The role of *FMT* and *FIS1A* in mitochondrial morphology and salt stress in *Arabidopsis thaliana*

Inaugural-Dissertation

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<tr>
<td>°C</td>
<td>degree Celsius</td>
</tr>
<tr>
<td>%</td>
<td>percent</td>
</tr>
<tr>
<td>cDNA</td>
<td>complementary DNA</td>
</tr>
<tr>
<td>Col-0</td>
<td>Columbia-0</td>
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<tr>
<td>DNA</td>
<td>deoxyribonucleic acid</td>
</tr>
<tr>
<td>dNTP</td>
<td>deoxynucleoside triphosphate</td>
</tr>
<tr>
<td>E. coli</td>
<td>Escherichia coli</td>
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<tr>
<td>e.g.</td>
<td>exempli gratia (Latin) for example</td>
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<tr>
<td>et al.</td>
<td>et alterni (Latin) and others</td>
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<tr>
<td>EtOH</td>
<td>ethanol</td>
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<td>Fig.</td>
<td>figure</td>
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<td>hr</td>
<td>hour</td>
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<tr>
<td>g</td>
<td>gram</td>
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<tr>
<td>GFP</td>
<td>green fluorescent protein</td>
</tr>
<tr>
<td>kb</td>
<td>kilo base pair</td>
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<tr>
<td>L</td>
<td>liter</td>
</tr>
<tr>
<td>M</td>
<td>Molar</td>
</tr>
<tr>
<td>min</td>
<td>minute</td>
</tr>
<tr>
<td>ml</td>
<td>mililiter</td>
</tr>
<tr>
<td>n</td>
<td>number</td>
</tr>
<tr>
<td>P value</td>
<td>Probability value</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
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<tr>
<td>rpm</td>
<td>revolutions per minute</td>
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<tr>
<td>Sec</td>
<td>second</td>
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<tr>
<td>T-DNA</td>
<td>transfer DNA</td>
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<tr>
<td>WT</td>
<td>wild type</td>
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<tr>
<td>µl</td>
<td>microliter</td>
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Nomenclature

The wild type genotype is written in italicized capital letters (e.g. FMT).
The mutant genotype is written in italicized lower case letters (e.g. fmt).
The polypeptide products of genes are written in non-italicized, capital letters (e.g. FMT).
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Abstract

Salt stress is known to have severe effects on plant health and fecundity, and mitochondria are known to be an essential part of the plant salt stress response. *Arabidopsis thaliana* serves as an excellent model to study the effects of salt stress as well as mitochondrial morphology. *Arabidopsis* contains several homologues to known mitochondrial proteins, including the fission protein FIS1A, and FMT, a homologue of the CLU subfamily. We sought to examine the effects of salt stress on knockout lines of FIS1A and FMT, as well as a transgenic line overexpressing FMT (*FMT-OE*) in columella cells in the root cap of *Arabidopsis*. *fmt* mutants displayed defects in both root and leaf growth, as well as a delay in flowering time. These mutants also showed a pronounced increase in mitochondrial clustering and number. *FMT-OE* mutants displayed severe defects in germination, including a decrease in total germination, and an increase in the number of days to germination. *fis1A* mutants exhibited shorter roots and slightly shorter leaves, as well as a tendency towards random mitochondrial clustering in root cells. Salt stress was shown to affect various mitochondrial parameters, including an increase in mitochondrial number and clustering, as well as a decrease in mitochondrial area. These results reveal a previously unknown role for FMT in germination and flowering in *Arabidopsis*, as well as insight into the effects of salt stress on mitochondrial morphology. FMT, along with FIS1A, may also help to regulate mitochondrial number and clustering, as well as root and leaf growth, under both control and salt-stressed conditions. This has implications for both FMT and FIS1A in whole-plant morphology as well as the plant salt stress response.
**Zusammenfassung**

regulieren, sowohl unter Kontroll- und Salzstressbedingungen. Dies hat Auswirkungen auf beide FMT und FIS1A in Ganzpflanzenmorphologie und für die Salzstressantwort.
1. Introduction

The driving force behind agriculture is an ever-increasing demand to grow food to sustain a rapidly increasing global population. The pressure to continually produce more crops in a wider variety of environments further drives the need for scientists to design and develop more stress-resistant crop plants. Crops that can endure the effects of exposure to multiple abiotic or biotic stresses while maintaining fecundity will prove to be the most useful for farmers operating in the growing global agricultural market.

