The versatility of plant organic acid metabolism in leaves is underpinned by mitochondrial malate–citrate exchange

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Abstract

Malate and citrate underpin the characteristic flexibility of central plant metabolism by linking mitochondrial respiratory metabolism with cytosolic biosynthetic pathways. However, the identity of mitochondrial carrier proteins that influence both processes has remained elusive. Here we show by a systems approach that DICARBOXYLATE CARRIER 2 (DIC2) facilitates mitochondrial malate–citrate exchange in vivo in Arabidopsis thaliana. DIC2 knockout (dic2-1) retards growth of vegetative tissues. In vitro and in organello analyses demonstrate that DIC2 preferentially imports malate against citrate export, which is consistent with altered malate and citrate utilization in response to prolonged darkness of dic2-1 plants or a sudden shift to darkness of dic2-1 leaves. Furthermore, isotopic glucose tracing reveals a reduced flux towards citrate in dic2-1, which results in a metabolic diversion towards amino acid synthesis. These observations reveal the physiological function of DIC2 in mediating the flow of malate and citrate between the mitochondrial matrix and other cell compartments.
Introduction

Malate is a prominent metabolite that occupies a pivotal node in the regulation of plant carbon metabolism. It is the mainstay of leaf respiration and metabolic redox shuttling between organelles (Scheibe, 2004). Early studies with isolated plant mitochondria demonstrated that exogenous malate can be translocated in exchange for an inorganic phosphate by an unknown butylmalonate-sensitive carrier into the mitochondrial matrix where it is then rapidly oxidized by both mitochondrial malate dehydrogenase (mMDH) and NAD-dependent malic enzyme (NAD-ME), generating oxaloacetate (OAA) and pyruvate as products (Figure 1; Macrae, 1971a, 1971b; Douce et al., 1973; Hatch and Kagawa, 1974; Wiskich, 1974, 1975; Day and Hanson, 1977a). OAA and the product of pyruvate, acetyl-CoA, can then be combined in the first step of the tricarboxylic acid (TCA) cycle to form citrate inside the mitochondrial matrix. However, a combination of enzyme kinetics (Kearney et al., 1972; Journet et al., 1981) and redox coupling reactions in the cytosol and mitochondria (Day and Hanson, 1977b; Wiskich and Day, 1982; Lance and Rustin, 1984) results in marked differences in the rate of malate oxidation and citrate production inside leaf mitochondria in the light and the dark. TCA cycle-driven respiration is largely inhibited in the light and mMDH generally operates in the direction of malate synthesis to facilitate redox coupling between the matrix and other cell compartments, resulting in net mitochondrial OAA import and malate efflux (Tcherkez et al., 2005, 2009; de Oliveira Dal'Molin et al., 2010; Bykova et al., 2014; Hüdig et al., 2015). At night, the requirement for exchanging reducing equivalents between mitochondria and chloroplasts via a malate valve (Kromer, 1995; Raghavendra and Padmasree, 2003; Scheibe, 2004; Selinski and Scheibe, 2019; Zhao et al., 2020) is believed to be minimal due to the cessation of both photorespiration and photoinhibitory conditions. The mitochondrial NAD-ME activity and its transcript abundance are highest in the dark (Tronconi et al., 2008); therefore, the synthesis and oxidation of malate is expected to be carried out by both mMDH and NAD-ME to support the synthesis of both citrate and ATP (Lee and Millar, 2016). These metabolic conditions would enable maximal citrate oxidation in the mitochondrial matrix (Igamberdiev et al., 2001; Tovar-Méndez et al., 2003), with excess citrate being exported for storage in the vacuole (Cheung et al., 2014). Mitochondrial citrate is also a major source of cytosolic acetyl-CoA (Schwender et al., 2006; Cheung et al., 2014) required for the elongation of fatty acids and the biosynthesis of isoprenoids and flavonoids (Fatland et al., 2002, 2005; Baud et al., 2003; Oliver et al., 2009). Exported citrate can also be metabolized by aconitase to sustain the TCA cycle and the glyoxylate cycle, as well as to support nitrogen assimilation by producing glutamate or glutamine as an end-product (Carrari et al., 2003; Gauthier et al., 2010; Foyer et al., 2011; Hooks et al., 2014).

Inhibition in the synthesis of citrate from malate-derived OAA in mitochondria and its export leads to floral sterility in potato (Solanum tuberosum) plants (Landschütze et al., 1995) and a change in nitrogen incorporation into amino acids (Kumagai et al., 2003). At night, the requirement for exchanging reducing equivalents between mitochondria and chloroplasts via a malate valve (Kromer, 1995; Raghavendra and Padmasree, 2003; Scheibe, 2004; Selinski and Scheibe, 2019; Zhao et al., 2020) is believed to be minimal due to the cessation of both photorespiration and photoinhibitory conditions. The mitochondrial NAD-ME activity and its transcript abundance are highest in the dark (Tronconi et al., 2008); therefore, the synthesis and oxidation of malate is expected to be carried out by both mMDH and NAD-ME to support the synthesis of both citrate and ATP (Lee and Millar, 2016). These metabolic conditions would enable maximal citrate oxidation in the mitochondrial matrix (Igamberdiev et al., 2001; Tovar-Méndez et al., 2003), with excess citrate being exported for storage in the vacuole (Cheung et al., 2014). Mitochondrial citrate is also a major source of cytosolic acetyl-CoA (Schwender et al., 2006; Cheung et al., 2014) required for the elongation of fatty acids and the biosynthesis of isoprenoids and flavonoids (Fatland et al., 2002, 2005; Baud et al., 2003; Oliver et al., 2009). Exported citrate can also be metabolized by aconitase to sustain the TCA cycle and the glyoxylate cycle, as well as to support nitrogen assimilation by producing glutamate or glutamine as an end-product (Carrari et al., 2003; Gauthier et al., 2010; Foyer et al., 2011; Hooks et al., 2014).
Acid transporters in Arabidopsis thaliana remained elusive. The identity and in vivo function for malate, OAA, citrate and alanine transporters are not clear. AlaAT, alanine aminotransferase; CS, citrate synthase; MDH, malate dehydrogenase; MPC, mitochondrial pyruvate carrier; PDC, pyruvate dehydrogenase complex.

By identifying homologues of yeast and mammalian carriers in Arabidopsis thaliana, several possible candidates that may contribute to mitochondrial malate transport in plants were identified (Picault et al., 2002; Palmieri et al., 2008; Monné et al., 2018). In contrast to the historical, well-established models that mitochondrial metabolite transporters are discrete and highly specific (Wisich, 1977; Klingenberg, 1979; LaNoue and Schoolwerth, 1979), these carriers appeared to lack substrate specificities under in vitro conditions—for example, dicarboxylate carrier (DIC) isoforms in proteoliposomes can rapidly exchange sulfate with phosphate, malate, OAA, and succinate, with an apparent low exchange activity in the presence of citrate, 2-oxoglutarate and fumarate (Palmieri et al., 2008). Although in vitro studies have been instrumental in revealing what transport activities can be mediated by a protein, they have limitations that turn out to be critical in the case of plant mitochondrial organic acid transporters. For instance, they cannot suitably consider the often unknown but clearly highly changeable metabolite pool sizes and fluxes that the transporters face in vivo. Reconstitution with a specific orientation of the transporter is not possible in most in vitro systems, but is critical for substrate specificity and respiratory physiology in vivo. Further, the specific local lipid environment and the pronounced electrochemical gradients (ΔΨ, ΔpH), both of which are likely to be central for inner mitochondrial membrane transport activity and specificity (Lee and Millar, 2016), can only be roughly considered. If we are to understand the roles that metabolite transporters fulfill in the living plant, additional information is needed about the influence of an individual transporter on net transport across a membrane within a physiologically meaningful context. Yet, in planta studies of mitochondrial metabolite transporter proteins that address the question of their in vivo influence on metabolism, requiring a systems perspective to consider their integration in the metabolic network, have been lacking.

In this study, we determine that DIC2 functions as a high affinity malate–citrate antiporter in Arabidopsis mitochondria, through evidence from the reverse genetic approach in combination with comprehensive in vitro, in organello, and in vivo analyses. We present strong evidence that plant DIC2 activity plays a critical role in dynamically coordinating anaplerotic metabolism with assimilatory and catabolic pathways between the mitochondria and other cellular compartments.

