (Aráujo et al., 2010; Engqvist et al., 2011). More generally, a reduced flow of carbon into the TCA cycle seems to lead to an increase of intermediates, probably due to a reduced overall activity of the TCA cycle (Aráujo et al., 2010; Yu et al., 2012; Huang et al., 2013; Pires et al., 2016).

Metabolites from sugar metabolism and the shikimate pathway together with homocysteine, a sulfur-containing amino acid, also accumulated more in the double-mutant than in the WT during DIS (Fig. 9). Interestingly, Law et al. (2018) have shown that genes encoding proteins involved in primary energy production (respiration, fermentation, and β-oxidation), amino acids, lipid or nucleotide catabolism, sulfur metabolism, and the shikimate pathway are overexpressed during senescence of individually darkened leaves. ProDH and proline may thus be crucial actors of this metabolic switch under senescence.

Proline generated by DIS can be metabolized by ProDH enzymes to produce P5C. This can also be synthesized by ornithine aminotransferase, the mRNA levels of which are also up-regulated during senescence (Buchanan-Wollaston et al., 2005; van der Graaff et al., 2006; Chrobok et al., 2016). We also found that the P5CDH protein, which allows the conversion of P5C to glutamate, was up-regulated in DIS (Fig. 6). Previous results for DLS have shown either no response or up-regulation (van der Graaff et al., 2006). No induction of P5CS was observed under our conditions, which is consistent with previous results reported for leaves subjected to DIS (van der Graaff et al., 2006). Thus, DIS triggers proline catabolism rather than proline biosynthesis and allows the generation of glutamate, which is the metabolic precursor of glutamine and asparagine that are involved in nitrogen remobilization during senescence (Finnemann and Schjoerring, 2000; Lin and Wu, 2004; Masclaux-Daubresse et al., 2010). Proline content has been shown to decrease during developmental senescence in wild-type Arabidopsis and in tobacco (Masclaux et al., 2000; Chrobok et al., 2016), which indicates an important role of ProDH in proline oxidation to release glutamate. Moreover, it has been shown that proline regulates the expression of Glutamine Synthase 1 (GS1) and Glutamate Dehydrogenase (GDH), which encode two enzymes involved in nitrogen remobilization during senescence (Masclaux-Daubresse et al., 2005), thus supporting an important role of this amino acid as a signal molecule during senescence.

Proline could originate from the degradation of cellular and cell wall proteins, which are notably rich in proline and hydroxy (Ihsan et al., 2017), as has also been described for the mammalian extracellular matrix (Pandhare et al., 2009). Degradation of cell wall proteins could therefore lead to an increase of free proline concentration in the cell that would act as an activator of ProDH transcription and as a substrate for both ProDH isoforms, contributing to nitrogen remobilization through glutamine generation from glutamate and its subsequent transport to sinks. In conclusion, our results indicate a dual role of ProDH under DIS. Proline oxidation allows it to be used as an alternative respiratory substrate and contributes to the production of glutamate and energy that can subsequently participate in the remobilization of nutrients from senescent tissues to developing plant organs.

**Supplementary data**

Supplementary data are available at *JXB* online.

Table S1. List of primers used for ddPCR.

Data Set S1. Raw data and statistical analysis of metabolite profiles in the leaf experiments.
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References