1.1 Effects of soil salinity on agriculture

The effects of stress on the proper development and growth of crops currently poses a severe threat to agriculture. One of the most damaging stresses is salinity, or increased levels of salt in the soil. All soil contains some level of salt, and many salts, such as nitrates, are essential plant nutrients. However, the increased use of irrigation and brackish water, as well as increased runoff and poor drainage, has led to high levels of excess salt in the soil. Additional sources of excess salt include inorganic fertilizers, manure, compost, mineral weathering, seawater intrusion into aquifers, and ice melters used on sidewalks and roads (Hasegawa et al., 2000, Carillo et al., 2011). In addition to sodium (Na\(^+\)), irrigation waters may also contain higher levels calcium (Ca\(^{2+}\)) and magnesium (Mg\(^{2+}\)). However, when the water evaporates, the Ca\(^{2+}\) and Mg\(^{2+}\) precipitate into carbonates, leaving behind high levels of Na\(^+\). Soil is considered saline when solution extracted from the soil has an electrical conductivity of 4dS m\(^{-1}\) (decisiemens per meter), where 4dS m\(^{-1}\) ≈ 40mM NaCl or more. In irrigated land, salt levels can vary both
spatially and seasonally, and additional factors such as temperature, pressure, and humidity, can also affect salt levels (Cardon, 2007). It has been estimated that more than 45 million hectares of crops had been damaged by salinization, and 1.5 million hectares were deemed unusable each year (Munns & Tester, 2008). This is predicted to result in up to a 50% reduction in arable land by the year 2050 (Pitman & Läuchli, 2002).

1.2 Effects of soil salinity on plants

Plants can be divided into two major categories for coping with salt tolerance. Glycophytes (salt-intolerant plants) evolved under conditions of low soil salinity and cannot grow or are severely inhibited at salt concentrations above 150mM NaCl. Most glycophytes can tolerate salt concentrations of ~50mM NaCl and below, although some can survive at higher concentrations. Halophytes (salt-tolerant plants) evolved in places with highly salinized soil, and can survive salinity in excess of 300-400mM (Hasegawa et al., 2000).

High salinity affects plants in two main ways: osmotic stress reduces the ability of the plant to extract water from the soil, and high concentrations of salts within the plant can cause damage to plant structures and impede many physiological and biochemical processes. Initial exposure to salt stress has an immediate effect on the plant, rapidly increasing the levels of osmotic and ionic stress. Osmotic stress occurs as a result of excess Na\(^+\) ions in the surrounding soil compared to internal concentrations in the root, which generates an external osmotic pressure that reduces water influx into the root. The result is a water deficit similar to those seen under drought conditions. This can lead to impaired growth and decreased viability, as water and key nutrients are unable to be
transported throughout the plant. Osmotic stress is believed to occur for the duration of salt exposure, resulting in increased stomatal closure and an inhibition of cell division and expansion. Long-term exposure to salt stress can also trigger ionic stress, which occurs as a result of increased Na\(^+\) accumulation in the leaves, which can disrupt protein synthesis and enzymatic activity, often triggering premature senescence in older leaves. This reduces the photosynthetic availability of the plant, further impairing plant growth (Hasegawa et al., 2000, Carillo et al., 2011). However, despite the damaging effects of salinity, plants have evolved a variety of mechanisms to counter-act salt stress over both the short- and long-term.

