MCU proteins dominate in vivo mitochondrial Ca\(^{2+}\) uptake in Arabidopsis roots

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Abstract

Ca\(^{2+}\) signaling is central to plant development and acclimation. While Ca\(^{2+}\)-responsive proteins have been investigated intensely in plants, only a few Ca\(^{2+}\)-permeable channels have been identified, and our understanding of how intracellular Ca\(^{2+}\) fluxes is facilitated remains limited. Arabidopsis thaliana homologs of the mammalian channel-forming mitochondrial calcium uniporter (MCU) protein showed Ca\(^{2+}\) transport activity in vitro. Yet, the evolutionary complexity of MCU proteins, as well as reports about alternative systems and unperturbed mitochondrial Ca\(^{2+}\) uptake in knockout lines of MCU genes, leave critical questions about the in vivo functions of the MCU protein family in plants unanswered. Here, we...
demonstrate that MCU proteins mediate mitochondrial Ca$^{2+}$ transport in planta and that this mechanism is the major route for fast Ca$^{2+}$ uptake. Guided by the subcellular localization, expression, and conservation of MCU proteins, we generated an mcu triple knockout line. Using Ca$^{2+}$ imaging in living root tips and the stimulation of Ca$^{2+}$ transients of different amplitudes, we demonstrated that mitochondrial Ca$^{2+}$ uptake became limiting in the triple mutant. The drastic cell physiological phenotype of impaired subcellular Ca$^{2+}$ transport coincided with deregulated jasmonic acid-related signaling and thigmomorphogenesis. Our findings establish MCUs as a major mitochondrial Ca$^{2+}$ entry route in planta and link mitochondrial Ca$^{2+}$ transport with phytohormone signaling.

Introduction

A characteristic of plant life is the ability to respond to environmental and developmental stimuli via flexible adjustments of metabolism, cell physiology, and the body plan. A signaling strategy that underpins a remarkable number of different responses is the modulation of intracellular free Ca$^{2+}$ concentration (White and Broadley, 2003; McAnish and Pittman, 2009; Kudla et al., 2018). Ca$^{2+}$ signaling mediates responses to abiotic stress factors, such as heat, cold, salt stress, low oxygen, biotic interactions such as pathogen attack and symbiosis, and developmental stimuli to drive processes such as pollen tube and root hair growth. The precise modulation of the timing and cellular location of Ca$^{2+}$ transients, as well as the physical association of different Ca$^{2+}$ responder proteins with specific interactors, offer a large combinatorial landscape that enables highly specific responses in gene expression, physiology, and metabolism (Carafoli and Krebs, 2016; Edel et al., 2017; Elie et al., 2020). While several of the many plant Ca$^{2+}$ sensors and responders of the calmodulin (CaM), CaM-like, calcineurin-B-like (CBL), CBL-interacting protein kinase (CIPK), Ca$^{2+}$-dependent protein kinase (CDPK/CPK), and Ca$^{2+}$/CaM-dependent protein kinase families have been studied down to the molecular details (recently reviewed by Kudla et al., 2018), only a few Ca$^{2+}$ channel proteins that mediate the Ca$^{2+}$ transients in the first place are currently known (Hedrich, 2012; Kong et al., 2020). Prominent examples include the cyclic nucleotide-gated channels (Charpentier et al., 2016; Leitão et al., 2019; Meena et al., 2019; Tian et al., 2019), glutamate receptor-like channels (Toyota et al., 2018; Wu’dick et al., 2018; Nguyen et al., 2018a), reduced hyperosmolarity-induced Ca$^{2+}$ increase channels (OSCa) (Hou et al., 2014; Yuan et al., 2014; Murthy et al., 2018; Thor et al., 2020), and Mid1-complementing activity, representing transmembrane proteins with potential for mechanosensing Ca$^{2+}$ channel activity (Nakagawa et al., 2007; Yamanaka et al., 2010; Yoshimura et al., 2021).

While studies of Ca$^{2+}$ signaling in plants have mainly focused on processes at the plasma membrane and in the cytosol and nucleus, it has been clear from early on that central processes of cellular Ca$^{2+}$ regulation occur at and in organelles, including the vacuole, endoplasmic reticulum, peroxisomes, chloroplasts, and mitochondria (reviewed in Stael et al., 2012; Resentini et al., 2021b), where Ca$^{2+}$-dependent proteins regulate key organelle functions (e.g. Weinl et al., 2008; Jin et al., 2009). Some of the earliest evidence for intracellular Ca$^{2+}$ transport in both plants and animals originates from Ca$^{2+}$ uptake experiments with isolated mitochondria (De Luca and Engstrom, 1961; Vasington and Murphy, 1962; Hodges and Hanson, 1965). Based on these studies, mitochondria had initially been postulated to act as major cellular Ca$^{2+}$ stores. Subsequent in vivo Ca$^{2+}$ measurements using aequorin in the mitochondria of Arabidopsis thaliana rosettes found that free Ca$^{2+}$ concentrations were only slightly higher in the matrix than in the cytosol and that mitochondrial Ca$^{2+}$ transients follow cytosolic Ca$^{2+}$ transients, albeit with different signatures (Logan and Knight, 2003). These observations suggested that the operation of a transport system at the mitochondrial membranes modulates uptake and generates specific mitochondrial Ca$^{2+}$ signatures (Logan and Knight, 2003; Loro et al., 2012).

Several Ca$^{2+}$ uptake pathways have been described for animal mitochondria (Sparagna et al., 1995; Buntinas et al., 2001; Bondarenko et al., 2013), but the Ca$^{2+}$ uniporter pathway was established as a major mode of Ca$^{2+}$ entry, and its pharmacological and electrophysiological properties were described in detail (Kirchok et al., 2004; Hajnoczky and Csordás, 2010). In contrast, the Ca$^{2+}$ fluxes measured in isolated plant mitochondria showed very different properties with respect to their pharmacology, the rates of uptake, and the requirement of phosphate, which is why the existence of a plant mitochondrial Ca$^{2+}$ uniporter had been debated (Dieter and Marme´, 1980; Zottini and Zannoni, 1993), albeit in the absence of in vivo data. When the physical identities of the first components of the mitochondrial Ca$^{2+}$ uniporter complex were elucidated in animals, that is, mitochondrial Ca$^{2+}$ uniporter (MCU) and mitochondrial Ca$^{2+}$ uptake (MICU) (Perocchi et al., 2010; Baughman et al., 2011; De Stefani et al., 2011), homologous genes were found also in plant genomes (Bick et al., 2012; Stael et al., 2012). Despite the conservation at the genetic level, noticeable differences became apparent, mirroring the functional differences observed earlier. For instance, multiple MCU copies exist in vascular plants, and a dominant-negative MCU paralog (MCUb; Raffaello et al., 2013) is lacking (Teardo et al., 2017). Similarly, the essential MCU regulator (EMRE), a component of the mitochondrial Ca$^{2+}$ uniporter complex in metazoans and some fungi required for its function (Pitts et al., 2020), is absent in plant genomes (Sancak et al., 2013; Wagner et al., 2016; Wang et al., 2019). Instead of three  

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encoding isoforms of the regulatory Ca\(^{2+}\)-binding EF-hand protein MCU in mammals, Arabidopsis only contains a single MCU locus (Ștael et al., 2012; Wagner et al., 2015; 2016). A functional investigation of Arabidopsis MICU showed that this protein has Ca\(^{2+}\)-binding activity and localizes to the mitochondrial inner membrane (Wagner et al., 2015). Arabidopsis micu knockout lines displayed deregulated mitochondrial Ca\(^{2+}\) signatures due to elevated concentrations of free matrix Ca\(^{2+}\), and enhanced uptake rates in planta, providing evidence for an inhibitory function of uniporter activity (Wagner et al., 2015).

Arabidopsis MCU1 and MCU2 localize to mitochondria (Tearo et al., 2017; Selles et al., 2018), while an additional localization to chloroplasts was reported for MCU6 in mesophyll cells (MCU6 was therefore referred to as chloroplast MCU [cMCU]) (Tearo et al., 2019). Arabidopsis MCU proteins were reported to show Ca\(^{2+}\) transport activity based on in vitro and heterologous analysis systems, that is, in planar lipid bilayers for MCU1 and MCU6 (Tearo et al., 2017, 2019), in human embryonic kidney 293 cells for MCU2 (Tsai et al., 2016), in yeast (Saccharomyces cerevisiae) for MCU1 and MCU2 (Selles et al., 2018), and in Escherichia coli for MCU6 (Tearo et al., 2019). These studies revealed that plant MCUs are sufficient for the formation of active channels, and there is no strict requirement for additional proteins, like for EMRE in metazoans. This is a feature that appears to be shared between plant MCUs and the MCUs of protozoa (Kovács-Bogdán et al., 2014; Chiurillo et al., 2017).

However, the critical question of whether MCUs are necessary for mitochondrial Ca\(^{2+}\) uptake in planta has remained unanswered, despite considerable efforts. As a consequence, the quantitative contribution of this putative Ca\(^{2+}\) uptake pathway relative to other potential routes of plant mitochondrial Ca\(^{2+}\) uptake (e.g. Carraretto et al., 2016a; reviewed in Wagner et al., 2016) also remains unclear. Furthermore, the physiological significance of MCU-mediated Ca\(^{2+}\) transport for other functions in plants has not been addressed in detail. Even though changes in the mitochondrial Ca\(^{2+}\) transient signature were observed in an Arabidopsis mcu1 knockout background, those effects were subtle, and any direct indicator of mitochondrial Ca\(^{2+}\) uptake rate remained unchanged (Tearo et al., 2017), suggesting either a lack of a prominent role of MCU1 in mitochondrial Ca\(^{2+}\) uptake in vivo or functional backup by the remaining mitochondrial MCU isoforms. For mcy2 knockout lines a fertility phenotype with reduced paternal transmission was observed and linked to defects in pollen tube germination and growth in vitro, which were, however, not evident in vivo (Selles et al., 2018). Whether the lack of MCU2 has an impact on mitochondrial Ca\(^{2+}\) transport in pollen or any other living tissue has remained unclear. The involvement of MCU6/cMCU in mediating in vivo Ca\(^{2+}\) flux into the chloroplast stroma has been reported, and phenotypes of Arabidopsis mcy6 lines were linked to deregulation in chloroplast Ca\(^{2+}\) dynamics (Tearo et al., 2019).

Here, we addressed the in vivo function of the plant MCU family. We used a reverse genetic approach to generate an Arabidopsis triple MCU knockout line of MCU1, MCU2, and MCU3 to overcome redundancy among MCU family members guided by Ca\(^{2+}\) imaging in root tips and the assessment of organismal phenotypes. Our findings indicate that MCU proteins mediate mitochondrial Ca\(^{2+}\) transport in planta and that this mechanism is the major route for rapid Ca\(^{2+}\) uptake.

**Results**

**The properties of the MCU family in Arabidopsis suggest functional redundancy**

We used Arabidopsis as a model to investigate the characteristics of the plant MCU family. MCU gene homologs are present across the plant kingdom and comprise small families in land plants (Tearo et al., 2017). A family of six Arabidopsis MCU homologs can be subdivided into two clades based on amino acid sequence identity: clade A (AtMCU1 [At1g09525], AtMCU2 [At1g37610]) and clade B (AtMCU3 [At2g23790], AtMCU4 [At4g36820], AtMCU5 [At5g42610], AtMCU6 [At5g66650]) (Figure 1, A and B; Supplemental File S1; Supplemental Data Set 1). All homologs share a predicted arrangement of an N-terminal signal peptide sequence and a loop containing a DVME amino acid motif, similar to the DIME motif found in the pore region of the mammalian MCU (Lee et al., 2015; Fan et al., 2020), flanked by two transmembrane helices (Supplemental Figure S1).