**Results**

DIC2 mutation causes decelerated vegetative growth rate that cannot be compensated by other mitochondrial dicarboxylate carrier isoforms

DIC2 is a mitochondrial protein based on interpretation of fluorescent microscopy images of transient (Van Aken et al., 2009) or stable expression of DIC2-GFP (Supplemental Figure S1) and Arabidopsis mitochondrial proteome analysis by mass spectrometry (Senkler et al., 2017; Fuchs et al., 2020). Three T-DNA insertion lines were obtained in order to study the possible physiological and metabolic consequences of a defect in DIC2 (At4g24570) function in planta. Only the homozygous dic2-1 line has a T-DNA insertion within the open reading frame of DIC2, resulting in the complete loss of DIC2 transcript (Figure 2A; Supplemental Figure S2A). Using selective reaction monitoring mass spectrometry to determine protein abundance (Taylor et al., 2014), we found that two different DIC2 peptides, detectable in Col-0, were not detectable in mitochondria isolated from dic2-1 plants (Figure 2A; Supplemental Figure S2B), confirming dic2-1 as a null mutant for DIC2. The homozygous dic2-1

![Figure 1 Regulation of malate oxidation leading to citrate production in Arabidopsis. Two malate-oxidizing pathways contribute to the formation of citrate in plant mitochondria. Malate can be oxidized into either OAA or pyruvate. OAA accumulation leads to the inhibition of the malate-oxidizing malate dehydrogenase reaction. The availability of acetyl-CoA for condensation with OAA by citrate synthase is another factor for the regulation of mitochondria malate oxidation. During the day, pyruvate-derived acetyl-CoA amount is limited due to the inhibition of pyruvate dehydrogenase phosphorylation and the reverse reaction of malate dehydrogenase is favored to support photorespiration. At night, NAD-dependent malic enzyme (NAD-ME) and dephosphorylated pyruvate dehydrogenase are activated, allowing optimal supply of acetyl-CoA and OAA for efficient citrate synthesis. While the identity of plant mitochondrial pyruvate carrier (MPC) has recently been confirmed (Le et al., 2021a), the exact molecular identity and in vivo function for malate, OAA, citrate and alanine transporters are not clear. AlaAT, alanine aminotransferase; CS, citrate synthase; MDH, malate dehydrogenase; MPC, mitochondrial pyruvate carrier; PDC, pyruvate dehydrogenase complex.
showed reduced root growth during seedling development (Supplemental Figure S2C) and a significantly decreased rate of rosette expansion (Figure 2B; Supplemental Figure S2D). From Day 42, the number of leaves emerged from Col-0 and dic2-1 was identical, but the mutant leaves were smaller and curly with a more rugose surface (Supplemental Figure S2F). dic2-1 could not reach the full rosette diameter of Col-0 after 1 week of bolting (Supplemental Figure S2G). dic2-1-specific phenotypes could be observed regardless of the photoperiod (Supplemental Figure S2H). These phenotypes could be restored by introducing DIC2 under the control of its native promoter (dic2-1/gDIC2) or a cauliflower mosaic virus 35S promoter (dic2-1/DIC2 OE; Figure 2B; Supplemental Figure S2D). We attempted to rescue the dic2-1 phenotype by expressing DIC1 and DIC3 in the dic2-1 background (dic2-1/DIC1 OE and dic2-1/DIC3 OE, respectively), but neither could complement the loss of DIC2, not even partially (Figure 2B). These results indicate that DIC2 does not share substrate preferences and/or specificities with DIC1 or DIC3 in planta.

**Net citrate export by energized mitochondria is compromised as a consequence of DIC2 loss**

To explore the transport function of DIC2, we first tested the hypothesis that malate is the main substrate for DIC2 in the context of intact isolated mitochondria, as inferred previously by a proteoliposome-based approach (Palmieri et al., 2008). To this end, we monitored the transport activity for $^{14}$C-malate by nonenergized isolated mitochondria in a minimal reaction medium that lacked added cofactors and ADP to allow malate to more slowly accumulate while preventing its conversion into other TCA cycle intermediates. Under such conditions, the initial uptake rate of 200-$\mu$M $^{14}$C-malate into mitochondria of dic2-1 was three-fold lower compared to Col-0 and dic2-1/gDIC2 (Figure 3A). Hence, DIC2 facilitates a significant malate uptake capacity in isolated Arabidopsis mitochondria and its loss limits this uptake, even in the presence of other potential carriers for malate transport.

Titrations with respiratory substrates at different concentrations revealed no obvious differences in the rate of internal or external NADH-driven O$_2$ consumption between Col-0 and dic2-1 (Supplemental Figure S3A). No change in the activity or distribution of electron transport chain (ETC) supercomplexes (Supplemental Figure S3B), the relative abundance of TCA cycle enzymes, pyruvate dehydrogenase or NAD-ME (Supplemental Figure S3C), or their $K_m$ and $V_{max}$ (Supplemental Table S2) were observed when mitochondria isolated from these genotypes were compared. This eliminated the possibility of a clear defect in the ETC or a specific step of the TCA cycle in dic2-1; instead, any shift in the distribution of specific metabolites between the matrix and extra-mitochondrial space in the mutant could reasonably be attributed directly to the absence of DIC2 and/or indirectly to the enzymatic regulatory compensation as the result of DIC2 loss.

Isolated mitochondria were then tested for their ability to consume malate under state III respiration condition and to generate and export metabolites synthesized from malate (Figure 3B; Supplemental Figure S4A). The rate of malate uptake by energized mitochondria was similar between the genotypes (Figure 3C), possibly due to compensation by other malate carriers. The abundance of pyruvate, generated through the action of NAD-ME from imported malate, increased linearly over time in mitochondria of Col-0 and dic2-1/gDIC2, whereas it leveled off in dic2-1 (Figure 3D).
Consistent with this, dic2-1 mitochondria exhibited a 20%–30% lower $V_{\text{max}}$ for malate-dependent state III $O_2$ consumption (Supplemental Figure S4C). However, pyruvate accumulation in the extramitochondrial space (EMS) was maintained in the mutant mitochondria in the same way as in Col-0 (Figure 3E), ruling out the possibility of a defect in pyruvate export. The apparent increases in fumarate export to the EMS and its accumulation in mitochondria appear to be compensatory effects to avoid prolonged overaccumulation of malate through its rapid removal by dehydration in the dic2-1 matrix (Figure 3, F and G), which would provide an explanation for the apparent lack of mitochondrial malate accumulation in dic2-1 compared to controls (Supplemental Figure S4A). These data pinpoint a reduced rate of malate oxidation in the matrix of malate-fed dic2-1 mitochondria as a plausible cause for the observed decrease in malate-dependent respiration and pyruvate formation.

Strikingly, upon malate feeding, there was a much higher rate of citrate accumulation inside dic2-1 mitochondria (Figure 3I), accompanied by a lower rate of export of citrate to the EMS when compared to Col-0 mitochondria (Figure 3H). We thus hypothesized that excess citrate in
**dic2-1** mitochondria was caused by compromised citrate export, which then prevented OAA condensation in the citrate synthase reaction and thus suppressed mMDH activity (Fahren et al., 1988). To confirm this, we tested the ability of mitochondria to directly consume citrate (Supplemental Figure S4B). Under such conditions, NAD-ME and mMDH activities became inhibited, as demonstrated by the low mitochondrial export rate of pyruvate. There was no change in citrate-dependent O$_2$ consumption mitochondria from all genotypes (Supplemental Figure S4D), further confirming that the mitochondrial citrate import rate was not affected and the reduction in malate-dependent respiration by **dic2-1** was likely caused by the citrate accumulation in the matrix triggering an OAA inhibition of mMDH. While the uptake rate of citrate by mitochondria was not altered significantly (Figure 3), the rate of citrate accumulation in **dic2-1** mitochondria was 2.3-fold higher than Col-0 or **dic2-1/gDIC2** (Figure 3K) without compromising product formation and accumulation rates (Supplemental Figure S4B). Taken together, while DIC2 is directly responsible for a portion of malate uptake, its loss also influences the export of citrate. However, the fact that citrate export was not completely abolished by the loss of DIC2 suggests that this critical export across the inner mitochondrial membrane is also not solely dependent on the presence of DIC2.

DIC2 preferentially exchanges malate against citrate in proteoliposomes

Our finding that DIC2 loss alters mitochondrial citrate export was unexpected, since proteoliposome-reconstituted DIC2 has been reported to have a relatively low citrate exchange activity against sulfate (Palmieri et al., 2008). It is possible that, in the absence of an electrochemical gradient across the liposomal membrane, DIC2 does not support an electrogenic exchange of a divalent dicarboxylate/sulfate/phosphate against a trivalent citrate at pH 7.0. To verify whether DIC2 acts as citrate transporter, or its loss rather indirectly contributes to a decrease in citrate export in organello, DIC2 was expressed heterologously in yeast and isolated microsomal membrane fraction was reconstituted into proteoliposomes (Figure 4A; Supplemental Figure S5A). Proteoliposomes preloaded with sulfate, phosphate, or citrate were incubated with TCA cycle organic acids (250 μM each) simultaneously at pH 6.0 (to shift the citrate buffer toward H-citrate$^+$) to assess DIC2 substrate specificity under a competitive condition. DIC2 did not demonstrate significant transport of any metabolites in the presence of internal sulfate or phosphate, including malate, succinate and OAA, when compared to the empty vector control (Supplemental Figure S5, B and C). In contrast, DIC2 showed a clear preference for malate when proteoliposomes were preloaded with citrate (Figure 4B).