### 1.3 Plant responses to salinity

Plants display a wide variety of responses to salinity, and as a result exhibit several whole-organism phenotypic changes. Some of these changes are side effects, while some occur as a direct response to salt stress. One example of a side-effect-derived change occurs when excess Na\(^+\) ions are actively shuttled out of the plant shoot and into the leaves, in order to allow for K\(^+\) accumulation in the shoot to help balance the K\(^+\)/Na\(^+\) ion ratio. However, when Na\(^+\) ions reach a critical level in the leaves they begin to stunt growth, eventually leading to leaf necrosis and eventual plant death (Hauser & Horie, 2010). Direct changes in response to salt stress include the reorganization of root system architecture (RSA), which allows for a rapid response to changes in NaCl in the soil (Malekpoor Mansoorkhani et al., 2014, Jones & Ljung, 2012). Other direct responses include a suppression of germination or a delay in flowering, most likely as a way of waiting until conditions are more ideal to grow or produce offspring seeds (Srivastava et
al., 2016, Conti et al., 2008). The genes and pathways that control these phenotypic changes are known in some cases, but the overall etiology underlying the various salt stress responses remain elusive. In addition, the roles of various organelles, including mitochondria, which are known to play a role in the stress response, remain poorly understood. Below is a discussion of some of the most well understood plant responses to stress, including salinity, and the role of mitochondria in each of these responses.

1.4 Mitochondrial response to stress

Mitochondria are known to play key roles in a variety of plant responses to salt stress. These responses usually involve highly conserved mechanisms, and include the hypersensitive response (HR) and programmed cell death (PCD), the SOS Pathway, the reorganization of Root System Architecture and salt-avoidance tropism.

1.4.1 The hypersensitive response and programmed cell death

A common response to stress is the plant hypersensitive response (HR). In response to pathogenic attack, cells will undergo programmed cell death (PCD) in the surrounding area under attack, disabling a virus from co-opting host machinery from neighbouring cells, eventually rendering the virus unable to reproduce, and thus eventually die. PCD is also a well-characterized mechanism in animals, and many of the basic regulatory mechanisms that underlie this response are similar in both plants and animals. Indeed, short-term salinity stress was shown to induce PCD in a manner similarly to animals (Andronis & Roubelakis-Angelakis, 2010). One shared feature between animal and plant PCD is the role of the mitochondria in the regulation of PCD. In both plants and animals,
mitochondria may initiate apoptosis in response to changes in various cellular regulators, such as cytosolic calcium and cellular pH, or indicators of cellular energy availability, such as ATP, ADP, NADH, and NADPH. Various other proteins may also be activated in response to stress and can associate with and modify the permeability of the outer mitochondrial membrane (OMM), including the opening of the mitochondrial permeability transition pore (mPTP). This leads to a decrease in the mitochondrial membrane potential and the release of various cell death activators from within the mitochondrion, including the apoptosis-inducing factor (AIF) and cytochrome c (Lam et al., 2001, Morel & Dangl, 1997, Heath, 2000).

1.4.2 The SOS Pathway

The SOS (Salt Overly Sensitive) pathway was originally identified in a genetic screen to find plants that were hypersensitive to salt stress (Wu et al., 1996, Zhu et al., 1998). The sos1, sos2, and sos3 mutants were shown to have severely impaired growth on media with an excess of Na\(^+\) or Li\(^+\) ions, or a deficit of K\(^+\) ions, but grew similarly to wild type plants under normal growth conditions. These mutants also grew normally under general osmotic or drought stress, which indicates that the SOS genes play a specific role in mediating the ionic response to salt stress in plants. The SOS pathway is activated when excess Na\(^+\) is sensed by the cell, leading to an increase of cytoplasmic Ca\(^{2+}\). This Ca\(^{2+}\) spike is sensed by SOS3, which activates SOS2, forming a SOS3-SOS2 kinase complex. This complex activates SOS1, an Na\(^+\)/H\(^+\) antiporter, which pumps excess Na\(^+\) from the cytoplasm and into extracellular space or the root medium (Ji et al., 2013). Mitochondria are known to buffer cytosolic calcium following a spike in concentration (Vandecasteele
et al., 2001), however, the extent of their role in the SOS pathway has not been well studied.