Based on sequence conservation and the observation that in vivo mitochondrial Ca\(^{2+}\) uptake was maintained in the absence of MCU1 (Tearo et al., 2017), we reasoned that MCU family members might be functionally redundant in mitochondria. To test this hypothesis and to generate a model system that overcomes this redundancy (Figure 1M), we first combined two Arabidopsis T-DNA lines by crossing, giving rise to the clade A null mutant mcu1 mcu2 (abbreviated mcu1 2). The absence of MCU1 and MCU2 expression, respectively, was previously established for both parental lines (Tearo et al., 2017; Selles et al., 2018) and was confirmed for the individual homozygous lines (Supplemental Figure S2, B–D) and for mcu1 2 (see paragraph below about Figure 3A). MCU1 and MCU2 were prioritized because they have diverged the least from the mammalian MCUs (Figure 1A), prior experimental evidence exists for their mitochondrial localization (Tearo et al., 2017; Selles et al., 2018), and their presence in the mitochondrial proteome has been experimentally confirmed (Wagner et al., 2015; Senkler et al., 2017; Fuchs et al., 2020). We further validated that both MCU1 and MCU2 are expressed in shoot and root tissues of 14-day-old wild-type seedlings (Figure 1C).

**Mitochondrial Ca\(^{2+}\) dynamics remain unperturbed in mcu1 2 roots at modest Ca\(^{2+}\) fluxes**

To investigate whether mitochondrial Ca\(^{2+}\) dynamics were altered by MCU ablation in planta, we introduced the
Figure 1 Mitochondrial matrix Ca\textsuperscript{2+} transients in response to stimulation with the auxin 1-naphthaleneacetic acid (NAA) in living root tips of mcu1, mcu1 2, and wild-type (Col-0) seedlings. A, Phylogenetic tree of A. thaliana and M. musculus proteins with homology to human MCU and MCU. Scale bar corresponds to a distance of 0.20 amino acid substitutions per site. Bootstrap values, as calculated using 1,000 replicates, are indicated at the nodes. Color code indicates clustering of corresponding groups. Amino acid sequences from GenBank. A. thaliana MCU1 (NP_001322910.1), A. thaliana MCU2 (NP_001077733.1), A. thaliana MCU3 (NP_179959.1), A. thaliana MCU4 (NP_195400.2), A. thaliana MCU5 (NP_199075.1), A. thaliana MCU6 (NP_201466.1), H. sapiens MCU (NP_612366.1), M. musculus MCU (NP_001028431.2), H. sapiens MCUb (NP_060388.2), and M. musculus MCUb (NP_080055.2). B, Amino acid sequence identity of MCU homologs in A. thaliana. Identity percentage, as
genetically encoded ratiometric Ca\(^{2+}\) sensor Yellow Cameleon 3.6 targeted to the mitochondrial matrix (4mt-YC3.6; Loro et al., 2012) into the mcu1 2 background (Supplemental Figure S3). YC3.6 was selected over Aequorin to allow imaging at high spatial resolution and over other genetically encoded fluorescent Ca\(^{2+}\) sensors based on its ratiometric nature and its pH stability; YC3.6 has proven to be reliable for direct comparisons of resting Ca\(^{2+}\) levels between different genetic backgrounds, as well as for monitoring Ca\(^{2+}\) dynamics in the plant mitochondrial matrix where pH can vary significantly (Schwarzländer et al., 2012; Wagner et al., 2015; Behera et al., 2018; Grenzi et al., 2021). Furthermore, we recently established the linear rate increase in YC3.6 Förster resonance energy transfer (FRET) ratio towards the start of a Ca\(^{2+}\) transient as a particularly direct measure of in vivo Ca\(^{2+}\) uptake activity (as opposed to other key characteristics of the transient, which may be affected more strongly by indirect/secondary factors, such as Ca\(^{2+}\) buffering or clearance) (Wagner et al., 2015). To assess mitochondrial Ca\(^{2+}\) dynamics, we focused on root tips as a model and hotspot of intracellular Ca\(^{2+}\) dynamics. We compared mcu1 2 to wild-type Col-0 seedlings, and we also included the mcu1 single mutant for which previous work could not resolve any changes in mitochondrial Ca\(^{2+}\) uptake rate in vivo (Teardo et al., 2017). We evoked Ca\(^{2+}\) transients by applying the synthetic auxin 1-naphthalene acetic acid (NAA, 0.01 mM) in a custom perfusion setup for in vivo fluorescence microscopy (Figure 1, D and E; Supplemental Movie S1), a system that we previously optimized to study subcellular Ca\(^{2+}\) transport in Arabidopsis mutants (Wagner et al., 2015; Behera et al., 2018). We quantitatively assessed the following key characteristics of the transient, which hold key information about the properties of mitochondrial Ca\(^{2+}\) management (Figure 1F): the steady-state FRET ratio (Figure 1C), which provides an integrated measure of Ca\(^{2+}\) uptake, export and buffering in the matrix at baseline; the linear rate of the FRET increase in the starting phase of the Ca\(^{2+}\) transient (Figure 1, H and I), which represents the most direct measure of in vivo Ca\(^{2+}\) uptake rate; the maximal FRET ratio (Figure 1J) as a measure of the maximal free Ca\(^{2+}\) accumulation reached at the peak of the transient (i.e. when uptake and clearance reach equal rates); the time that passes from the start of the transient to the maximum (\(t_{\text{max}}\) Figure 1K); and the time that passes after the peak to return to 50% of the amplitude (\(t_{1/2 \text{ down}}\) Figure 1L) as an indicator of Ca\(^{2+}\) clearance efficiency, which may be modulated in response to primary changes in matrix Ca\(^{2+}\) accumulation. Before the NAA treatment (time 0–145 s), baseline matrix free Ca\(^{2+}\) levels were indistinguishable between lines (Figure 1, D, E, and G). Also, after the start of NAA perfusion, the characteristics of the mitochondrial Ca\(^{2+}\) transients did not differ between the three lines in any of their key properties (Figure 1, H–L). The absence of any differences between the lines indicates that, despite the lack of MCU1 and MCU2, NAA-induced mitochondrial Ca\(^{2+}\) uptake remained unaffected, and the mitochondrial uptake capacity was not limiting in any of the lines investigated.

Absence of MCU1 and MCU2 limits mitochondrial Ca\(^{2+}\) uptake at high Ca\(^{2+}\) fluxes

We next hypothesized that the mitochondrial Ca\(^{2+}\) uptake machinery becomes limiting only at sufficiently high in vivo Ca\(^{2+}\) flux (Figure 1M). To reveal putative differences in Ca\(^{2+}\) transport capacity, we aimed to drive Ca\(^{2+}\) transport activity, that is, the actual Ca\(^{2+}\) flux across the mitochondrial inner membrane, closer to the capacity limit of the uptake machinery. For this purpose, we used extracellular ATP (eATP) treatments at 0.01, 0.1, and 2.0 mM as a model stimulus to trigger mitochondrial Ca\(^{2+}\) transients in a dose-dependent manner ranging from low to high Ca\(^{2+}\) fluxes (Loro et al., 2012; Wagner et al., 2015; Matthus et al., 2019). Exposure to eATP triggered matrix Ca\(^{2+}\) transients in the root tips of all lines, and the elicited transients increased proportionally with increasing eATP concentration (Figure 2, A, H, and O). To achieve a systematic and quantitative

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**Figure 1 (Continued)** calculated using the Basic Local Alignment Search Tool (BLAST), is displayed by numerical values and color code (green, high identity; orange, low identity). C, Reverse transcription–quantitative PCR (RT–qPCR) analysis of MCU1 and MCU2 mRNA abundance in shoots and roots of 14-day-old Col-0 seedlings. Gene expression was normalized to SAND family protein expression for each sample. Error bars = s.d., n = 5–6. D, FRET ratios over time in root tips of mcu1 1, mcu1 2, and Col-0 seedlings expressing the mitochondrial Ca\(^{2+}\) biosensor YC3.6, acquired under continuous perfusion and treated with 0.01 mM NAA for 3 min, as indicated by the black box on the x-axis. Whole frames as illustrated in part E were used for the quantification. D’, same as (D) but y-axis scale and range adjusted. E, False-color images illustrating FRET ratios in root tips of seedlings expressing the mitochondrial YC3.6 in the indicated genetic backgrounds imaged under continuous perfusion shown at steady-state at 100 s (upper images) and during the Ca\(^{2+}\) transient induced by treatment with 0.01 mM NAA for 3 min at 430 s (lower images). A representative set of replicates of a complete time-series each is shown in Supplemental Movie 1. To compose the panel, images were cropped and image margins, when depicted, are colored in black. Low FRET ratios indicating low Ca\(^{2+}\) are shown in blue, and high FRET ratios indicating high Ca\(^{2+}\) are shown in red. Scale bar = 50 µm. F, Schematic representation of the extracted kinetic parameters of the sensor response depicted as letters in parenthesis and shown in parts (G)–(L). G, Steady-state FRET ratios before stimulus application in Col-0, mcu1 1, and mcu1 2. H, FRET ratio increase following 0.01 mM NAA application; linear region was selected by maximal goodness-of-fit measure of linear regression (\(R^2\)) for Col-0. I. Linear rate representing the slope of regression of part (H). J, Maximal peak FRET ratio after 0.01 mM NAA application. K, Time required to reach maximal FRET ratio after stimulus application. L, Time required to pass maximal ratio amplitude during recovery after the stimulus. n ≥ 9; error bars = s.d. No significant differences between Col-0 and the mutant lines were observed (One-way ANOVA with Dunnett’s multiple comparisons test). M, Experimental rationale based on the hypothesis that MCU capacity for Ca\(^{2+}\) transport exceeds in vivo Ca\(^{2+}\) uptake activity from the cytosol/intermembrane space into the mitochondrial matrix. Hence, revealing in vivo MCU function requires the Ca\(^{2+}\) transport capacity to be limited below the required activity by knocking out a critical number of MCU genes.
Figure 2 Genetic ablation of both MCU1 and MCU2 alters mitochondrial matrix Ca\textsuperscript{2+} transients induced by stimulation with eATP. A, FRET ratios over time in root tips of mcu1, mcu1 2 and Col-0 seedlings expressing the mitochondrial YC3.6, acquired under continuous perfusion and treated with 0.01 mM eATP for 3 min. Whole frames as illustrated in (B) were used for the quantification. B, Ratiometric images illustrate FRET ratios in root tips of seedlings expressing 4mt-YC3.6 in the indicated genetic backgrounds acquired under continuous perfusion and shown at steady-state at 100 s (upper images) and during the Ca\textsuperscript{2+} transient induced by treatment with 0.01 mM eATP for 3 min at 320 s (lower images). A representative set of replicates of a complete time-series each is shown in Supplemental Movie 2. To compose the panel, images were cropped and image (continued)
comparison, we analyzed the individual characteristics of the eATP-induced transients as for NAA (Figures 1, D–L and 2). No difference between the lines was observed at 0.01 mM eATP (Figure 2, A–G; Supplemental Movie S2). At 0.1 mM eATP, the mcu1 2 double null mutant showed a tendency toward lower rates of increases in matrix free Ca\(^{2+}\) levels, but this effect was not statistically significant (Figure 2, H–N; Supplemental Movie S3). At 2 mM eATP, however, the matrix Ca\(^{2+}\) transient was markedly different in the mcu1 2 line (Figure 2, O–U; Supplemental Movie S4), which showed decreased rates of Ca\(^{2+}\) level elevation (Figure 2, Q and R) and lower maximal Ca\(^{2+}\) accumulation (Figure 2S). The time to reach the peak of the transient and the recovery remained unchanged (Figure 2, T and U).