Malate–citrate exchange by DIC2 was saturable in a time- and concentration-dependent manner (Figure 4C; Supplemental Figure S5D), with an apparent $K_m$ of 48 μM for malate (Figure 4D) which is 10-fold lower than in a previous report (0.51 mM; Palmieri et al., 2008). While the concentration of malate and citrate in the intermembrane space is currently unknown, such a $K_m$ value is below their expected cytosolic amount in plants (Gerhardt and Heldt, 1984; Gerhardt et al., 1987; Winter et al., 1994) and within the expected concentration range of most organic acids in intact mammalian mitochondria (10–50 μM; Chen et al., 2016). Our results also indicate that citrate is transported by DIC2 as a divalent anion or a trivalent form with cotransport of a proton. Together with the data from isolated mitochondrial assays, we concluded that DIC2 is a biologically relevant high affinity and directional malate$_{in}$/citrate$_{out}$ transporter.

**DIC2 function affects leaf dark respiration by modulating NAD homeostasis**

We then set out to determine how defects in mitochondrial malate import and citrate export operate in planta and if they could explain the observed phenotypes of **dic2-1**. In total we investigated three scenarios where leaf mitochondria operate under different flux modes: in the light, during the transition from light to dark and in the dark. First, we carried out chlorophyll fluorescence and infrared gas-exchange analyses at different light intensities, as changes in photosynthetic performance can cause many metabolic perturbations and mask other primary metabolic effects in mutants. There was no change in the CO$_2$ assimilation rate (Figure 5A), stomatal conductance, transpiration rate, photosynthetic electron transport rate, photosystem II quantum yield, or nonphotochemical quenching in **dic2-1** leaves (Supplemental Figure S6, A–E). When plants were shifted to high light for 3 and 16 h, we did not detect any changes in the maximum photochemical efficiency of PSII ($F_v/F_m$) among **dic2-1**, Col-0, and **dic2-1/gDIC2** (Supplemental Figure S6F). These results indicated that the loss of DIC2 has no direct impact on photosynthetic performance.

Second, we examined the transition from light to darkness where leaf mitochondria undergo several rapid changes in metabolic activity as photorespiration ceases and the noncyclic mode of the mitochondrial TCA cycle is activated. Upon transfer of illuminated leaves to darkness, **dic2-1** exhibited a more rapid release of CO$_2$ in the first 30–40 s before reaching a higher rate of steady-state respiratory CO$_2$ release compared to Col-0 and **dic2-1/gDIC2** (Figure 5B). The sharp CO$_2$ release upon sudden darkness is indicative of a post-illumination burst (PIB) of respiration, an estimate of the degree of respiratory glycine oxidation (Rawsthorne and Hylton, 1991). To examine if photorespiratory activities were altered in the mutant, we grew the mutant under high CO$_2$ (0.2%) to suppress photorespiration but no rescue of the phenotype was achieved (Supplemental Figure S6G). These data indicate that an altered photorespiratory activity after dark shift is not the primary cause for the change in PIB response by **dic2-1**. Since malate/OAA utilization in mitochondria could control the degree of glycine oxidation by regulating NADH/NAD$^+$ (e.g. via a malate shuttle; Journet et al., 1981), we hypothesized that a PIB increase in **dic2-1** could be linked to an altered distribution of TCA cycle metabolites between mitochondria and cytosol (and possibly
plastids), leading to changes in metabolic and/or NAD redox state in these compartments. To capture any rapid and transient changes in NAD redox state during a sudden dark shift, we utilized a fluorescent protein biosensor Pедерп-мCherry which reports cytosolic NADH/NAD^+ through the ratio of tSapphire to mCherry fluorescence (log_{10}(tS/mC)), where a higher log_{10}(tS/mC) corresponds to a more reduced NAD pool (Steinbeck et al., 2020). We observed equally high NADH/NAD^+ ratios in Col-0 and dic2-1 plants during illumination (Figure 5C). However, upon transfer to darkness, the expected decline in NADH/NAD^+ ratios in the first 100 s was significantly slower in dic2-1 compared to Col-0. Both differences in PIB and NAD redox status upon rapid transition to darkness indicate some defect in switching between light and dark flux modes by dic2-1 mitochondria.

Third, we examined metabolic changes to dic2-1 leaves during the night. DIC2 expression steadily increased over the course of nighttime, followed by a decline from the peak at the end of dark photoperiod to the lowest level in the light (Supplemental Figure S6H). Intriguingly, while there was no difference in the NADH/NAD^+ ratio between the two genotypes over the course of the dark–light cycle (Figure 5D), the leaf nighttime respiration rate (R_N) remained consistently higher in the mutant (Figure 5E). Given that primary metabolites are crucial determinants for regulating transcript abundance during a diurnal cycle (Gibon et al., 2006), there appears to be a metabolic rearrangement in the mutant to avoid any detrimental impact of DIC2 absence on NAD redox equilibrium, albeit at the expense of heightened respiration (Figure 5B and E) and slower vegetative growth.
DIC2 function causes an altered 2-OG metabolism in the mutant. Branched chain amino acids (BCAAs), which use 2-OG as co-substrate to initiate their catabolism in mitochondria, were also more abundant in dic2-1 compared to Col-0 and dic2-1/gDIC2 plants 1 h after the shift to darkness, including two- to four-fold higher abundances of pyruvate, citrate, and isocitrate, and a remarkable 10-fold increase in 2-oxoglutarate abundance (2-OG). Several amino acids linked to 2-OG utilization also changed in abundance in dic2-1. In particular, the glutamate pool in dic2-1 was larger than in Col-0 and dic2-1/gDIC2 plants 1 h after the dark shift and during the light photoperiod, while GABA and glutamine abundances remained unchanged. Aspartate and alanine, which can be converted into OAA and pyruvate respectively in a reversible reaction that requires glutamate/2-OG transamination, were also generally higher in abundance in dic2-1 plants at night. Such increases in cellular 2-OG utilization in dic2-1 would be consistent with increases in citrate retention and subsequent 2-OG and/or glutamate formation and export by mutant mitochondria (Figure 3I). Overall, the metabolite profiles revealed that the loss of DIC2 function causes an altered 2-OG metabolism in the dark, which may provide a metabolite buffer to compensate for a defect in mitochondrial DIC2 activity while cytosolic NAD redox state is maintained.

We next further investigated the cause of the increased RN in dic2-1. An increased nighttime consumption rate of succrose, but not of glucose or fructose, by dic2-1 was observed (Figure 6B; Supplemental Figure S7B). The expression of several nutrient-responsive senescence markers, SAG101 (He and Gan, 2002), WRKY53 (Miao and Zentgraf, 2007), and SEN1 (Oh et al., 1996), was also highly upregulated in the mutant in the dark but not in the light (Supplemental Figure S7C), suggesting nighttime starvation. These results, combined with a lack of photosynthetic differences (Figure 5A; Supplemental Figure S6, A–E), suggest that an accelerated depletion rate of carbon stores via respiration leads to nighttime starvation in dic2-1. Thus, changes in the cytosol and/or mitochondria to maintain metabolic homeostasis required for fundamental cellular functions.

### 2-oxoglutarate sharply accumulates in dic2-1 leaves upon shift to darkness

To gain further insight into how DIC2 function is integrated into metabolism and influences NAD redox state in vivo, we carried out metabolite profiling of leaf samples collected from 6-week-old plants at different time points of a diurnal cycle (Figure 6A; Supplemental Figure S7A). The metabolite profiles of malate, fumarate, and succinate over a diurnal cycle were generally similar between genotypes. Also, a similar pattern in daytime glycerol and serine accumulation was observed, despite a clear difference in PIB (Figure 5B), further confirming that DIC2 function does not directly affect photosynthetic flux. The most notable metabolite level changes in dic2-1 plants compared to Col-0 and dic2-1/gDIC2 were observed in the first hour after the sudden shift to darkness, including two- to four-fold higher abundances of pyruvate, citrate, and isocitrate, and a remarkable 10-fold increase in 2-oxoglutarate abundance (2-OG). Several amino acids linked to 2-OG utilization also changed in abundance in dic2-1. In particular, the glutamate pool in dic2-1 was larger than in Col-0 and dic2-1/gDIC2 plants 1 h after the dark shift and during the light photoperiod, while GABA and glutamine abundances remained unchanged. Aspartate and alanine, which can be converted into OAA and pyruvate respectively in a reversible reaction that requires glutamate/2-OG transamination, were also more abundant in dic2-1 after 1 h of darkness, while aspartate-derived threonine accumulated in the mutant throughout the diurnal cycle. Branched chain amino acids (BCAAs), which use 2-OG as co-substrate to initiate their catabolism in mitochondria (Hildebrandt et al., 2015), were generally higher in abundances in dic2-1 plants at night. Such increases in cellular 2-OG utilization in dic2-1 would be consistent with increases in citrate retention and subsequent 2-OG and/or glutamate formation and export by mutant mitochondria (Figure 3I). Overall, the metabolite profiles revealed that the loss of DIC2 function causes an altered 2-OG metabolism in the dark, which may provide a metabolite buffer to compensate for a defect in mitochondrial DIC2 activity while cytosolic NAD redox state is maintained.