1.4.3 Reorganization of Root System Architecture and salt-avoidance tropism

The shape and structure of the roots, as well as their spatial configuration within the soil, determines the root system architecture (RSA) of a plant. The RSA of an individual plant is determined by the unique and heterogeneous distribution of edaphic resources (Malamy, 2005, de Dorlodot et al., 2007). In response salt stress, the RSA is altered such that primary root elongation is inhibited, while lateral root (LR) formation increases in response to lower concentrations of NaCl, but is inhibited at higher concentrations (Wang et al., 2008, Zolla et al., 2010). These responses are mediated by changes in cell length and number (Duan et al., 2015), and mitochondria are known to play a role in this regulation (van der Merwe et al., 2009).

In addition to changes in RSA, plant roots may change their direction of growth to avoid excess salt in the soil. Roots primarily grow downwards towards the gravity vector, a phenomenon known as positive gravitropism, or tropic growth. Although it is not known exactly how roots recognize the gravity vector, the “starch statolith hypothesis” posits that amyloplasts in the columella cells of the root cap sediment to the “bottom” of the cell, directing the orientation of growth (Fig. 1A,B) (Sato et al., 2015). When it is necessary to avoid NaCl ions in the soil, roots can exhibit negative gravitropism and grow against the gravity vector, a process known as salt-avoidance tropism, which helps minimize exposure to stress. How the root is able to sense excess salt and subsequently change its direction of growth is not well understood. Sun et al. (2008) found that, upon
exposure of the root to salt stress, two main responses were initiated: 1) rapid degradation of amyloplasts, followed by 2) root bending resulting in negative gravitropism. Amyloplast degradation may be regulated by the SOS pathway, and root bending is likely triggered by PIN2, an auxin efflux carrier, that asymmetrically distributes auxin in the root leading to root curvature (Ottenschlager et al., 2003). However, it is highly likely that other proteins, including those involved with mitochondria, act to regulate salt-avoidance tropism and root bending, as well as reorganization of RSA. Zhang et al. (2015) recently discovered a mitochondrial-localized protein, SSR1, which regulates root growth and architecture, and is required for PIN2 trafficking. This indicates a clear role for mitochondria and their associated proteins in the regulation of root system architecture, with implications for a role in the changes of this architecture in response to salt stress.
**Figure 1: Mechanistic action of tropic growth in the *Arabidopsis thaliana* root.** A) Following a 90° turn, statoliths in columella cells start to fall to the bottom of the cell, and are fully sedimented by 5 min. B) Diagram of an *Arabidopsis* root; columella cells are labelled in green (adapted from Barrada et al. (2015); Sato et al. (2015)). Scale bar = 50 µm.

**1.5 Mitochondrial clustering in response to stress**

A less well-understood response of mitochondria to stress is that of mitochondrial clustering. Mitochondrial trafficking and movement is known to be essential for proper mitochondrial and cellular function, but under stress conditions, mitochondria display altered motility and distribution, which can have deleterious consequences for an organism (Chen & Chan, 2009, Nunnari & Suomalainen, 2012). In plants, mitochondrial clustering and/or arrest of mitochondrial motility has been recognized as a response to various abiotic and biotic stresses, including in response to the application of reactive
oxygen species, heat shock (Scott & Logan, 2008), methyl jasmonate (Zhang & Xing, 2008), oxylipin, 9-hydroxy-10,12,15-octadecatrienoic acid (Vellosillo et al., 2013), and UV light exposure (Gao et al., 2008). However, the mechanisms that give rise to this mitochondrial clustering are not known. Knockouts of the highly conserved gene CLU (CLUstered mitochondria) was shown to induce mitochondrial clustering in a variety of eukaryotes, including Dictyostelium, Saccharomyces cerevisiae, Drosophila, and Arabidopsis (Zhu et al., 1997, Fields et al., 1998, Cox & Spradling, 2009, El Zawily et al., 2014). However, the role of mitochondrial clustering in plants in response to stress, including salt stress, has not previously been investigated.