These data show that mitochondrial Ca\(^{2+}\) uptake capacity became limiting in the mcu1 2 line under high rates of cytosolic Ca\(^{2+}\) flux. Since such a limitation was not observed in the mcu1 line, the limitation was due to either the combination of the two alleles or to mcu2 alone. We decided against trying to pinpoint the specific quantitative in vivo contribution to the individual MCU loci since such an approach has proven futile in systems where total in vivo capacity exceeds the required in vivo activity (as is the case for most enzymes of central plant metabolism). Rather, we focused on further constraining total MCU capacity in order to investigate the significance of the MCU family. Since the mitochondrial Ca\(^{2+}\) uptake rate was reduced by approximately one-third in mcu1 2 root tips (Figure 2R), there was considerable remaining capacity for mitochondrial Ca\(^{2+}\) uptake, which was mediated by either the remaining MCU homologs or by alternative transport systems.

### The mcu1 2 3 triple mutant shows a dramatic reduction in total MCU expression

To address the hypothesis that remaining genes from the MCU family provide functional backup in the mcu1 2 background, we carefully selected an additional MCU locus to generate a triple knockout line. Only the clade A members MCU1 and MCU2 could be detected in mitochondrial proteomes (Wagner et al., 2015; Senkler et al., 2017; Fuchs et al., 2020). Yet, direct evidence for MCU3, MCU4, and MCU6 expression at the protein level comes from a recent mass spectrometry-based proteomic analysis of whole Arabidopsis tissues, in which only MCU5 was not found (Mergner et al., 2020). For a quantitative account of the relative expression of the MCU homologs, we measured their transcript abundance in roots and shoots by reverse transcription–quantitative PCR (RT–qPCR). In roots, MCU2, MCU3, and MCU4 were the most highly expressed MCU transcripts, with similar levels of abundance (Figure 3A), while MCU3 was the most abundant MCU transcript in shoots (Supplemental Figure S4A). MCU5 was below the detection limit in both roots and shoots. These data are in line with public mRNA-seq data (GENEVESTIGATOR, Hruz et al., 2008; Klepikova et al., 2016) (Supplemental File S2).

Based on the high MCU3 transcript abundance in both roots and shoots, we selected MCU3 to further limit total in vivo MCU capacity. Notably, MCU3 shares the highest amino acid sequence identity with MCU1 and MCU2 relative to the other members of MCU clade B (Figure 1B) and exclusively localizes to the mitochondria, which we validated by subcellular localization analysis of an MCU3-GFP construct (Supplemental Figure S5). Therefore, we crossed an mcu3 T-DNA insertion line into the mcu1 2 background to generate mcu1 mcu2 mcu 3 (abbreviated mcu 1 2 3 hereafter; Supplemental Figure S6). RT–qPCR analysis of root and shoot tissue confirmed the absence of the MCU1, MCU2, and MCU3 transcripts in the mcu1 2 3 background (Figure 3A, Supplemental Figure S4A). The expression levels of MCU5 and MCU6 were similarly low in wild-type Col-0, mcu1, mcu1 2, and mcu 1 2 3. Surprisingly, the abundance of the MCU4 transcript decreased with increasing MCU ablation. Consequently, combined MCU transcript expression was reduced down to 22% in roots and 15% in shoots in the mcu 1 2 3 background (Figure 3A; Supplemental Figure S4A).

### Tissue gradients of mitochondrial Ca\(^{2+}\) at baseline are perturbed in mcu 1 2 3 roots

To directly assess whether reduced MCU expression gives rise to changes in mitochondrial Ca\(^{2+}\) dynamics in mcu1 2 3 seedlings, we introduced the mitochondrial Ca\(^{2+}\) biosensor 4mt-YC3.6 (Loro et al., 2012) into the triple mutant background. Exclusive mitochondrial localization of the sensor was confirmed by co-localization with MitoTracker (Supplemental Figure S7). Strikingly, we observed an apparent difference in the FRET ratio of the biosensor, indicating lower resting concentrations of matrix free Ca\(^{2+}\) in the root tips of mcu1 2 3 seedlings (Figure 3B). While root tips of Col-0 showed a characteristic tissue gradient of matrix Ca\(^{2+}\) levels that increased from the root cap toward the elongation zone (see also Figures 1, E and 2, B, I, and P) (Wagner et al., 2015), this gradient was noticeably flattened in mcu1 2 3, but not in mcu1 or mcu1 2 (Figure 3B).

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**Figure 2 (Continued)**

Margins, when depicted, are colored in black. C, FRET ratio increase following 0.01 mM eATP application; linear region was selected by maximal goodness-of-fit measure of linear regression (R\(^2\)) for Col-0 (R\(^2\) > 0.998 for mcu1 and R\(^2\) > 0.992 for mcu1 2) (D) Linear rate (ratio change per second) representing the slope of regression of (C). E, Maximal peak FRET ratio after 0.01 mM eATP application. F, Time required to reach maximal FRET ratio after stimulus application. G, Time required to pass half-maximal ratio during recovery after the stimulus. H–N, Analogous analysis for 0.1 mM eATP as in (A–G). Note that in (I), white image sectors within the root tissue region represent saturated pixels, which were removed from the ratiometric analysis. A representative set of replicates of a complete time-series each is shown in Supplemental Movie 3. O–U, Analogous analysis for 2.0 mM eATP as in (A–G). A representative set of replicates of a complete time-series each is shown in Supplemental Movie 4. Scale bars = 50 μm. n ≥ 6; error bars = SD. *P < 0.05, **P < 0.01 (One-way ANOVA with Dunnett’s multiple comparisons test).
explore the effect of these tissue-specific changes, we analyzed the spatial distribution of FRET ratios by subdividing the sides of the root tip, where mitochondrial Ca\(^{2+}\) dynamics are particularly active (see Figures 1, E and 2, B, I, and P), into nine equally sized regions of interest (ROIs; Figure 3C). We excluded the root cap from the analysis, since it showed consistently low levels of mitochondrial free Ca\(^{2+}\) in all genetic backgrounds, and it does not typically show any pronounced response to stimuli that trigger Ca\(^{2+}\) transients in the other tissues. In each of the nine ROIs, the matrix-free Ca\(^{2+}\) concentration was lower in mcu1 2 3 roots compared to the wild-type control (Figure 3C), demonstrating that MCU activity is required to set basal matrix free Ca\(^{2+}\) concentrations in Arabidopsis. Furthermore, matrix Ca\(^{2+}\)
levels progressively increased from ROI1 to ROI9 in Col-0 roots (Figure 3C), adding up to a marked difference, as quantified for ROI2 and ROI9 (Figure 3C). This trend was maintained in mcu1 and mcu1 2 but was completely abolished in mcu1 2 3 (Figure 3, C and C). To control for general alterations in root tip architecture, we assessed the cellular organization in the root tips of mcu1 2 3 and Col-0 (Supplemental Figure S4B). The overall root tip architecture was unchanged in mcu1 2 3, as indicated by the size and number of cell layers of the meristematic zone (Supplemental Figure S4, B–D) and the finding that MCU1, 2 and 3 were expressed in the cell types that are present in the ROIs (Supplemental Figure S4E), validating the ROI comparison between the different genetic backgrounds.

Mitochondrial Ca$^{2+}$ uptake capacity is strongly impaired in mcu1 2 3 root tissues

We next hypothesized that the shift in basal matrix free Ca$^{2+}$ was due to the severe impairment of Ca$^{2+}$ uptake in mcu1 2 3. To obtain direct in vivo information on the uptake, we monitored root tip Ca$^{2+}$ transients triggered by 2.0 mM eATP in ROI2 and ROI9 (Figure 4; Supplemental Movie S5). Different signatures of the transients were observed between the different ROIs, with more pronounced transients of higher uptake rates and amplitude in ROI2 than in ROI9 (Figure 4, A, B, and H). Yet, in both ROIs, as well as all ROIs in between, the impairment of the transient (i.e. uptake rate and maximal FRET ratio) in mcu1 2 was exacerbated in mcu1 2 3 (Figure 4, C–E, I–K). In ROI9, the uptake rate was close to 0 in mcu1 2 3, and the transient was nearly completely abolished (compromising the accurate measurement of transient characteristics; Figure 4, I–M). In ROI2, a transient still occurred in mcu1 2 3, but the uptake rate was diminished to <25% that of Col-0 (Figure 4, B–D). Ca$^{2+}$ clearance efficiency, as indicated by $t_{1/2}$ downwa was perturbed in ROI2 but not in ROI9 (Figure 4, G and M). Systematic analyses using a weaker dose of the eATP stimulus that triggered a weaker matrix Ca$^{2+}$ transient (0.1 mM eATP) yielded similar results (Supplemental Figure S8). Only at the weakest transient (0.01 mM eATP) did the differences between the genotypes start to become obscured (Supplemental Figure S9).

The data not only demonstrate that MCU proteins mediate mitochondrial Ca$^{2+}$ uptake in vivo but also that MCUs underpin the dominant mechanism for rapid matrix uptake in Arabidopsis root tips. The residual uptake activity in the mcu1 2 3 line might have been due to the remaining MCUs or to alternative mechanisms, which will need to be resolved in the future. In either case, those activities clearly make only minor contributions to mitochondrial Ca$^{2+}$ uptake capacity in the root tip and cannot compensate for the loss of MCU1, MCU2, and MCU3 when cytosolic Ca$^{2+}$ level elevations are rapid and pronounced. The altered mitochondrial Ca$^{2+}$ dynamics of the mcu1 2 3 line were independent of the model stimulus that evoked the primary cytosolic Ca$^{2+}$ elevation, as validated by auxin application (0.01 mM NAA; Supplemental Figure S10, A–F) to trigger a mild (non-limiting) Ca$^{2+}$ response and an acute osmotic challenge via sequential administration and washout of 200 mM sorbitol to trigger a pronounced Ca$^{2+}$ elevation at washout (Supplemental Figure S10, G–L; as optimized previously for guard cells and roots; Loro et al., 2012; Corso et al., 2018).