![Figure 5](attachment://figure5.png)
metabolite abundances noted above were accompanied by a higher sucrose catabolism in the mutant at night. When leaf discs were incubated in the dark with uniformly labeled $^{14}C$-malate or $^{14}C$-leucine, we found that dic2-1 showed higher $^{14}C$O$_2$ emissions than Col-0 and dic2-1/gDIC2 from malate (Figure 6C) but not leucine (Supplemental Figure S7D). The observed increase in BCAAs accumulation in dic2-1 in the dark (Figure 6E; Supplemental Figure S8C) may thus be contributed by increased TCA cycle fluxes into biosynthetic pathways and/or an elevated proteolysis, while BCAAs breakdown for respiration remained unchanged. Thus, these data indicated that the loss of DIC2 results in increased sucrose utilization (or export for use by other tissues) and TCA cycle-facilitated respiration to compensate for a failure to maintain the homeostasis of organic acid oxidation.

**A defect in organic acid use causes accelerated leaf senescence of the DIC2 knockout during prolonged darkness**

To further examine the extent to which DIC2 transport function influences whole plant sugar-sensitive organic acid use, we subjected Arabidopsis plants to darkness for 15 days. The mutant exhibited an accelerated decline in $F_v$/$F_m$ beginning around 7 days of darkness (Supplemental Figure S8A). Leaves of dic2-1 were more yellowed compared with the Col-0 and the dic2-1/gDIC2 after 10 days (Figure 6D; Supplemental Figure S8B). When 15 days dark-treated plants were transferred back to the normal short-day cycle, only Col-0 and dic2-1/gDIC2 could recover after 7 days. To identify the TCA cycle metabolites that were affected the most by the absence of DIC2, we carried out a time-course measurement of organic acid contents over 12 days of prolonged darkness (Figure 6E; Supplemental Figure S8C). When $F_v$/$F_m$ began to decline more rapidly in dic2-1 on Day 7 (Supplemental Figure S8A), only malate, 2-OG, and citrate were significantly more accumulated. While all TCA cycle intermediates were significantly increased in the mutant 10 days after dark treatment, malate and 2-OG accumulated at least 10 times higher in abundance in the mutant than Col-0 and dic2-1/gDIC2. Citrate abundance in the Col-0 and dic2-1/gDIC2 progressively declined from Days 3 to 12 of darkness, whereas it remained unchanged in dic2-1 plants throughout the treatment. Notably, the accumulation of these metabolites by dic2-1 in planta would be consistent with a failure to properly regulate mitochondrial malate and citrate metabolism. Throughout a diurnal cycle, pool sizes of these metabolites were unchanged; it was only when plants were exposed to dark-induced starvation that altered patterns in the utilization of specific organic acids manifested (Figure 6A).

DIC2 modulates metabolite turnover through TCA cycle and amino acid metabolism, which supports citrate export at night

The observed decrease in malate-dependent O$_2$ consumption by isolated mitochondria (Supplemental Figure S4C) did not explain the faster dark respiration of intact leaves (Figures 5, E and 6, C). This could be due to the absence of any extramitochondrial metabolism, which in vivo maintains metabolite supply for sustaining mitochondrial transport activities, TCA cycle and respiration in response to rearranged metabolism and transport in the mutant. To account for the apparent homeostasis in metabolite pool sizes (Figure 5A), we postulated that there could be a flux change in certain steps of metabolism to compensate for the reduced malate import and citrate export by dic2-1 mitochondria.

To determine if dic2-1 metabolizes carbon differently, we traced the labeling of U-$^{13}C$-glucose into the TCA cycle and closely related amino acids in leaf discs in the dark for 8 h. The $^{13}C$-tracing data were expressed on the basis of per O$_2$ molecule consumed to take into consideration the observed differences in organic acid use-dependent CO$_2$ release and dark respiration rate (Supplemental Data Set S1). Consistent with DIC2 function as a mitochondrial citrate exporter, dic2-1 displayed a decreased rate of citrate labeling over the course of dark incubation (Figure 7A). On the other hand, the citrate pool remained stable in the dark in dic2-1 (Figure 6A) possibly due to compensation by altered citrate turnover rates in other compartments. A decrease in OAA availability from cytosolic phosphoenolpyruvate (PEP) carboxylase could also be ruled out, since labeling of m + 3 aspartate (a proxy for labeled OAA) was not altered (Supplemental Figure S9, A and B). Decreased citrate labeling was accompanied by a significant increase in the abundance of labeled 2-OG due to a higher total pool in the mutant (Figure 7B; Supplemental Data Set S1), which coincided with an increased rate of $^{13}C$ incorporation into glutamine (Figure 7C). These increases could be facilitated by a higher mitochondrial glutamate efflux rate since there was a higher initial 2-OG accumulation rate in substrate-fed dic2-1 mitochondria (Supplemental Figure S4, A and B), implying that an enhanced flux into glutamine was necessary to remove excess mitochondrial 2-OG by mitochondrial glutamate transamination reactions in concert with a cytosolic glutamine synthase (Bernard and Habash, 2009). The downstream extra-mitochondrial biosynthetic pathway of aspartate, threonine, and asparagine, were increased in total label abundances in the mutant mainly due to a significantly higher incorporation as m + 2 species via the TCA cycle (Figure 7, D and E; Supplemental Figure S9, C–H and Supplemental Data Set S1), while the total amount of succinate, fumarate, and malate labeled via the TCA cycle (m + 2) and/or PEP-OAA interconversion (m + 3) did not change (Figure 7F; Supplemental Figure S9, I–L). These results are consistent with a metabolic diversion of excess malate that was not consumed by mitochondria for citrate synthesis, export, and storage into aspartate.

The loss of DIC2 causes a significant metabolic rearrangement towards amino acid synthesis with less citrate available from the TCA cycle for storage at night (Supplemental Figure S9M). To further examine if citrate
Figure 6 Quantitative analysis of metabolites associated with the TCA cycle in a diurnal cycle and during prolonged darkness in the DIC2 knockout mutant. A. Plants were grown under short day conditions for 6 weeks and leaf discs were collected at 1, 4, 8, 12, and 15 h after dark shift and 1, 4, and 7 h after light shift. Metabolites in this figure were analyzed by LC–MRM–MS (n ≥ 6). Metabolites are colored according to their accumulation pattern in dic2-1 in a diurnal cycle: Orange, accumulates at night; Brown, accumulates during light and dark shift; Green, accumulates predominantly during the day; Blue, accumulates throughout a diurnal cycle; Purple, accumulates only after the first hour of dark shift; gray, metabolite not measured. B. Sucrose levels in leaf discs from Col-0, dic2-1, and dic2-1/gDIC2 collected at different time points of a diurnal cycle as quantitatively determined by GC–MS against authentic standards (n ≥ 6). C. CO2 evolution of leaf discs incubated in uniformly labeled 14C-malate (1, 4, and 7 h after light shift. Metabolites in this figure were analyzed by LC–MRM–MS (Figure 6). D. Images of 5- to 6-week-old, short-day grown plants treated with 0 and 10 days of extended darkness. Values for maximum quantum efficiency of photosystem II (Fv/Fm) are shown below each genotype (n = 6). E. Changes in citrate, 2-OG and malate content in Arabidopsis plants after 0, 3, 7, 10, and 12 days of extended darkness treatment (n ≥ 6). Individual data points were overlaid in dots in bar graphs. Each data point represents mean and each error bar represents ± se where appropriate. Asterisks indicate a significant change for dic2-1 versus Col-0 and dic2-1 versus dic2-1/gDIC2 comparisons as determined by one-way ANOVA with Tukey post hoc test (*P < 0.05; **P < 0.01).
production and/or consumption by non-TCA cycle pathways could be altered to compensate for the reduction in mitochondrial citrate supply when DIC2 is absent, we measured the transcript levels of peroxisomal citrate synthase (CYS2 and CYS3; Pracharoenwattana et al., 2005) and cytosolic ATP-dependent citrate lyase (ACLA-1, ACLA-2, ACLA-3, ACLB-1, and ACLB-2; Fatland et al., 2002) in leaves collected at 15 h into the dark photoperiod (Figure 7G). We found that the expression of both CYS2 and CYS3 was significantly higher in dic2-1 than Col-0 and dic2-1/gDIC2 by two-fold. In contrast, the expression of ACL genes in the mutant did not show consistent change, with only ACLA-2 and ACLA-3 exhibiting significant decrease and increase in transcript abundance, respectively, in the mutant when compared to other genotypes. If we assume that the expression of genes in citrate metabolism is controlled by citrate in a similar manner to other metabolic pathways observed previously (Gibon et al., 2006), it is estimated that the loss of DIC2 triggered a change in citrate contribution from 61% mitochondria to 64% peroxisomes (Supplemental Figure S9M).
In summary, these data helped to explain how a defect in mitochondrial malate/citrate exchange in \textit{dic2-1} is significantly overcome in a whole plant metabolic context to establish day and night homeostasis and retain a viable plant albeit with stunted growth rate. Only at day to night transitions and prolonged darkness do the consequences of these unusual metabolic fluxes yield temporary gross imbalances in metabolite pools.