We tried to construct independent multi-knockout lines by clustered regularly interspaced short palindromic repeats and CRISPR-associated protein 9 (CRISPR–Cas9)-mediated genome editing but were not successful. Independent lines are typically required to test for the potential involvement of second site mutations and to validate the link between genetic locus and phenotype. In the specific system combining loci of one gene family, the link between the MCU loci and the mitochondrial Ca$^{2+}$ transport phenotype is demonstrated by the exacerbation of the phenotype from mcu1 2 to mcu1 2 3. It is highly unlikely that such a specific phenotype arises by chance, which rules out potential contributions by secondary site mutations with similarly high confidence as genetically independent lines. To validate this experimentally, we generated lines expressing MCU2 cDNA (35S:MCU2) in either the mcu1 2 3 background or the Col-0 wild-type background (Supplemental Figure S11, A and B). The MCU2 transcript was detected at high abundance (Supplemental Figure S11C), allowing for two independent lines each to be used for in vivo Ca$^{2+}$ analysis using 2 mM ATP as the stimulus (Supplemental Figure S12). Even though the lines analyzed were in the T2 generation, that is, segregating for the 35S:MCU2 insertion locus, the mitochondrial Ca$^{2+}$ dynamics in mcu1 2 3 35S:MCU2 roots were rescued, as evident from the average uptake rates (Supplemental Figure S12), demonstrating that the impairment of MCU function was indeed causative for the observed phenotype in mitochondrial Ca$^{2+}$ uptake. Interestingly, the roots of Col-0 35S:MCU2 seedlings showed even higher uptake rates than the wild-type, as well as increased matrix resting levels, confirming the correlation between MCU abundance due to elevated MCU expression and Ca$^{2+}$ uptake capacity. The finding that introducing MCU2 alone led to a functional rescue adds evidence for functional redundancy among MCU1, 2 and 3. Functional redundancy among the three MCU genes is further supported by the observation that mitochondrial Ca$^{2+}$ uptake in the mcu3 single mutant was similar to that of the wild-type (Supplemental Figure S12). This control shows that the combined absence of three MCU proteins, rather than MCU3 alone, is responsible for the major impairment of Ca$^{2+}$ uptake capacity in mcu1 2 3 (as compared to mcu1 2, where the effect is much milder; Figure 4).

Cytosolic Ca$^{2+}$ dynamics remain unperturbed in mcu1 2 3 roots

We next hypothesized that diminished mitochondrial Ca$^{2+}$ uptake would modify the primary Ca$^{2+}$ transient in the cytosol. Cytosolic Ca$^{2+}$ signatures are shaped by mitochondrial Ca$^{2+}$ transport, which acts as a capacitor, and these
signatures were shown to be modified by the impairment of the mitochondrial Ca\(^{2+}\) uniporter complex in mammalian cell systems (Drago et al., 2012; Logan et al., 2014; Rasmussen et al., 2015).

We expressed the NES-YC3.6 Ca\(^{2+}\) sensor in the cytosol (Krebs et al., 2012) in the mcu1 2 3 background and monitored the resting state of cytosolic free Ca\(^{2+}\), as well as Ca\(^{2+}\) transients elicited by 2.0 mM eATP (Figure 5A; Supplemental Movie S6). mcu1 2 3 and Col-0 showed identical resting states of cytosolic free Ca\(^{2+}\) (Figure 5B), and no change was observed in any of the characteristics of the Ca\(^{2+}\) transient in ROI2 and ROI9 either (Figure 5, C–N). We also systematically analyzed ROI1 and ROI3 to ROI8 and did not detect any difference in the cytosolic Ca\(^{2+}\) transient (Supplemental Figure S13). These data strongly suggest that even severe impairment of Ca\(^{2+}\) uptake into the mitochondria (Figure 4, H–J) does not alter the Ca\(^{2+}\) transients in the cytosol of root tips (Figure 5, I–K). The contribution made by the mitochondria may be sufficiently minor to be buffered or compensated for by other mechanisms. As a result, the impaired Ca\(^{2+}\) dynamics phenotype is highly specific to the mitochondrial matrix.

Impaired mitochondrial Ca\(^{2+}\) uptake in mcu1 2 3 is not associated with changes in gross plant development under a broad spectrum of conditions. Despite the dramatic cell physiological phenotype of diminished matrix Ca\(^{2+}\) uptake in the mcu1 2 3 line, the plants appeared to grow and develop normally. The systematic assessment of plant growth in soil, including quantifying rosette size (Supplemental Figure S14A), photosynthetic efficiency (Supplemental Figure S14B), and silique length (Deng et al., 2016) (Supplemental Figure S14C) did not reveal any apparent differences. The same was true for seedlings grown on agar plates, as assessed for leaf area and root length (Supplemental Figure S15). Also, the external
challenge of seedlings with different cues to induce osmotic, ionic, hormonal, and redox stress did not lead to differential growth behavior (Supplemental Figure S16). Since mitochondria have been shown to be critical for optimal tip growth, we specifically assessed tip-growing root hairs and pollen tubes for their developmental and growth properties (Supplemental Figure S17) (Schiefelbein et al., 1992; Monshausen et al., 2008; Zhang and Turner, 2008; Schoenaers et al., 2017). However, mcu1 2 3 behaved like Col-0 for all of the tested parameters. The impairment of mitochondrial electron transport and the induction of retrograde signaling by the Complex III inhibitor antimycin A (Wagner et al., 2019; Fuchs et al., 2022) did not result in any growth responses in seedlings (Supplemental Figure S18). In addition, no differences were detected in recovery after submergence (in darkness or continued light-dark cycles) and hypoxia stress, which we employed due to their impact on mitochondrial respiratory function and cytosolic Ca^{2+} levels (Wagner et al., 2019; Igamberdiev and Hill 2018) (Supplemental Figure S19). The characteristic gradual Ca^{2+} accumulation that occurs under hypoxia in the cytosol and the mitochondria of seedlings over several hours was unchanged (Supplemental Figure S19, D and E), indicating sufficient residual capacity for the low rates of mitochondrial Ca^{2+} transport involved. Whole plant darkening led to a small but significant difference in chlorophyll content between rosettes of Col-0 and the mcu1 2 3 line after 11 days of treatment (Supplemental Figure S20).

mcu1 2 3 seedlings show deregulated jasmonic acid homeostasis and impaired touch signaling
To validate the absence of gross phenotypic consequences at a global gene expression scale or to pinpoint the impact of disrupted MCU capacity and matrix free Ca^{2+} dynamics, we performed a comparative transcriptome analysis of roots from hydroponically grown mcu1 2 3 and Col-0 seedlings without additional stimulation. Five replicates of pooled seedling roots were analyzed each. The vast majority of

Figure 5 Cytosolic Ca^{2+} transients in root tips induced by treatment with eATP are unchanged in mcu1 2 3. A, Representative FRET images of Col-0 and mcu1 2 3 root tips expressing the cytosolic Ca^{2+} sensor NES-YC3.6 imaged under continuous perfusion and shown at steady-state before stimulus application at 100 s (upper images) and at 180 s (lower images) during the Ca^{2+} transient induced by treatment with 2.0 mM eATP for 3 min. A representative set of replicates of a complete time-series each is shown in Supplemental Movie 6. Scale bar = 50 μm. B, Steady-state FRET ratios in the roots before the application of 2.0 mM eATP. C, FRET ratio dynamics of ROI2 of seedlings expressing NES-YC3.6 acquired under continuous perfusion and treated with 2.0 mM eATP for 3 min. D, FRET ratio increase following 2.0 mM eATP application; linear region was selected by maximal goodness-of-fit measure of linear regression (R^2) for Col-0. E, Linear rate representing the slope of regression of (D). F, Peak FRET ratio after 2.0 mM eATP application. G, Time required to reach maximal FRET ratio after stimulus application. H, Time required to pass half-maximal ratio amplitude during recovery. I–N, Analogous analysis for ROI9 as in (C)–(H). n = 8; error bars = SD. No significant differences between Col-0 and mcu1 2 3 were observed (Student’s t test).
transcripts remained unchanged between the genotypes, confirming the absence of major pleiotropic rearrangements (Figure 6A). However, a specific set of 172 genes showed differential abundance (P-adjusted < 0.05; −1.5 ≤ fold change ≥ 1.5), the majority of which (159) were repressed in mcu1 2 3 compared to Col-0 (Supplemental Data Sets 2 and 3 for the complete lists of transcripts). The ontologies of the differentially expressed genes pointed toward functions in the response to stress and to external stimuli (Figure 6B). At the center of the different categories was a remarkable enrichment of transcripts involved in jasmonic acid (JA) metabolism (Zhang and Turner, 2008).

To explore the connection between MCU function and the transcriptional regulation of JA metabolism, we compared the 172 differentially abundant transcripts in mcu1 2 3 with the recently reported core set of 82 genes regulated by the JA-activated transcription factor MYC2 (i.e. the “MYC2 regulon” Van Moerkercke et al., 2019). Fourteen genes were shared in both datasets, representing a strong statistical enrichment, and all 14 transcripts were repressed in mcu1 2 3 (Supplemental Figure S21A; Supplemental Data Set 4). To test the hypothesis of altered JA homeostasis, we selected a set of established JA-related transcripts involved in JA biosynthesis (LOX, OPR3, and AOS) (Mueller, 1997; Wasternack, 2007; Schaller and Stintzi, 2009; Beaugelin et al., 2019), JA catabolism (JAO2 and JAO4; Caarls et al., 2017; Smirnova et al., 2017), and JA-mediated signaling pathways (MYC2, JAZs, NPR3, WRKY53, and GRX480) (Huang et al., 2017; Ruan et al., 2019; Van Moerkercke et al., 2019) and assessed their expression in the mcu1 2 3 root transcriptome, thus also including those transcripts that did not pass the criteria of P adjusted < 0.05 and −1.5 ≤ fold change ≥ 1.5 of the global analysis. All of these JA-related transcripts were repressed in mcu1 2 3 (Figure 6C). We selected four of the JA-related transcripts (OPR3, MYC2, JAZ10, and JAO4), as well as two stress marker transcripts (CPK28 and WRKY40) that were significantly repressed in the mcu1 2 3 transcriptome dataset for quantification by RT–qPCR in seedlings that were grown independently of the material used for the transcriptome analysis. All six transcripts were present at lower levels in mcu1 2 3 versus Col-0 (Figure 6D; Supplemental Figure S21B, B–D). An assessment of transcripts encoding Group VII ERF transcription factors, ethylene signaling components, and targets of mitochondrial retrograde regulation (De Clercq et al., 2013) did not reveal any concerted response (Supplemental Figure S21E).

To investigate whether the transcriptional fingerprint reflected changes in the contents of JA signaling intermediates, we quantified 12-oxo-phytodienoic acid (OPDA), dinor-oxo-phytodienoic acid (dn-OPDA), JA, and jasmonoyl-L-isoleucine (JA-Ile) in the roots of hydroponically grown mcu1 2 3 and Col-0 seedlings. JA and JA-Ile levels were doubled in the roots of mcu1 2 3 compared to Col-0 roots (Figure 6E), which was unexpected considering the positive feedback control of JA biosynthesis. OPDA and dn-OPDA were also present at higher levels in the mutant, but that change was variable between independent experimental repetitions (Figure 6E; Supplemental Figure S21D). To determine whether this inconsistent pattern was due to regulation at the protein level, we analyzed ALLENE OXIDASE CYCLASE (AOC) protein levels by immunoblotting (Stenzel et al., 2003). These levels were largely unchanged between the mcu backgrounds and the Col-0 control (Supplemental Figure S21C). Taken together, the transcriptome analysis and the analytical phythohormone measurements uncovered deregulated JA metabolism and signaling as well as stress responses in roots of the mcu1 2 3 mutant, pointing to an unexpected link between mitochondrial Ca2+ dynamics and JA homeostasis.