**Discussion**

Malate import and citrate export are two of the most critical fluxes that mitochondria contribute to the rest of the cellular metabolic network in plants (Fernie et al., 2004; Sweetlove et al., 2010; Lee et al., 2016). Malate plays a pivotal role in mediating metabolic redox exchange between cellular compartments (Scheibe, 2004; Selinski and Scheibe, 2019; Zhao et al., 2020), while citrate produced in mitochondria is the major source of cellular citrate when the TCA cycle operates at night in mature green leaves. To date, several mitochondrial malate carriers, including DIC2, were found to have a broad substrate preference for dicarboxylates, and limited ability to transport tricarboxylates and/or amino acids with close resemblance to dicarboxylates in vitro (Picault et al., 2002; Palmieri et al., 2008; Monné et al., 2018). However, neither their actual contributions to mitochondrial metabolite transport nor the knockon consequences of their loss on mitochondrial metabolism in organello or in planta have been verified. Determining if these carriers are generalists or specialists in vivo and how they influence metabolism is critical in any future attempt to modify mitochondrial substrate use, or to understand if transport is a point of control in metabolic models of plant cell function. Here, we redefined DIC2, not as a generalist but as a specialist high-affinity mitochondrial malate/citrate carrier influencing both these functions in mitochondrial anaplerotic metabolism based on multiple lines of in vivo and in vitro evidence (Figure 8). While our data reveal that other carriers can facilitate these two functions, they are insufficient to maintain metabolism unperturbed. Transport of sulfate, phosphate, and other dicarboxylates by DIC2 may occur at low affinity under in vitro or specific conditions that enable these metabolites to accumulate at several magnitudes above malate and citrate levels; however, such activity is unlikely to occur in vivo given that malate and citrate are primary substrates for redox valves to connect subcellular compartments and the preferred chemical species for organic acid storage (Igamberdiev and Eprintsev, 2016).

Given the prior evidence of broad DIC1 and DIC3 substrate preference in vitro and high amino acid identity

![Figure 8 Schematic model of mitochondrial malate–citrate exchange for the maintenance of citrate metabolism at night. In wild-type leaves (left), DIC2 facilitates the export of citrate in exchange for malate (thick purple line) and operates in conjunction with other organic acid transporters (including other malate and citrate carriers) to support mitochondrial anaplerotic metabolism in the dark. Citrate exported from mitochondria is mostly transferred to the vacuole, with the remainder being converted to acetyl-CoA in the cytosol for the synthesis of secondary metabolism products, such as isoprenoids and flavonoids. The DIC2 loss-of-function mutant (\textit{dic2-1}, right) significantly reduces citrate export from mitochondria but has little impact on cellular organic acid pools, including citrate. This is due to the activation of a number of compensatory mechanisms to minimize major metabolic disruptions, including (in green solid lines for single-step reactions or dashed lines for multistep pathways): increased citrate supply from \(\beta\)-oxidation of fatty acids in peroxisomes, faster sucrose depletion rate to increase the overall carbon supply, a higher rate of malate transport by non-DIC2 carriers and diversions of excess mitochondrial citrate and cytosolic malate into amino acid biosynthesis. However, these metabolic rearrangements fail to alleviate or cause the stunted phenotype of \textit{dic2-1} plants, indicating that DIC2 activity is important for maintaining an overall metabolic homeostasis, particularly at night and during starvation.](image-url)
among the DIC homologs, our finding that they could not complement the dic2-1 phenotype (Figure 2B) was unexpected. However, it is not unprecedented; for example, PAPST1 and PAPST2, which are 78% identical in amino acid sequence, are responsible for 3'-phosphoadenosine 5'-phosphosulfate (PAPS)/3'-phosphoadenosine 5'-phosphate (PAP) transport and PAP/ATP(D)P exchange, respectively, based on both in vitro and in vivo evidence (Gigolashvili et al., 2012; Ashykzhmina et al., 2019). In a similar way, while a previous proteoliposome-based kinetic study showed that Arabidopsis DIC1 and DIC2, the closest homologs of yeast DIC, transport a variety of different dicarboxylates (Palmieri et al., 2008), such an approach alone could not accurately predict apparent substrate affinity and specificity (Figure 4), directionality of transport in intact mitochondria (Figure 3) or the in vivo consequences of DIC2 loss on transport or metabolism of other organic acids (Figures 5–7). Further experiments are required to assess how exactly DIC1 and DIC3 differ from DIC2 in terms of substrate preferences and transport kinetics.

Given the importance of mitochondria-supplied citrate to metabolism in green tissues (Gauthier et al., 2010; Cheung et al., 2014; Igamberdiev, 2020), it is expected that a complete block in mitochondrial citrate synthesis and export would result in severe reduction in post-germination vegetative growth. Indeed, a simultaneous loss of NAD-ME, mitochondrial pyruvate carrier and alanine aminotransferase activities severely disrupts seedling development, most likely due to limited supply of pyruvate for mitochondrial citrate synthesis (Le et al., 2021a). Mutants lacking the activity in one of the subunits of mitochondrial pyruvate dehydrogenase complex (mPDC) or both mMDH isozymes, which catalyze acetyl-CoA and OAA formation, respectively, required for citrate synthesis, are significantly slower in vegetative growth (Tomaz et al., 2010; Yu et al., 2012; Ohbayashi et al., 2019). Here, we also observed a reduction in vegetative growth in dic2-1, with decreases in citrate export by isolated mitochondria (Figure 3.H and K) and in vivo 13C-glucose labeling into citrate (Figure 7A) when compared to the wild-type. Unlike mMDH and mPDC mutants, however, the steady-state pool of organic acids, including citrate, was surprisingly stable in dic2-1 (Figure 5A). It is very likely that the increase in citrate retention in the dic2-1 matrix triggered an increase in nighttime sucrose depletion (Figure 6B) to maintain the organic acid pools, resulting in higher RN and malate-dependent CO2 evolution (Figures 5, E and 6, C). Another possible pathway to compensate for a reduction in mitochondrial citrate supply is by altering fatty acid turnover via the last step of peroxisomal β-oxidation (Figure 8), given that dic2-1 phenotypes are observed mostly during darkness and the transcripts of two peroxisomal citrate synthases (CYS2 and CYS3) are upregulated in the mutant (Figure 7G) and are more abundant in wild-type plants at night (Gibson et al., 2004; Smith et al., 2004) and during prolonged darkness (van der Graaff et al., 2006). However, possible side effects of such a strategy include a loss of synchronization in fatty acid turnover with peroxisomal citrate synthesis and export, higher triacylglycerol utilization, and lipid peroxidation (Fan et al., 2017), which reasonably explain the increased susceptibility of dic2-1 to dark-induced senescence (Figure 6, D and E). The requirement for an equilibrium between organelar citrate synthesis and fatty acid breakdown to prevent excess reactive oxygen species accumulation would also explain why a strong DIC2 transcriptional response is commonly observed during abiotic stress treatments, including touch and sound vibration (Ghosh et al., 2016; Van Aken et al., 2016), phosphate deficiency (Lin et al., 2011), light-to-dark transition (Lee et al., 2005), and cold, drought, and UV stresses (Kilian et al., 2007). Overall, our data strongly indicate that DIC2 transport function is integrated into whole plant metabolism as an important supplier of citrate for nighttime vacuole storage, and its absence severely limits metabolic flexibility for maintaining energy supply and redox balance in response to a sudden dark shift or a prolonged sugar starvation.

The existence of multiple modes of TCA cycle substrate transport has been proposed (Wiskich, 1977; LaNoue and Schoolwerth, 1979) but the lack of clarity as to the identity of components responsible has hindered our understanding on how they cooperate to support the integration of mitochondrial carbon metabolism into the wider cellular metabolic network (Lee and Millar, 2016). In the case of dic2-1, a residual rate of mitochondrial malate import and citrate export is maintained, suggesting that other carriers exist to partially compensate for the absence of DIC2 activity. It is also remarkable that in vivo cytosolic NADH/NAD+ ratios in dic2-1 leaves did not differ from wild-type throughout the night with only a glimpse of change during a light-to-dark transition (Figure 5, C and D). The homeostasis of cytosolic NAD redox state in dic2-1 could be maintained, among other possible strategies, by (1) diverting of excess organic acids to energy-consuming biosynthesis of specific amino acids (Figures 7, G and 8) and/or (2) remobilizing excess NADH into the intermembrane space of mitochondria where it can be oxidized by external NADH dehydrogenases, thereby raising leaf respiration in the dark (Figure 5E). Studies of mitochondrial carrier mutants reported to date have been unable to directly link growth and/or metabolic phenotype with gene function (Toka et al., 2010; de Souza Chaves et al., 2019; Feitosa-Araujo et al., 2020). It is possible that, apart from functional redundancy, changes in metabolic fluxes occur without altering the overall metabolite pool or physiology, which can only be captured by sensitive measurements under specific external conditions.

In conclusion, we have refined the current understanding of how DIC2 influences mitochondrial organic acid transport and metabolism in Arabidopsis, and pinpoint how its absence leads to growth retardation. The importance of mitochondrial organic acid exchange carriers in plants has been discussed for decades, but it is only in the last decade that their identities are beginning to be unraveled through heterologous systems. By utilizing a systems approach based on reverse genetics, we demonstrate that a thorough carrier analysis in heterologous systems is only the start to reveal the in vivo role of a carrier such as DIC2. Given the lack of
genetic clarity of many plant mitochondrial carriers, it is reasonable to suspect that the true substrate specificities and in vivo metabolic influence of many other carriers have yet to be unraveled. A complete understanding of the identities of mitochondrial metabolite carriers and their substrate preferences with kinetic characteristics and transport orientation in vivo will assist in improving metabolic models, as well as refining their individual role in underpinning the remarkable flexibility of plant cell metabolism.

Materials and methods

Plant material and growth conditions

The DIC2 T-DNA insertion lines were obtained from GABI-Kat (dic2-1 GK-217B05, dic2-2 GK-833F11, and dic2-3 GK-047F03; Kleinboelting et al., 2012). These T-DNA insertion lines were screened for homozygous insertion by PCR of genomic DNA using the primer pairs according to Supplemental Table S3. Inverse PCR was carried out according to Teschner et al. (2010). The construct for genomic complementation with transgene gDIC2 was generated by amplifying the DIC2 genomic region (including ~2-kb upstream of start codon and ~1-kb downstream of stop codon) from wild-type Col-0 DNA, and the resulting ~4-kb fragment was cloned into pCAMBIA1380. Overexpression constructs were generated by amplifying the coding sequence of DIC2, DIC1, or DIC3 and inserting each of these fragments into pB7WG2 by Gateway cloning. The DIC2-GFP construct was generated by inserting DIC2 coding sequence without the stop codon into pUBC-DEST (Grefen et al., 2010) by Gateway cloning. All cloning primers were listed in Supplemental Table S3. Agrobacterium-mediated transformation of each construct was carried out by floral dipping (Zhang et al., 2006).

Arabidopsis seedlings and plants were grown under a light intensity of 100 μmol m⁻² s⁻¹ (tubular fluorescent lighting) after seeds were stratified at 4°C in the dark for 2–3 days. Plants were grown on compost supplemented with Perlite and Vermiculite (3:1:1) under continuous, short, or long photoperiod as indicated. To analyze seedling phenotypes, seeds were surface-sterilized, stratified and grown vertically for 10 days on agar plates containing half-strength MS medium, 2 mM 2-(N-morpholino)ethanesulfonic acid (MES) at pH 5.7 and 1% (w/v) agar under short-day conditions. For the isolation of mitochondria, surface-sterilized seeds were grown in Murashige and Skoog (MS) medium (1/2-strength MS medium, 2 mM MES, pH 5.7) supplemented with 1% (w/v) sucrose and 0.1% (w/v) agar for 14–16 days with gentle agitation (40–60 rpm) under long-day conditions.

Subcellular localization analysis by confocal laser scanning microscopy

A Nikon A1Si confocal microscope equipped with the following excitation and emission wavelength settings was used: 488-nm excitation and 525-nm emission for GFP, 560-nm excitation and 595-nm emission for TMRM, and 640-nm excitation and 700-nm emission for chlorophyll autofluorescence. DIC2 localization images were acquired by a NIS element AR software package (version 4.13.01, Build 916) using a 20× lens (Nikon CFI Plan Apo VC 20x 0.75 N.A.) and a 60× lens (Nikon CFI Plan Apo VC 60x 1.20 N.A. WA) and were processed using ImageJ software package.

Transcript analysis by quantitative real-time polymerase chain reaction

RNA isolation was performed using the Spectrum Plant Total RNA Kit (Sigma-Aldrich) in conjunction with on-column DNAse treatment to remove genomic DNA. cDNA was synthesized using the iScript cDNA Synthesis Kit (Bio-rad). Quantitative polymerase chain reaction was performed in a LightCycler 480 II Real Time PCR System (Roche) using the QuantNova SYBR Green RT-PCR Kit (Qiagen) according to the manufacturer’s instructions with reference genes elongation factor-1α (At5g60390) and SAND family protein (At2g28390; Czechowski et al., 2005). Relative transcript abundance was obtained by normalizing to two reference genes and the normalized values were then combined by geometric mean. All primers are listed in Supplemental Table S3.

Isolation of mitochondria, O₂ electrode measurements, enzyme activity assays, blue native electrophoresis, and metabolite uptake assays by silicone oil centrifugation

Mitochondria were isolated from 2-week-old Arabidopsis seedlings as described previously (Sweetlove et al., 2007). Single substrate-dependent O₂ consumption by purified mitochondria was measured at pH 6.8 in a computer-controlled Clark-type O₂ electrode unit according to Lee et al. (2010). In vitro activities of TCA cycle enzymes in isolated mitochondria were measured as described by Huang et al. (2015).

Digitonin-solubilized mitochondrial proteins were separated on a blue-native polyacrylamide gel electrophoresis (PAGE) according to Eubel et al. (2003), using a 4.5%–16% gradient. In-gel Complex I activity staining was carried out as outlined by Sabar et al. (2005).

Time-course measurements of substrate uptake by isolated mitochondria were carried out according to Lee et al. (2014) with modifications specified in Le et al. (2021b). Briefly, for uptake in the absence of cofactors, freshly prepared mitochondria (100 μg) were resuspended in 200-μL minimal transport buffer (250-MM sucrose, 10-MM TES, 0.5-MM ATP, pH 7.2). Uptake assays were initiated by adding radiolabeled U-[¹⁴C]-malate (200 μM, Perkin Elmer) at 25°C. After the specified incubation time, the reaction was stopped immediately by rapid centrifugation (at 13,000g for 3 min at room temperature) through a 90-μL silicone oil layer (AR200, Sigma Aldrich) into the bottom sedimentation layer containing 20 μL of 10% (v/v) perchloric acid. Fractions above (supernatant) and below (containing the mitochondria pellet) the silicone oil interface were transferred to separate scintillation vials and mixed with 3 mL of Ultima Gold scintillation fluid (PerkinElmer). Radioactivity...
was detected by a Beckman LS 6500 Scintillation Counter. For energized uptake, isolated mitochondria (100 μg) were resuspended in 190-μL respiratory transport medium (225-mM sucrose, 5-mM KH₂PO₄, 10-mM TES, 10-mM NaCl, 4-mM MgSO₄, 0.1% (w/v) BSA, 2-mM NAD, 12-μM CoA, 0.2-mM thiamine pyrophosphate, and 0.5-mM ADP, pH 6.8) and uptake was initiated by adding 50-μM or 500-μM substrate as stated. Reactions were stopped as described above, except the bottom layer contained 20 μL of 0.5-M sucrose, pH 1.5. Fractions above (extra-mitochondrial space) and bottom (pellet, mitochondria) were collected, and metabolites were methanol-extracted and detected by selective reaction monitoring mass spectrometry (liquid chromatography–selective reaction monitoring–mass spectrometry [LC–SRM–MS], see below).

**Proteoliposome transport assay**

The coding sequence of DIC2 without the stop codon was introduced into the yeast expression vector pYES-DEST52 (ThermoFisher Scientific) by Gateway Cloning. pYES-DIC2 or pYES-DEST52 (empty vector control) was transformed into the uracil-auxotrophic Saccharomyces cerevisiae strain INVSc1 (ThermoFisher Scientific) by lithium acetate transformation. To prepare microsomes, 500 mL of transformed yeast culture was inoculated overnight at 30°C according to Rautengarten et al. (2016) with slight modifications. Briefly, yeast pellet was washed several times in H₂O and was resuspended in 10-mL resuspension buffer (50-mM potassium phosphate, pH 7.1, 1.4-M sorbitol, 10-mM NaN₃, and 40-mM 2-mercaptoethanol) supplemented with 2,000 units of Lyticase (Sigma-Aldrich), followed by a 37°C incubation for 1 h with gentle shaking. The resulting spheroplasts were collected by centrifugation at 2,500g for 5 min at room temperature, washed once with wash buffer (0.8-M sorbitol, 10-mM triethanolamine/acetic acid (pH 7.2), 1-mM EDTA, 1-mM PMSF, and protease inhibitor tablet [Roche]) and lysed in 5-mL wash buffer using a Dounce homogenizer. The lysate was centrifuged at 1,500g for 5 min at 4°C to remove pellet containing large debris and the supernatant was centrifuged again at 3,000g for 5 min at 4°C. The resulting supernatant was centrifuged at 100,000g for 75 min to collect the crude microsomal fraction. This fraction was resuspended in reconstitution buffer (10-mM Tricine-KOH [pH 6.0], 50-mM potassium glutonate, and 20% glycerol).