Recent observations have suggested that the integrated effects of mitochondrial regulation, Ca2+, and JA are involved in touch responses and thigmomorphogenesis (Van Aken et al., 2016; Van Moerkercke et al., 2019; Xu et al., 2019; Darwish et al., 2022), prompting us to assess the responses of mcu1 2 3 and Col-0 plants to a reoccurring touch stimulus (Darwish et al., 2022) (Figure 6, F–K). The resulting changes in rosette area, bolting time and inflorescence height consistently suggested an impaired thigmomorphogenesis response in the mcu1 2 3 mutants. While the touch stimulation of the shoot has no direct equivalent in the root, we devised a method for analyzing the mechanoresponse of root growth using an agar penetration assay (Mousavi et al., 2021). No differences in root growth were observed between mcu1 2 3 and the Col-0 control (Supplemental Figure S22). The discovery of a link between mitochondrial Ca2+ uptake, JA signaling and touch responses, as well as the potential deregulation of JA feedback control, opens the door for future work to resolve the connection in mechanistic detail and to unravel the impact of mitochondrial Ca2+ uptake capacity on processes linked to JA homeostasis and touch signaling.

Discussion

The major route of plant mitochondrial Ca2+ uptake is via MCU channels

Reverse genetics and in vivo Ca2+ imaging demonstrated that MCU proteins mediate Ca2+ uptake into the mitochondrial matrix in living Arabidopsis tissues. Exploiting the combinational ablation of three members of the MCU family and challenging these mutants with increasing intracellular Ca2+ fluxes, as elicited by NAA and eATP (Figures 1–4; Supplemental Figures S8 and S9), pinpointed the redundant functionality of MCU family proteins, which could be overcome by combining mutant lines. The additive Ca2+ uptake phenotype from mcu1 2 to mcu1 2 3 provides a clear demonstration that functional redundancy was overcome and that MCU capacity decreased beyond the normally required activity in vivo. These findings resolve the long-standing question of the function of a Ca2+ uniporter in plant mitochondria and unify the previous observations of (1) mitochondria-specific Ca2+ signatures in planta (Logan and Knight, 2003; Loro et al., 2012); (2) the activation of
Figure 6 Deregulated jasmonic acid homeostasis and impaired thigmomorphogenesis response in mcu1 2 3 seedlings. A, Volcano plot of the seedling root transcriptome of mcu1 2 3 compared to Col-0. Log2 fold change (FC, x-axis) is plotted against the –log10 P-value (y-axis). Downregulated transcripts (FC ≤ −1.5 and P-adjusted < 0.05): blue; upregulated transcripts (FC ≥ 1.5 and P-adjusted < 0.05): red; not differentially abundant transcripts: gray. B, Gene ontology (GO) enrichment analysis for transcripts that were down regulated in mcu1 2 3. The 10 categories with the highest enrichment are sorted by decreasing false discovery rate. Numerator: number of differentially abundant transcripts in the transcriptome dataset. Denominator: number of total genes curated for the GO term. C, Heat map of the relative abundance of JA-related transcripts from the transcriptome dataset shown in panel (A). FC values of mcu1 2 3 normalized to Col-0 (set to 1). The color gradient from white to black represents values from 1.0 (expression like Col-0) to 0.30 (expression lower than Col-0). D, Expression of selected JA-related transcripts, and UBQ10 and SAND FP for reference, as measured by RT–qPCR in roots of Col-0 and mcu1 2 3 seedlings. Relative expression was calculated as the ratio between the abundance of the respective transcript and of the UBIQUITIN-CONJUGATING ENZYME 21 (UBC21) reference transcript in the same sample. Relative expression is normalized to Col-0 (set to 1, black dotted line). n = 5–6; error bars = SD. * P < 0.05; ** P < 0.001 (Student’s t test). E,
mitochondrial Ca$^{2+}$ uptake when an Arabidopsis homologue of the metazoan MCU regulator MICU is absent (Wagner et al., 2015); and (3) the ability of MCU1 and MCU2 to act as Ca$^{2+}$ channels in vitro and in recombinant systems (Tsai et al., 2016; Teardo et al., 2017; Selles et al., 2018). The concomitant ablation of MCU isoforms 1, 2, and 3 (Figure 3A; Supplemental Figure S4A) diminished the in vivo Ca$^{2+}$ uptake down to residual rates (Figure 4; Supplemental Figures S8 and S9) and clearly demonstrated the dominant role of MCU proteins in rapid mitochondrial Ca$^{2+}$ uptake. Conversely, the 3SS:MCU2 lines showed the opposite effect, that is, increased matrix free Ca$^{2+}$ at baseline and increased Ca$^{2+}$ uptake rates (Supplemental Figure S12), suggesting that (1) MCU expression and mitochondrial Ca$^{2+}$ uptake capacity are directly linked and (2) even MCU abundance in the wild-type can be limiting for transport capacity. While the mcu1 2 3 line established here may be considered a strong knockdown line of total MCU capacity specifically of the mitochondria, a full knockout line may be generated in the future to also abolish any remaining MCU capacity. MCU5 did not appear to be expressed under any condition studied so far, but MCU4 and/or MCU6 may indeed provide the remaining Ca$^{2+}$ transport capacity observed here. However, including MCU6 in a higher-order knockout line would be complicated by the recent observation that MCU6 also localizes to chloroplasts (Teardo et al., 2019) (even though MCU6 expression in the plastid envelope exclusively in green tissue is surprising and deserves further investigation of the underlying regulation of protein targeting). A higher-order mutant line that includes MCU6 knockout could give rise to organismal effects that may arise from both the mitochondria and chloroplasts, complicating the mechanistic dissection of phenotypes. Hence, in addition to a sextuple mcu1 2 3 4 5 6 line, the sextuple line complemented with plastid-specific MCU6 appears to be most suitable to remove any residual mitochondrial MCU capacity while ruling out primary effects beyond the mitochondria.

Reverse genetics of proteins with redundant functions cannot reveal the relative quantitative contributions of the individual proteins in the wild-type in a straightforward manner. This is particularly true for a system in which the overall in vivo capacity exceeds the required in vivo activity. The clade A proteins MCU1 and MCU2, but no clade B proteins, were detected by mitochondrial proteomics (Wagner et al., 2015; Senkler et al., 2017; Fuchs et al., 2020) and also showed a higher peptide coverage than clade B proteins in tissue-specific proteomics (Mergner et al., 2020), suggesting that MCU1 and MCU2 are the most abundant MCU proteins in Arabidopsis mitochondria. Yet, the mcu1 2 3 triple mutant showed a stronger impairment in mitochondrial Ca$^{2+}$ uptake than the mcu1 2 double mutant (Figures 3 and 4). This observation suggests that the absence of MCU1 and MCU2 (or MCU2 on its own), abolishes any excess uptake capacity, rendering the remaining uptake system limiting, which gives rise to a strong additional impairment of uptake rate in response to MCU3 ablation. Loss of MCU3 alone is not responsible for the strong impairment of matrix Ca$^{2+}$ uptake in the mcu1 2 3 background, since the mcu3 line showed Ca$^{2+}$ dynamics similar to those of the Col-0 control (Supplemental Figure S12). The idea that the three MCU proteins act redundantly and that the MCU capacity is increasingly limited in higher-order mutants is supported by the finding that defects in Ca$^{2+}$ uptake are dependent on the intensity of the stimulus and that the impairment of the Ca$^{2+}$ uptake rate was already pronounced in mcu1 2 root tips when exposed to 2 mM eATP (down to 65% of Col-0 in ROI2 and 44% of Col-0 in ROI9; Figure 4, D and J). It can be concluded that (1) MCU proteins provide the dominant path for Ca$^{2+}$ to rapidly enter the mitochondria in root tips and (2) any remaining MCU4, MCU5, and MCU6 proteins combined provide at most a minor fraction of the total capacity for rapid mitochondrial Ca$^{2+}$ uptake (24% in ROI2; 10% in ROI9; Figure 4, D and J). However, it was also evident that very slow rates of Ca$^{2+}$ accumulation in the matrix, which were characteristic during hypoxia (Supplemental Figure S19, D and E), were not affected in mcu1 2 3 seedlings. In that case, it appears that the remaining capacity is not limiting.

We cannot and do not aim to assess the relative quantitative contributions of the individual MCU loci to mitochondrial Ca$^{2+}$ uptake capacity. Instead, we provide a system that offers different degrees of limited total MCU capacity in vivo. We focused our investigations on Arabidopsis root tips of young seedlings as a model system for highly standardized in vivo analyses, but it is possible that other tissues express alternative transporters. A direct impact on mitochondrial Ca$^{2+}$ dynamics in green tissues was not systematically assessed here. Yet, circumstantial support for the functional importance of MCU1, MCU2, and MCU3 in the shoot comes from the finding that the transcript responses were similar in the roots and shoots of mcu1 2 3 (Figure 6D; Supplemental Figure S21D). Future work will need to

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**Figure 6 (Continued)**

Quantification of OPDA, dn-OPDA, JA, and JA-Ile in roots of Col-0 and mcu1 2 3 seedlings. *n* = 5; error bars = SD. *P* < 0.05 (Student’s *t* test). F, Representative images of Col-0 wild-type and mcu1 2 3 plants with and without touch treatment (touched, untouched) applied for 4 weeks twice daily from 14 days after transfer to the growth room. G, Percentage of bolting plants over the growth period. T, touched; UT, untouched. *n* = 11–13 plants. H, Day of bolting of Col-0 and mcu1 2 3 plants. T, touched; UT, untouched. Boxes extend from the 25th to 75th percentiles, the line in the box represents median, and the whiskers represent the minimum and maximum values. *n* = 11–13 plants. I, Stem height of Col-0 and mcu1 2 3 plants 39 days after sowing. T, touched; UT, untouched. *n* = 10 plants. Error bars = SEM. J, Representative images of mcu1 2 3 and Col-0 wild-type 27 days after sowing. T, touched; UT, untouched. K, Rosette area of mcu1 2 3 and Col-0 wild-type plants 27 days after sowing. T, touched; UT, untouched. *n* = 11–13 plants. Error bars = SEM. Different letters indicate significant differences between all samples at *P* < 0.05 (two-way ANOVA with posthoc Tukey’s test).
explore the potential importance of individual MCU isoforms in different tissues as well as potential alternative mitochondrial Ca\textsuperscript{2+} uptake pathways (Wagner et al., 2016; Carraretto et al., 2016a). Since the suppression of MCU was also found to impair mitochondrial Ca\textsuperscript{2+} transients in several animal model systems (De Stefani et al., 2011; Pan et al., 2013; Xu and Chisholm, 2014; Holmström et al., 2015; Tufi et al., 2019), the dominant role of MCU amongst the mitochondrial Ca\textsuperscript{2+} uptake mechanisms appears to be evolutionarily conserved among plants and metazoans.

MCUs set the baseline of plant mitochondrial free Ca\textsuperscript{2+} concentrations
Reduced matrix free Ca\textsuperscript{2+} concentrations at resting state were previously observed in mcu loss-of-function lines of fruit fly (Drosophila melanogaster) and mouse (Mus musculus) (Pan et al., 2013; Holmström et al., 2015; Tufi et al., 2019). An assessment of the characteristic tissue-specific pattern of resting matrix free Ca\textsuperscript{2+} levels in Arabidopsis root tips (Figures 1, E, 2, B, I, P, 3, B, and 4, A; Supplemental Figures S8, A and S9, A; Supplemental Movies S1–S5) (Loro et al., 2012; Wagner et al., 2015) revealed that the matrix free Ca\textsuperscript{2+} gradient from the division zone to the elongation zone was flattened in the absence of MCU1, MCU2, and MCU3 (Figure 3, B and C). This effect suggests that MCU-mediated Ca\textsuperscript{2+} uptake is a decisive factor in setting free Ca\textsuperscript{2+} levels even at steady-state and that this role of MCUs is evolutionarily conserved between animals and plants.