For liposome preparation, 600 mg of soybean L-α-phosphatidylcholine (Sigma) was dissolved in 10-mL chloroform and evaporated under a stream of nitrogen for at least 45 min. The resultant film was resuspended for 2 h at 42°C in reconstitution buffer (10-mM Tricine-KOH, pH 6.0, 50-mM potassium glutonate, and 20% glycerol). Prior to reconstitution, solubilized L-α-phosphatidylcholine was extruded at 42°C using an Avanti Mini-Extruder (Avanti Polar Lipids) fitted with a 100-nm polycarbonate membrane. Approximately 400 μg microsomal proteins were reconstituted into extruded lipid at 1:13 (lipid: protein) in the presence of 10-mM internal loading substrate, 50-mM octyl-β-glucoside, and Bio-beads (BioRad Laboratories) with gentle agitation for 2 h at room temperature. Unincorporated components were removed from substrate-loaded proteoliposomes by passing through Sephadex G50 (medium, GE Healthcare) several times.

Aliquots of 200-μL proteoliposomes were incubated with specified substrates (pH adjusted to 6.0) at 25°C for indicated times. Substrates that remained on the outside of proteoliposomes were rapidly removed by passing the reaction through a pre-packed 2-mL Sephadex G50 column. Proteins and metabolites in the resulting supernatant were extracted by chloroform/methanol extraction. Following liquid removal and sample resuspension, 3-nitrophenylhydrazine-derivatised metabolites were quantified by LC–SRM–MS (see below). Extracted proteins were separated on mini-PROTEAN TGX stain-free gels and visualized directly without staining using the Gel Doc XR+ with Image Lab software (Bio-Rad Laboratories). Proteins were then transferred onto a PVDF membrane for immunodetection with the monoclonal anti-V5 antibodies (1:10,000 dilution; ThermoFisher Scientific, Catalogue number R96025) using the Amersham Imager 680 (GE Healthcare).

**Leaf gas exchange, chlorophyll fluorescence, and respiration measurements**

Measurements of gas-exchange parameters were carried out using a LI-6400 XT infrared gas analyser (Li-Cor). After at least 2 h of illumination, measurements were carried out using fully developed leaves from at least 8-week-old short-day grown plants. Prior to each measurement, leaf was acclimated in the enclosed 60-mm² chamber to 22°C, relative humidity 70%, and a CO₂ concentration of 400 ppm with light intensity of 250 μmol m⁻² s⁻¹. A series of light intensities (0, 50, 100, 250, 500, 700 μmol m⁻² s⁻¹) was applied and the following parameters were then recorded: CO₂ assimilation rate, stomatal conductance, and transpiration rate. For the analysis of post-illumination bursts, leaf was equilibrated in a chamber to 22°C and relative humidity 70%, with light intensity of 1,000 μmol m⁻² s⁻¹ and CO₂ concentration of 100 ppm. CO₂ assimilation rate in the light was monitored for 3 min before dark shift (i.e. photosynthetic active radiation [PAR] = 0 μmol m⁻² s⁻¹) and CO₂ evolution rate was monitored for 2.5 min.

Chlorophyll fluorescence measurements were carried out using the IMAGING-PAM Maxi Version Chlorophyll Fluorometer (WALZ). Plants were acclimated in the dark for at least 15 min before Fv/Fm, PSII quantum yield during illumination (Y(II)), photosynthetic electron transport rate, and nonphotochemical quenching were measured at increasing actinic light intensities and plotted as light response curves. Dark respiration was measured in dark-adapted, fully expanded leaves using a Q2 O₂ sensor (Astec-Global) as previously described (O’Leary et al., 2017). O₂ concentration in a sealed 850-μL capacity tube with a single leaf disc (6.5-mm diameter, with a total of 40–60 mg fresh weight) floated on leaf respiration buffer (10-mM HEPES, 10-mM MES, and 0.2-mM CaCl₂·2H₂O, pH 7.2) was measured. For leaf ¹⁴CO₂
evolution measurements, leaf discs were floated on leaf respiration buffer containing ~0.5 KBq mL\(^{-1}\) of U-\(^{14}\)C-malate or U-\(^{14}\)C-leucine (Perkin Elmer). Evolved \(^{14}\)CO\(_2\) trapped in KOH-soaked filter paper and total radiolabel extracted from leaf discs were measured by a liquid scintillation counter (Beckman Coulter).

**Analysis of NAD redox dynamics**

All experiments were carried out with leaf discs of 6-week-old *A. thaliana* plants grown under long-day conditions (16-h light/8-h dark). Microscopy procedures described in Wagner et al. (2015) were used as guideline for ratiometric sensor monitoring with the modification that samples were mounted between two cover slips (22 \(\times\) 40 mm, VWR). Dynamic changes under dark-light conditions were monitored as described by Elsaesser et al. (2020) with a confocal laser scanning microscope (Zeiss LSM 780, attached to an Axio Observer.Z1; Carl Zeiss Microscopy) using the 10 \(\times\) objective (Plan-Apochromat, 0.3 M27). Excitation was set at 405 nm (T-Sapphire) and 543 nm (mCherry), emission was collected at 499–544 nm (T-Sapphire) and 588–624 nm (mCherry). The ZEN Experiment Designer software module (Carl Zeiss) was used to program time series with varying scanning cycles and illumination times. A custom-made device with an implemented cold white LED stripe (Pferdekaemper, Menden, Germany) was used to apply actinic light with an intensity of 220 \(\mu\)mol m\(^{-2}\) s\(^{-1}\). Time series with Pperedox-mCherry were performed with the following regime: (1) 15 min in the dark of 30 scans with 30-s interval; (2) 15 min of illumination with 8 loops of 15-s light followed by 3.2-s scan and 26 loops of 30 s light followed by 3.2-s scan; (3) 15 min in the dark of 30 scans with 30-s interval.

For in vivo measurements of NAD redox changes, leaf discs stably expressing cytosolic Pperedox-mCherry were subjected to multiwell plate reader-based fluorimetry, as described in de Col et al. (2017). Plant material was submerged in 200-\(\mu\)L assay medium (10-mM HEPES, 10-mM MES, and 0.2-mM CaCl\(_2\), pH 7.2) in a transparent 96-well plate (NUNC). Pperedox-mCherry fluorescence was recorded in a CLARIOstar microplate reader (BMG Labtech, Ortenberg, Germany). The chromophores T-Sapphire and mCherry were excited at 400 \(\pm\) 10 nm and 570 \(\pm\) 10 nm, and emission collected at 515 \(\pm\) 7.5 nm and 610 \(\pm\) 5 nm, respectively. Samples were analyzed employing top optics with focal height of 8.0 mm, incubation temperature at 25°C, well-multichromatic monitoring. 50 flashes per cycle and double orbital shaking at 400 rpm for 10 s before each measurement cycle. Background fluorescence of control plants without sensor expression was recorded in parallel and subtracted from all data before analysis. Mounting of the leaf discs was performed, except for the use of a green LED head torch, in the dark at the end of the night shortly prior to the night-to-day transition. After recording the baseline, plates were exposed to actinic light under white LEDs with a photon flux density of 120 \(\mu\)mol m\(^{-2}\) s\(^{-1}\) for in total 8 h, except for 3-min-long interruptions allowing for fluorometric sensor recording at indicated time points. For the subsequent dark phase, the plate was kept inside the plate reader (dark) and measurements were performed every 30 min.

**Metabolite extraction from plant tissues**

Plant tissues (~25 mg for general metabolite analysis or ~50 mg for isotope labeling experiments) were collected at specified time points and immediately snap-frozen in liquid nitrogen. Samples were ground to fine powder and 500 \(\mu\)L of cold metabolite extraction solution (90% [v/v] methanol, spiked with 2-\(mg\)-\(\mu\)L\(^{-1}\) ribitol, 6-\(mg\)-\(\mu\)L\(^{-1}\) adipic acid, and 2-\(mg\)-\(\mu\)L\(^{-1}\) and \(^{13}\)C-leucine as internal standards). Samples were immediately vortexed and shaken at 1,400 rpm for 20 min at 75°C. Cell debris was removed by centrifugation at 20,000g for 5 min at room temperature. For each sample, 100 \(\mu\)L (400 \(\mu\)L for isotope labeling experiments) of supernatant was transferred to a new tube and either proceeded to derivatization for LC–MS analysis or dried using a SpeedVac.

**Analyses of sugars by gas chromatography–mass spectrometry**

For gas chromatography–mass spectrometry analysis of sugars, dried samples were dissolved in 20 \(\mu\)L of pyridine in methoxyamine hydrochloride (20 mg mL\(^{-1}\)) and incubated at 30°C for 90 min at 1,200 rpm. Then, 30 \(\muL\) of N-methyl-N-(trimethylsilyl)-trifluoroacetamide was added to each sample and allowed to react for 30 min at 37°C at 1,400 rpm. The derivatized metabolite samples were analyzed by an Agilent 6890 gas chromatograph coupled with a 7683B Automatic Liquid Sampler and a 5973N mass selective detector, and fitted with an Agilent VF-5ms capillary column (30 \(\mu\)m \(\times\) 0.25 mm, 0.25 mm internal diameter) and a 10 m integrated guard column. The helium carrier gas flow rate was 1 mL·min\(^{-1}\). For the 58.5-min temperature gradient, the GC oven was held at the initial temperature of 70°C for 1 min, increased to 76°C at the rate of 1°C·min\(^{-1}\) and then to 325°C at the rate of 6°C·min\(^{-1}\) where it was held at 325°C for 10 min. The inlet temperature, MS source temperature and quadrupole temperature were 300°C, 230°C, and 150°C, respectively. The MS detector m/z scan range was 40–600. Data were analyzed using MassHunter Quantitative Analysis Software (version B07.01, Build 7.1524.0). Metabolites were quantified by comparing the integrated peak area with a calibration curve obtained using authentic standards, and normalized against fresh weight and ribitol internal standard.