Mitochondrial Ca\textsuperscript{2+} uptake does not shape cytosolic Ca\textsuperscript{2+} signatures in roots
In contrast to the mitochondrial matrix, MCU ablation had no measurable effect on cytosolic Ca\textsuperscript{2+} transients in Arabidopsis root tips (Figure 5; Supplemental Figure S13). Since our analysis was limited in resolution to tissue averages and due to our focus on root tips, differences in other tissues and at the cellular level or in cytosolic microdomains remain possible. In line with the observations here, the opposite situation, that is increased mitochondrial Ca\textsuperscript{2+} uptake capacity in Arabidopsis mcu lines did not show any detectable impact on cytosolic Ca\textsuperscript{2+} dynamics either (Wagner et al., 2015). In contrast, mitochondrial Ca\textsuperscript{2+} uptake activity in animal systems, as mediated by the mitochondrial Ca\textsuperscript{2+} uniporter, was reported to act as a capacitor and to shape the cytosolic Ca\textsuperscript{2+} signatures, at least for specific cell types (De Stefani et al., 2011; Drago et al., 2012; Prudent et al., 2013; Rasmussen et al., 2015). Cytosolic free Ca\textsuperscript{2+} levels may be particularly strongly buffered to compensate for changes in Ca\textsuperscript{2+} sequestration activity by the mitochondria. A contribution to this buffering capacity may be made by other organelles that are known to actively participate in intracellular Ca\textsuperscript{2+} dynamics in plants, for example, the vacuole, the endoplasmic reticulum, and the plastids (Logan and Knight, 2003; Iwano et al., 2009; Bonza et al., 2013; Schönknecht, 2013; Loro et al., 2016; Corso et al., 2018; Sello et al., 2018; Shkolnik et al., 2018; Frank et al., 2019; Teardo et al., 2019; Dindas et al., 2021; Resentini et al., 2021a). The impact of impaired mitochondrial Ca\textsuperscript{2+} transport on other organelles is beyond the scope of this study but deserves further attention, especially in the light of the observed JA-related phenotype.

Impairment of mitochondrial Ca\textsuperscript{2+} uniport in vivo causes a spectrum of organismal phenotypes
The biochemical properties of the mitochondrial Ca\textsuperscript{2+} uniporter complexes from different metazoan species have been resolved to the highest detail by X-ray and cryo-EM studies, resulting in a comprehensive understanding of the structural basis of their function and regulation (Baradaran et al., 2018; Fan et al., 2018, 2020; Nguyen et al., 2018b; Wang et al., 2019), pharmacology (Arduino et al., 2017; Paillard et al., 2018; Di Marco et al., 2020), and the functions of their components in positioning within the mitochondrial membrane (Gottschalk et al., 2019). Also, a growing number of pathologies have been linked to defects in uniporter function in mammals and other metazoans (Logan et al., 2014; Kwong et al., 2015; Mammucari et al., 2015; Antony et al., 2016; Lewis-Smith et al., 2016; Liao et al., 2017; Debattisti et al., 2019). The functional in vivo significance of the uniporter remains difficult terrain, however, and has been under debate (Murphy et al., 2014; Pendin et al., 2014). Specifically, an mcu knockout mouse line was viable and showed surprisingly mild phenotypes (Pan et al., 2013; Pendin et al., 2014; Holmström et al., 2015). Severe phenotypic impairment or lethality was predicted for animal models based on the function of matrix Ca\textsuperscript{2+} dynamics in controlling cell death pathways and respiratory activity (reviewed in [Wagner et al., 2016]). In different mcu mouse models, the functions of even cardiac tissues and brown fat were largely uncompromised (Luongo et al., 2015; Flicker et al., 2019). Consistently mild phenotypes were observed in response to impaired mitochondrial Ca\textsuperscript{2+} uptake in fruit fly (Tufi et al., 2019) and nematode (Caenorhabditis elegans) (Xu and Chisholm, 2014), while severe developmental defects occurred in zebrafish and Trypanosoma brucei lacking MCU (Huang et al., 2013; Prudent et al., 2013).

Despite the severe impairment in mitochondrial Ca\textsuperscript{2+} uptake and baseline maintenance in the Arabidopsis mcu1 2 3 line, plant growth and development were not visibly compromised in the large majority of instances that we assessed (Supplemental Figures S4, B–D; S14–19; S22). The absence of an obvious organismal phenotype despite a pronounced cell physiological phenotype is reminiscent of recent observations in Arabidopsis chloroplasts lacking two POLLUX ion channel family proteins, PLASTID ENVELOPE ION CHANNEL 1 and 2 (PEC1/2) (Völknert et al., 2021). The strict retention of at least one MCU gene in genomes across the plant phylogeny indicates a selective advantage provided by MCU genes. For single loss-of-function lines of MCU1 and MCU2, impaired root growth and decreased male fertility associated with decreased in vitro pollen tube germination and growth were previously reported, respectively (Teardo...
et al., 2017; Selles et al., 2018), in the absence of drastic impairment of mitochondrial Ca²⁺ uptake. We were unable to observe similar phenotypes in the mcu1 2 3 line (Supplemental Figures S15 and S17). If the phenotypes reported for the single mutants were associated with mitochondrial Ca²⁺ uptake, the arising phenotypes may be expected to be at least the same in the mcu1 2 3 line, where mitochondrial Ca²⁺ uptake is severely compromised. Since this was not observed, the phenotypes may be independent of Ca²⁺ transport function, and the effect may have been lost by compensation through yet unknown factors in the triple mutant. Alternatively, the reported phenotypes may arise only under specific experimental conditions, such as those in the previous studies, and those conditions may not have been replicated with the required precision here.

Although broad phenotyping efforts did not reveal major effects on growth and development of the mcu1 2 3 line under most conditions, we observed slightly less chlorophyll degradation in response to whole plant darkening (Supplemental Figure S20). This difference is in line with the increased transcript abundance of MCU1 and 3 that was previously observed during developmental and dark-induced senescence (Chrobok et al., 2016; Law et al., 2018). Since whole plant darkening will initially trigger a metabolic starvation response, followed by the induction of senescence (Law et al., 2018), the specific involvement of MCU in dark starvation and/or leaf senescence deserves future dissection, along with the analysis of mitochondrial Ca²⁺ dynamics in mature leaves that face this treatment. While it is not possible to comprehensively cover growth conditions, biological processes, and genotype–environment interactions, the observed phenotypes provide a handle on the physiological importance of mitochondrial Ca²⁺ transport. How exactly these phenotypes arise deserves focused analysis. Mechanistic analysis is likely to offer insight into the role of plant mitochondrial Ca²⁺ uptake in the integration of external stimuli, hormonal regulation, and growth responses.

Decreasing MCU capacity in plants selectively affects JA homeostasis

The lack of global reprogramming of the mcu1 2 3 seedling root transcriptome (Figure 6A) is in line with the absence of obvious gross pleiotropic phenotypes under a range of conditions that we tested. Instead, the selective enrichment and concerted repression of JA-related transcripts and doubling in basal JA and JA-Ile levels hint at a link between mitochondrial Ca²⁺ dynamics and JA signaling (Figure 6, B–E; Supplemental Figure S21). It seems likely that the decreased responsiveness to touch in mcu1 2 3 plants, as manifested by less pronounced thigmomorphogenesis responses (Figure 6, F–K), is linked to deregulated JA homeostasis. However, this hypothesis will need to be tested. The observation of increased chlorophyll content in mcu1 2 3 rosettes after whole plant darkening (Supplemental Figure S20) might also be linked to the deregulation of JA. JA was previously found to induce leaf senescence via the repression of Rubisco activase (Shan et al., 2011), and key photosynthetic proteins were repressed in response to wounding in a JA-dependent manner (Geller et al., 2011). The mechanistic nature of the unexpected link to JA, and the apparent uncoupling of the positive feedback regulation of JA, lay the groundwork for further study, especially since our understanding of the relationship between mitochondria and plant hormonal regulation is still in its infancy (Berkowitz et al., 2016). The interaction between Ca²⁺ signaling and JA signal transduction is known to occur in the cytosol and nucleus, where JA induces Ca²⁺ mobilization to modulate the expression of JA-response genes (Sun et al., 2010; Yan et al., 2018), and CPKs as well as CBL–CIPK complexes can act as regulators of JA signaling (Munemasa et al., 2011; Förster et al., 2019). Indeed, Ca²⁺ signaling is required for the induction of JA biosynthesis (Wasternack, 2007; Toyota et al., 2018; Meena et al., 2019), and eATP signaling involving intracellular Ca²⁺ transients (as exploited in this work) was shown to affect JA signaling to optimize plant defense responses (Tripathi et al., 2018). Interestingly, basal JA pathway activity is markedly dependent on endomembrane cation flux capacities, as observed in mutants with altered activity of the Ca²⁺-regulated channel TPC1 in the tonoplast (Bonaventure et al., 2007; Lenglet et al., 2017).

Despite the lack of a similar mechanistic framework as for the relationship between Ca²⁺ and JA, mitochondria act as hubs that sense and integrate environmental and cellular signals to trigger specific responses to various stimuli to initiate signaling pathways that modulate nuclear gene expression to optimize growth and development (Ng et al., 2014). Genetic or chemical perturbations of mitochondrial functions result in transcriptional programming linked to plant hormone functions (Van Aken and Whelan, 2012; Zhang et al., 2014) and alter phytohormone levels (e.g. abscisic acid, gibberellic acid, and metabolites of the auxin indole-3-acetic acid) (Xu et al., 2019).

Conversely, several transcripts encoding mitochondrial proteins, such as OM66 and DIC2, have been found to be induced by mechanical signaling, which is largely JA-dependent, and mitochondrial metabolite levels change shortly after mechanical treatment (Aken et al., 2016). Interestingly, mechanical stimuli that trigger JA signaling were also shown to trigger characteristic mitochondrial Ca²⁺ transients in Arabidopsis seedlings (Logan and Knight, 2003; Teardo et al., 2015). Mitochondrial OM64 deficiency in rice (Oryza sativa) causes resistance to sucking and piercing insects by stimulating JA production and signaling (Guo et al., 2020). These findings, as well as the current findings, suggest a tight interaction between mitochondria, JA responses, and touch signaling, although the exact causality of these events remain to be determined.

The key advance of this study lies in the demonstration that a conserved mitochondrial Ca²⁺ uniporter system operates in planta and dominates matrix Ca²⁺ dynamics. Several critical questions about the role of mitochondrial
Ca\textsuperscript{2+} signaling in plants arise that can now be addressed due to the ability to gradually constrain mitochondrial Ca\textsuperscript{2+} transport capacity. Most importantly, plant Ca\textsuperscript{2+} signaling has been extensively studied but with a focus on transport across the plasma membrane and the tonoplast and on Ca\textsuperscript{2+} sensing in the cytosol and nucleus. A comprehensive understanding of cellular Ca\textsuperscript{2+} signaling will only be achieved, however, if we also integrate the other cell compartments into this picture (Kudla et al., 2018; Resentini et al., 2021b). Several different functions have been associated with mitochondrial Ca\textsuperscript{2+} transport in mammals. It will now be feasible to dissect which of those functions have been conserved and which plant-specific functions have been acquired during evolution. Already in this study we identified an unexpected connection with JA signaling and the touch response. Obvious questions to address in the future include how the MCU-JA-touch link is underpinned mechanistically and what additional downstream consequences JA deregulation may have on plant fitness.

Materials and methods

Plant lines, transformation, and culture conditions

Arabidopsis thaliana Col-0 mutant lines mcu1 (SALK_082151 (Teardo et al., 2017)), mcu2 (WiscDsLox393-396L9; Selles et al., 2018), and mcu3 (SALK_019312) were obtained from the Nottingham Arabidopsis Stock Centre and validated by genotyping and sequencing. The primers used for the genotyping are listed in Supplemental Table S1. Double and triple mutants were obtained by crossing, and wild-type control lines were obtained by segregation. The generation of Col-0 lines expressing mitochondrial 4mt-YC3.6 and cytoplasmic NES-YC3.6 was described before (Krebs et al., 2012; Loro et al., 2012). The mcu1, mcu1 2, and mcu1 2 3 lines were transformed with 4mt-YC3.6 or NES-YC3.6 using the floral dip method (Clough and Bent, 1998). For each construct, independent transgenic lines were selected based on antibiotic resistance and fluorescence. For mcu1 2 3, all experiments were recorded using two independent biosensor lines. To generate the 3SS:MCU2 lines, Col-0 and mcu1 2 3 plants expressing 4mt-YC3.6 were transformed by floral dip using the pB7GW2/MCU2 construct. T1 transformants of the Col-0 background were selected using BASTA resistance as a marker (5 days after germination in soil treated with BASTA 3 times every other day). Surviving individuals were assessed by genotyping for the presence of the 3SS:MCU2 T-DNA. Since the mcu1 2 3 line is already resistant to BASTA due to the presence of a BASTA resistance marker gene in the T-DNA in the endogenous MCU2 locus, BASTA application was used to enrich for individuals carrying the 3SS:MCU2 insertion. Individuals that were markedly larger than the majority after BASTA application were selected and subsequently assessed by genotyping for the presence of the 3SS:MCU2 T-DNA and by RT–qPCR for transgene expression.

Seeds were surface sterilized and stratified at 4°C in the dark for 2 days. Seedling culture was performed in growth chambers set to long day conditions (16 h at 21°C and 90 μE m\textsuperscript{2} s\textsuperscript{-1}, 8 h at 17°C and darkness; Philips TL5 HO 49W/840 [MASTER] Cool White tubes). For Ca\textsuperscript{2+} imaging, seedlings were grown vertically for 4–5 days (Loro et al., 2012) on plates containing half-strength Murashige and Skoog medium (MS; Duchefa, Haarlem, the Netherlands (Murashige and Skoog, 1962)) supplemented with 0.1% (w/v) sucrose and 0.05% (w/v) 2-(N-morpholino)ethanesulfonic acid (MES), adjusted to pH 5.8 (KOH), and 0.8% (w/v) plant agar (Duchefa, Haarlem, the Netherlands). For RT–qPCR analysis of MCU gene expression, 14-day-old seedlings were grown vertically on half-strength MS medium supplemented with 0.1% (w/v) sucrose, 0.05% (w/v) MES and 0.8% (w/v) plant agar, pH 5.8 (KOH). For microarray experiments, RT–qPCR analyses of JA- and stress-related genes and phytohormone measurements, seedlings were cultured on a mesh in sterile hydroponics boxes (Phytotray II; Sigma-Aldrich, Darmstadt, Germany) filled with 150 mL half-strength MS medium supplemented with 0.1% (w/v) sucrose and 0.05% (w/v) MES, pH 5.8 (KOH). After 20 days, roots were separated from the shoot with a sharp blade, immediately frozen in liquid nitrogen, and stored at −80°C before RNA extraction. For subcellular localization studies of MCU3-GFP, Agrobacterium suspensions were infiltrated into the leaves of 4- to 5-week old Nicotiana benthamiana plants with a needleless syringe, and transformed areas were used for imaging after 2–4 days.

Plant phenotyping analyses

The individual culture and treatment conditions for the phenotypic analyses are described in the respective figure legends. The touch treatments to assess thigmomorphogenesis were carried out as described recently (Darwish et al., 2022).

Molecular cloning

The coding sequence of MCU3 was amplified from Arabidopsis Col-0 cDNA using primers from Supplemental Table S1 and inserted into the pDONR207 vector (Invitrogen, Carlsbad, CA, USA) using Gateway BP-Clonase II (Invitrogen) according to the manufacturer’s instructions. Gateway LR recombination was used with destination vector pUBC-GFP-Dest (Grefen et al., 2010) to generate MCU3-GFP under the control of the UBQ10 promoter. The construct was introduced into Agrobacterium tumefaciens AGL1 cells by electroporation and used for transient expression in N. benthamiana leaves (Sparkes et al., 2006).

Analogously, to generate 3SS:MCU2 lines, the MCU2 coding sequence was amplified from Col-0 cDNA using primers from Supplemental Table S1 and inserted into the pDONR207 vector (Invitrogen) using Gateway BP-Clonase II (Invitrogen) according to the manufacturer’s instructions. Positive clones were confirmed by sequencing. Gateway LR recombination was used with pB7GW2 (Karimi et al., 2002) as a destination vector to place MCU2 under the control of the CaMV35S promoter. The construct was then introduced into A. tumefaciens C58C1 cells.
Phylogenetic analysis
The amino acid sequences of human MCU and MCUb (UniProt ID Q8NE86 and QENWR8, respectively) were used to retrieve homologous sequences in A. thaliana and M. musculus through a BlastP search (Supplemental Data Set 1). Sequences were aligned by Muscle (Edgar, 2004) in MEGA version 6 (Tamura et al., 2013) with default parameters (Supplemental File S1) and clustered in an unrooted maximum likelihood tree using the JTT matrix-based method (Jones et al., 1992), uniform rates among sites (MEGA version 6). The tree with the highest log likelihood (~5697.26) is shown in Figure 1A. The bootstrap values indicated at the nodes were from 1,000 repetitions.

RNA extraction and RT–qPCR analysis
Total RNA was extracted from pooled shoots or roots of 20-day-old seedlings grown under hydroponic conditions or from 2-week-old seedlings vertically grown in Petri dishes using a Nucleospin RNA Plant kit, treated with DNase I (both from Macherey-Nagel, Düren, Germany), and used for RT–qPCR analyses of JA- and stress-related genes or MCU gene expression, respectively. RNA (500 ng) was reverse transcribed using an iScript cDNA Synthesis Kit (BioRad, Hercules, CA, USA). A “no RT”-reaction, in which the RNA was subjected to the same conditions of cDNA synthesis but without reverse transcriptase, was included as a negative control in all RT–qPCR analyses. RT–qPCR was performed with SYBR Green detection (KAPA SYBR Fast qPCR Master Mix, Roche, Mannheim, Germany) in a C1000 Touch Thermal Cycler using a Nucleospin RNA Plant kit, treated with DNase I (both from Macherey-Nagel, Düren, Germany), and used for RT–qPCR analyses of JA- and stress-related genes or MCU gene expression, respectively. RNA (500 ng) was reverse transcribed using an iScript cDNA Synthesis Kit (BioRad, Hercules, CA, USA). A “no RT”-reaction, in which the RNA was subjected to the same conditions of cDNA synthesis but without reverse transcriptase, was included as a negative control in all RT–qPCR analyses. RT–qPCR was performed with SYBR Green detection (KAPA SYBR Fast qPCR Master Mix, Roche, Mannheim, Germany) in a C1000 Touch Thermal Cycler using the CFX 96 Real-Time System (Bio-Rad, Hercules, CA, USA) and specific primers (Supplemental Table S1). Data were analyzed using the DDC_T method/comparative CT method. The transcript levels were normalized to that of the SAND family protein (SAND FP, At2g28390; Czechowski et al., 2005), and the relative expression levels for each analyzed gene were determined by comparative CT methods (Schmittgen and Livak, 2004). All RT–qPCR reactions were run in triplicates using total RNA preparations from independent pools of plants that were grown side-by-side under identical experimental conditions.

Transcriptome analysis
Gene expression profiling was performed using Arabidopsis AraGene_1.0.0-st-type arrays from Affymetrix. RNA was extracted from the samples as described for RT–qPCR analysis from five independent replicates of pooled root tissue. Biotinylated antisense cDNA was prepared according to the Affymetrix standard labeling protocol with a GeneChip WT Plus Reagent Kit and a GeneChip Hybridization, Wash and Stain Kit (both from Affymetrix, Santa Clara, USA). Afterwards, hybridization on the chip was performed in a GeneChip Hybridization oven 640, dyed in the GeneChip Fluidics Station 450, and scanned with a GeneChip Scanner 3000. All of the equipment used was from the Affymetrix Company (Affymetrix, High Wycombe, UK).

Bioinformatic analysis of transcriptome data
A Custom CDF version 22 with TAIR based gene definitions was used to annotate the arrays (Dai et al., 2005). The raw fluorescence intensity values were normalized by applying quantile normalization and robust multichip average (RMA) background correction. An analysis of variance (ANOVA) model, using genotype and treatment as fixed effects, was performed to identify differentially expressed genes using a commercial software package (SAS JMP10 Genomics, version 6, SAS Institute, Cary, NC, USA). A false positive rate of $a = 0.05$ with false discovery rate (FDR) correction was taken as the level of significance. Downregulated genes (FC $≤ −1.5$ and $P$-adjusted $< 0.05$) in $muc1 2 3$ were analyzed for gene ontology (GO) term enrichment using GOrilla (http://cbl-gorilla.cs.technion.ac.il/) (Eden et al., 2009).

Time lapse in vivo Ca2+ imaging of root tips
Four to 5-day-old seedlings were used for root imaging as optimized and described previously (Loro et al., 2012; Wagner et al., 2015; Behera et al., 2018). Briefly, seedlings were gently placed into a dedicated custom perfusion chamber and stabilized with cotton wool soaked in imaging solution (5 mM KCl, 10 mM MES, and 10 mM CaCl₂, pH 5.8 (Tris)). The root was continuously perfused with imaging solution, whereas the shoot was not submerged. The seedlings were kept under continuous perfusion for 10 min before ratiometric image acquisition as an optimization of a previous protocol (Teardo et al., 2017) in order to minimize the potential impact of specimen manipulation on the measurements. Treatments were performed by supplementing the imaging solution with 0.01 mM NAA (from a 10.74 mM stock solution) or with sodium adenosine triphosphate (Na₂ATP, working solution of 0.01, 0.1, or 2 mM from a 200 mM stock solution buffered at pH 7.4 with NaOH). The 200 mM D-sorbitol solution was prepared as described for standard imaging solution with the direct addition of the sugar powder (5 mM KCl, 10 mM MES, 10 mM CaCl₂, and 200 mM D-sorbitol, pH 5.8 (Tris)). The solutions were administrated for 3 min under running perfusion and then washed out. The experiments conducted using wild-type and mutant seedlings were performed alternatingly. An inverted fluorescence microscope (Ti-E Nikon) was used for time lapse acquisition with a CFI PLAN APO 20 × VC dry objective for stimulation with NAA and eATP, or a CFI Plan Fluor 4 × dry objective for stimulation with D-sorbitol. Excitation light was produced by a Prior Lumen 200 PRO fluorescent lamp (Prior Scientific) at 440 nm (436/20 nm) set to 20%. Images were collected with a dual charge-coupled device camera (ORCA-D2; Hamamatsu). For Cameleon analysis, the FRET CFP/YFP optical block A11400-03 (emission 1 483/32 for CFP and emission 2 524/27 for FRET) with a dichroic mirror (Hamamatsu) was used for the simultaneous measurement of CFP and cpVenUS acquisitions. Images were acquired every 5 s. For the roots of seedlings expressing the mitochondrial Cameleon, the exposure times were 100–500 ms (depending on the magnification used) with a 4 × 4 pixel binning, while...
for the cytosolic Cameleon, the exposure times were 100 ms with a 2 x 2 pixel binning. Filters and dichroic mirrors were from Chroma Technology. NIS-Elements (Nikon) was used as a platform to control the microscope, illuminator, and camera.