**Analyses of organic acids and amino acids by multiple reaction monitoring using LC–SRM–MS**

For LC–MS analysis of organic acids, sample derivatization was carried out based on a previously published method with modifications (Han et al., 2013). Briefly, for each 100 \(\muL\) of sample, 50 \(\muL\) of 250-mM 3-nitrophenylhydrazine in 50% methanol, 50 \(\muL\) of 150-mM 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide in methanol, and 50 \(\muL\) of 7.5% pyridine in 75% methanol were mixed and allowed to react on ice for 60 min. To terminate the reaction, 50 \(\muL\) of 2-mg mL\(^{-1}\) butylated-hydroxytoluene in methanol was added,
followed by the addition of 700 μL of water. Derivatized organic acids were separated on a Phenomenex Kinetex XB-C18 column (50 × 2.1 mm, 5 μm particle size) using 0.1% (v/v) formic acid in water (solvent A) and methanol with 0.1% formic acid (solvent B) as the mobile phase. The elution gradient was 18% B at 1 min, 90% B at 10 min, 100% B at 11 min, 100% B at 12 min, 18% B at 13 min, and 18% B at 20 min. The column flow rate was 0.3 mL-min⁻¹, and the column temperature was maintained at 40°C. The Agilent 6430 Triple Quadrupole mass spectrometer (QQQ-MS) was operated in negative ion mode in the SRM mode.

For measuring amino acids, dried samples were resuspended in 500-μL HPLC-grade water before they were filtered to remove insoluble debris. Metabolites were separated on an Agilent Poroshell 120 Bonus-BP column (100 × 2.1 mm, 2.7-μm internal diameter) using 0.1% formic acid in water (solvent A) and acetonitrile with 0.1% formic acid (solvent B) as the mobile phase. For the analysis of amino acids and sugars, the elution gradient was 0% B at 1 min, 1% B at 4 min, 10% B at 6 min, 100% B at 6.5 min, 100% B at 8 min, 0% B at 8.5 min, and 0% B at 15 min. The column flow rate was 0.25 mL-min⁻¹, and the column temperature was kept at 40°C. The QQQ-MS was operated in positive ion mode in SRM mode.

A 0.5 μL or a 15 μL aliquot of each sample was injected and analyzed by an Agilent 1100 HPLC system coupled to a QQQ-MS equipped with an electrospray ion source. Data acquisition and LC-MS control were done using the Agilent MassHunter Data Acquisition software (version B.02.00). Separation of metabolites was performed using a Luna C18 column (Phenomenex; 150 × 2 mm, 3-μm particle size). The mobile phase consisted of 97:3 water: methanol with 10-mM tributylamine and 15-mM acetic acid (solvent A) and 100% methanol (solvent B). The gradient program was 0% B 0 min, 1% B 5 min, 5% B 15 min, 10% B 22 min, 25% B 24 min, 27% B 35 min, 29% B 80 min, 95% B 81 min, 95% B 82 min, 0% B 83 min, and 0% B 97 min. The flow rate was 0.2 mL-min⁻¹, with column temperature kept at 35°C, autosampler was cooled to 10°C and injection volume was 30 μL. The Q-TOF was operated in MS mode with negative ion polarity using the following operation settings: capillary voltage, 4,000 V; drying N2 gas and temperature, 10 L-min⁻¹ and 250°C respectively; Nebulizer, 30 psi. Fragmentor, skimmer, and octopole radio frequency (Oct1 RF Vpp) voltages were set to 110 V, 65 V, and 750 V, respectively. The scan range was 70–1,200 m/z and spectra were collected at 4.4 spectra/s, which corresponded to 2,148 transients/spectrum. MS scan peaks of all the possible mass isotopologs were integrated using MassHunter Quantitative Analysis Software (version B.07.01, Build 7.1.524.0). Extracted peak matrices were processed using IsoCor (Millard et al., 2019) to correct for the contribution of naturally occurring isotopes. Relative isotopomer abundance for each metabolite and percentages of 13C enrichment were calculated according to Araújo et al. (2014). Metabolite quantitation was carried out based on calibration curves obtained with unlabeled authentic standards.

Dynamic 13C-glucose labeling of Arabidopsis leaf discs and analysis of labeled metabolites by time-of-flight mass spectrometry

Leaf discs (~50 mg) were prepared from short-day grown (8-h light/16-h dark) plants 1 h before the end of a normal light photoperiod. They were floated on leaf respiratory buffer containing 20-mM U-13C-glucose (99% purity, Sigma Aldrich). At the specified incubation time, leaf discs were briefly washed with respiratory buffer to remove excess labeled glucose and frozen in liquid nitrogen for metabolite extraction as stated above.

Analyses of unlabeled and labeled metabolites were performed using an Agilent 1100 HPLC system coupled to an Agilent 6510 Quadrupole/Time-of-Flight (Q-TOF) mass spectrometer equipped with an electrospray ion source. Data acquisition and LC-MS control were carried out using the Agilent MassHunter Data Acquisition software (version B.02.00). Separation of metabolites was performed using a Luna C18 column (Phenomenex; 150 × 2 mm, 3-μm particle size). The mobile phase consisted of 97:3 water: methanol with 10-mM tributylamine and 15-mM acetic acid (solvent A) and 100% methanol (solvent B). The gradient program was 0% B 0 min, 1% B 5 min, 5% B 15 min, 10% B 22 min, 25% B 24 min, 27% B 35 min, 29% B 80 min, 95% B 81 min, 95% B 82 min, 0% B 83 min, and 0% B 97 min. The flow rate was 0.2 mL-min⁻¹, with column temperature kept at 35°C, autosampler was cooled to 10°C and injection volume was 30 μL. The Q-TOF was operated in MS mode with negative ion polarity using the following operation settings: capillary voltage, 4,000 V; drying N2 gas and temperature, 10 L-min⁻¹ and 250°C respectively; Nebulizer, 30 psi. Fragmentor, skimmer, and octopole radio frequency (Oct1 RF Vpp) voltages were set to 110 V, 65 V, and 750 V, respectively. The scan range was 70–1,200 m/z and spectra were collected at 4.4 spectra/s, which corresponded to 2,148 transients/spectrum. MS scan peaks of all the possible mass isotopologs were integrated using MassHunter Quantitative Analysis Software (version B.07.01, Build 7.1.524.0). Extracted peak matrices were processed using IsoCor (Millard et al., 2019) to correct for the contribution of naturally occurring isotopes. Relative isotopomer abundance for each metabolite and percentages of 13C enrichment were calculated according to Araújo et al. (2014). Metabolite quantitation was carried out based on calibration curves obtained with unlabeled authentic standards.

Relative quantitation of mitochondrial protein abundances by LC–MRM–MS

Multiple reaction monitoring (MRM) was carried out exactly as described previously (James et al., 2019), except trypsin was added to the protein samples in a mass ratio of 1:20. Peptide abundances from each sample were normalized against Voltage-dependent anion channel (VDAC) in which its abundance was identical between mitochondria from Col-0, dic2-1, and dic2-1/dic2 based on immunoblotting. Transitions used for multiple reaction monitoring of VDAC, DIC2, TCA cycle enzymes, pyruvate dehydrogenase, and NAD-ME are provided in Supplemental Data Set S2.

Statistical analysis

All statistical analyses were performed either with Excel (Student’s t test) or RStudio (analysis of variance (ANOVA) with Tukey Posthoc analysis, Shapiro–Wilk normality test, and Levene’s test) or RStudio (analysis of variance (ANOVA) with Tukey Posthoc analysis, Shapiro–Wilk normality test, and Levene’s test). Kinetic parameters Vₚₚ and Vₚₚₚ were determined from the Michaelis–Menten equation using the Enzyme Kinetic module in SigmaPlot (version 12.5). Statistical tests and replicate number are as indicated in figure legends; all statistical evidence is provided in Supplemental Data Set S3.
Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: DIC1 (At2g22500), DIC2 (At4g25470), DIC3 (At5g09470), VDAC1 (At3g01280), SAND (At2g28390), EF-1α (At5g03930), CSY2 (At3g58750), CSY3 (At2g42790), ACLA-1 (At1g0670), ACLA-2 (At1g0810), ACLA-3 (At1g09430), ACLB-1 (At3g06650), ACLB-2 (At5g49460), SEN1 (At4g35770), WRKY53 (At2g28390), and SAG101 (At5g14930). All data generated or analyzed during this study are included in this published article and its [supplementary information files](https://www.plantcell.org/content/33/19/3700.full).
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