Ratiometric image data analysis
Individual ratiometric images and time-lapse imaging data were analyzed using a custom MatLab program package (Fricker, 2016). Unless otherwise indicated, the fluorescence intensity was determined over ROIs, which corresponded to the entire root tip. The cpVenus and CFP emissions of the analyzed ROIs were used for the ratio (R) calculations (cpVenus/CFP). Background subtraction was performed independently for both channels before calculating the ratio. To analyze the sorbitol treatments, we employed the Template Matching Plugin (normalized cross-correlation configuration in the Drop menu of Fiji’s Matching method) for ImageJ and Fiji software. The plugin was necessary to eliminate root-tip movements due to the change in cells turgor, as determined by examining the change in the osmotic potential of the solution, hence eliminating measurement artifacts due to specimen movement.

Confocal microscopy
Confocal imaging was performed with a Leica SP5 inverted laser scanning microscope using a 25 x lens (HCX IRAPO L, 0.95 numerical aperture, water immersion, zoom 1.5 was used) or a 63 x lens (HCX PL APO, 1.20 numerical aperture, water immersion, zoom 10 x was used). To visualize cell wall organization, roots were stained by incubating in the dark for 5 min in 1 µg mL⁻¹ propidium iodide (P4864; Sigma-Aldrich, Darmstadt, Germany) solution (from a 2 mg mL⁻¹ stock solution diluted in ddH₂O) and rinsed two times in water (Alassimone et al., 2010). Propidium iodide was monitored with a 514 nm excitation wavelength and a 564–660 nm bandpass emission filter. For colocalization studies with MitoTracker, plant material was submerged for 30 min with 200 nM MitoTracker Orange CMTMros (M7510, Invitrogen). Fluorescence of cpVenus and MitoTracker was measured at 514 nm; emission was collected at 520–550 and 570–620 nm, respectively.

Confocal imaging of MCU3-GFP was performed with a Leica TCS SP8X laser scanning microscope using a 63 x lens (HC PL APO CS2, 1.20 numerical aperture, water immersion). GFP and chlorophyll were excited at 488 nm using a flexible pulsed white-light laser, and emission was collected using hybrid detectors at 505–530 and 650–700 nm, respectively. Minor chloroplastic signal in the GFP channel was observed when using standard confocal microscopy settings as described in a previous report (Carraretto et al., 2016b) but was entirely removed by the time gating function of the microscope, which allows fluorophores to be separated out based on their fluorescence lifetimes. For the data shown in Supplemental Figure S5, GFP detection was time-gated at 0.3–12.0 ns (Kodama, 2016).

Photosynthetic efficiency
Measurements of efficiency of photosystem II (Fv/Fm) were performed on true leaves of 32-day-old plants as described previously (Teardo et al., 2017).

Phytohormone measurements
Measurements of OPDA, dnOPDA, JA, and JA-Ile were performed on roots of 20-day-old seedlings grown on mesh in sterile hydroponics boxes. Phytohormones were quantified simultaneously using a standardized Ultra-performance liquid chromatography–tandem Mass Spectrometry (UPLC–MS/MS)-based method according to Balcke et al. (2012). In brief, 50 mg of frozen samples were extracted with 500 µL methanol supplied with [²H₃]OPDA, [²H₃]JA, and [²H₃]JA-Ile (50 ng each) as internal standards. After centrifugation, the supernatant was diluted with nine volumes of water and subjected to solid phase extraction on HR-XC (Chromabond, Macherey-Nagel, Düren, Germany). Elution was done with 900 mL acetonitrile, and 10 µL of the eluate were subjected to UPLC–MS/MS. The phytohormone contents were calculated using the ratio of analyte and internal standard peak heights, thereby calculating the content of dnOPDA using [²H₃]OPDA as an internal standard.

Immunoblotting
Proteins were extracted from frozen seedling tissue (50 mg) by adding 200 µL of extraction buffer (25 mM Tris–Cl pH 6.8, 1% SDS, 1% (v/v) β-mercaptoethanol, 1% (v/v) Halt protease inhibitor cocktail [Sigma-Aldrich, #78442]) and heating at 95°C for 5 min. After centrifugation for 10 min, the supernatant was collected and proteins quantified using Coomassie Plus Reagent (Sigma-Aldrich, #23236). Ten microgram total protein was mixed with 2 x Laemmlli Buffer (1:1) and heated at 96°C for 10 min to dissolve AOC trimers. Proteins were separated by SDS–PAGE (12% acrylamide) and transferred to a PVDF membrane. Detection of protein bands was done by staining with Ponceto S. The membrane was blocked with 5% (w/v) BSA in TBST (20 mM Tris–Cl pH 7.8, 150 mM NaCl, 0.05% [v/v] Tween) and immunostained using anti-AtAOC antibody (1:5,000, Stenzel et al., 2003) and a goat anti-rabbit IgG antibody conjugated with alkaline phosphatase (1:5,000, Sigma-Aldrich). Chemiluminescence detection was performed with ImmunStar AP Substrate (ThermoFisher Scientific, Waltham, MA, USA, #1705018) for 5 min and visualized using a Fusion FX Imaging system (Vilber, www.vilber.com).

Statistical analysis
The statistical analyses applied to the individual datasets are detailed in the respective figure legends, and a summary is provided in Supplemental Data Set 5. All statistical tests were conducted using Microsoft Excel and GraphPad Prism.

Accession numbers
Sequence data from this article can be found in the GenBank/EMBL libraries under the following accession numbers: AGC2-1, At3g25250; AOS, At5g42650; CPK28,
At5g66210; GRX480, At1g28480; LOX3, At1g17420; JAZ1, At1g19180; JAZ2, At1g74950; JAZ3, At3g17860; JAZ5, At1g17380; JAZ6, At1g72450; JAZ8, At1g30135; JAZ9, At5G20900; JAZ10, At5g13220; JAZ12, At1g70700; MCU1, At1g09575; MCU2, At1g57610; MCU3, At2g23790; MCU4, At4g36820; MCU5, At5g42610; MCU6, At5g66650; MYC2, At1G23640; NPR3, At5g45110; OPR3, At2g06050; SAND family protein, At2g28390; WRKY40, At1g80840; WRKY53, At4g23810; UBC21, At5g25760; UBO10, and At4g05320.

The following materials are available in the online version of Supplemental data.

Supplemental Figure S1. Protein alignment and predicted topology of MCUs in A. thaliana.

Supplemental Figure S2. Generation and validation of the mcu1 2 line.

Supplemental Figure S3. Subcellular localization of 4mt-YC3.6 in Col-0 and mcu1 2.

Supplemental Figure S4. Primary root tip architecture is unchanged in mcu1 2 3 seedlings.

Supplemental Figure S5. MCU3-GFP localizes to mitochondria in N. benthamiana epidermal cells.

Supplemental Figure S6. Generation and validation of the mcu1 2 3 line.

Supplemental Figure S7. Subcellular localization of 4mt-YC3.6 in mcu1 2 3.

Supplemental Figure S8. The mitochondrial matrix Ca\(^{2+}\) transient triggered by 0.1 mM eATP shows progressive impairment with progressive ablation of MCUs.

Supplemental Figure S9. The mitochondrial matrix Ca\(^{2+}\) transient triggered by 0.01 mM eATP shows progressive impairment with progressive ablation of MCUs.

Supplemental Figure S10. The mitochondrial matrix Ca\(^{2+}\) transients induced by 0.01 mM NAA or 200 mM D-sorbitol show consistent impairment in mcu1 2 3 compared to Col-0 as for other stimuli.

Supplemental Figure S11. Generation and validation of 35S:MCU2 lines.

Supplemental Figure S12. The mitochondrial matrix Ca\(^{2+}\) transient triggered by 2.0 mM eATP in root tips of mcu3 and 35S:MCU2 lines.

Supplemental Figure S13. Cytosolic Ca\(^{2+}\) transients in root tips induced by treatment with 2.0 mM eATP are unchanged for all ROIs in mcu1 2 3.

Supplemental Figure S14. Gross plant development is unchanged in mcu1 2 3.

Supplemental Figure S15. Seedling development shows no obvious changes in mcu1 2 3.

Supplemental Figure S16. Primary root growth shows no obvious changes in mcu1 2 3 under different external treatments.

Supplemental Figure S17. Pollen tube germination and growth, and root hair development are unchanged in mcu1 2 3.

Supplemental Figure S18. mcu1 2 3 and Col-0 seedling growth in the presence of antimycin A (AA).

Supplemental Figure S19. Submergence and low oxygen responses in Col-0 and mcu1 2 3 plants.

Supplemental Figure S20. Whole plant darkening to induce starvation and senescence in Col-0 and mcu1 2 3 plants.

Supplemental Figure S21. Repression of JA homeostasis in mcu1 2 3 seedlings.

Supplemental Figure S22. Root penetration growth of Col-0 and mcu1 2 3 in media with different agar concentrations.

Supplemental Table S1. List of primers.

Supplemental Data Set 1. Sequence information used for phylogenetic analysis.

Supplemental Data Set 2. List of 159 downregulated genes in mcu1 2 3 compared to Col-0.

Supplemental Data Set 3. List of 13 upregulated genes in mcu1 2 3 compared to Col-0.

Supplemental Data Set 4. List of 14 genes repressed in mcu1 2 3 and present in the MYC2 regulon dataset.

Supplemental Data Set 5. Summary of statistical analyses.

Supplemental File S1. Sequence alignments for phylogenetic analysis.

Supplemental File S2. MCU transcript abundance in different Arabidopsis organs and during development as observed in publicly available transcriptome analyses and curated by the Genevestigator platform.

Supplemental Movie S1. Mitochondrial Ca\(^{2+}\) transients in response to stimulation for 3 min with 0.01 mM NAA in living Arabidopsis root tips of Col-0, mcu1, and mcu1 2.

Supplemental Movie S2. Mitochondrial Ca\(^{2+}\) transients in response to stimulation for 3 min with 0.01 mM eATP in living Arabidopsis root tips of Col-0, mcu1, and mcu1 2.

Supplemental Movie S3. Mitochondrial Ca\(^{2+}\) transients in response to stimulation for 3 min with 0.10 mM eATP in living Arabidopsis root tips of Col-0, mcu1, and mcu1 2.

Supplemental Movie S4. Mitochondrial Ca\(^{2+}\) transients in response to stimulation for 3 min with 2.0 mM eATP in living Arabidopsis root tips of Col-0, mcu1, and mcu1 2.

Supplemental Movie S5. Mitochondrial Ca\(^{2+}\) transients in response to stimulation for 3 min with 2.0 mM eATP in living Arabidopsis root tips of Col-0 and mcu1 2 3.

Supplemental Movie S6. Cytosolic Ca\(^{2+}\) transients in response to stimulation for 3 min with 2.0 mM eATP in living Arabidopsis root tips of Col-0 and mcu1 2 3.

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