1.6 The FMT/CLU and FIS1A genes in eukaryotes

The first member of the CLU family to be identified was cluA in Dictyostelium, and this gene was found to be necessary for the correct dispersion of mitochondria within the cell (Zhu et al., 1997). Fields et al. (1998) demonstrated that CLU1, a functional homologue of cluA in Saccharomyces cerevisiae (S. cerevisiae), performed a similar function. clu1Δ cells, which had their CLU1 genes deleted, formed loose clusters of mitochondria within the cytoplasm. Cox and Spradling (2009) characterized the CLU gene in Drosophila, known as clueless, and found that clueless mutants exhibited mitochondrial clustering within cells. Flies that were homozygous for the clu defect (cluD08713 or cluD04554) lived for only 3-7 days, were smaller than WT flies, sterile, and could not fly.

The CLU homologue in Arabidopsis, FMT (friendly mitochondria), was originally identified by Logan et al. (2003) in a mutant screen to find candidate genes involved in mitochondrial dynamics and morphology in higher plants. FMT is 26% identical and
41% similar to the *Dictyostelium* cluA protein and, like cluA, also contains a TPR (tetratricopeptide repeat)-like domain. FMT is 20% identical and 34% similar to the *S. cerevisiae* Clu1p protein. All CLU homologues that have been studied possess a TPR domain, and it remains the most highly conserved portion of the CLU gene throughout its evolution between species. Tetratricopeptide repeats are found in genes in all species, and are known for their ability to mediate protein interactions between partner proteins. In plants, they are found in a variety of genes involved in stress and hormone signalling. One example is *TTL1* (Tetratricopeptide-repeat thioredoxin-like 1), which is known to be a positive regulator of the ABA- (abscisic acid) mediated stress response. Knockouts of *TTL1* increased salt and osmotic sensitivity during seed germination and in later development (Rosado et al., 2006). *Drosophila* clueless was also found to bind nuclear-encoded mitochondrial mRNAs through its TPR domain and direct them to the mitochondrial outer membrane where they could potentially be positioned for co-translational import into mitochondria (Sen et al., 2015).

Electron microscopy analysis of leaf tissue of fmt mutant plants initially revealed the similar phenotype of mitochondrial clustering that was observed in other species (Logan et al., 2003). Further analysis by El Zawily et al. (2014) revealed that FMT might play a role in intermitochondrial association and quality control. It was hypothesized that FMT may function as a fusion protein, as there are currently no known homologues to conserved fusion proteins in plants. However, plant fission proteins are highly conserved in plants, including DRP and FIS1A. Mitochondrial fission proteins also play an important role in the stress response by facilitating the division of a partially damaged
mitochondrion into one healthy and one damaged mitochondrion that can be targeted for degradation (Youle & van der Bliek, 2012).

The mitochondrial division machinery used by *Arabidopsis* is conserved across animals, plants, and fungi (for review see Praefcke & McMahon, 2004). In plants, dynamin-like proteins (DLPs) have been shown to be necessary for the division of mitochondria (Aung & Hu, 2012). DRP3A and DRP3B (previously known as ADL2a and ADL2b, respectively) in *Arabidopsis* are homologous to the Dnm1 and Drp1/DLP1 proteins found in yeast and mammals, respectively. These proteins are part of a DRP subclade that is well conserved across eukaryotic species and contain the GTPase, MD, and GED domains (Miyagishima et al., 2008). DRP3A and DRP3B both localize to mitochondria and were shown to play a dual role in the final scission of both organelles (Aung & Hu, 2012). *Arabidopsis* also contains two proteins homologous to FIS1 in *S. cerevisiae* and humans: FIS1A (also known as BIGYIN1), and FIS1B (also known as BIGYIN2) (Mozdy et al., 2000, Tieu & Nunnari, 2000, Smirnova et al., 2001, James et al., 2003, Youle & Karbowski, 2005). Similar to their yeast and mammalian counterparts, these plant proteins localize to the outer mitochondrial membrane (OMM) and play a key role in mitochondrial division (Logan, 2010). *Arabidopsis fis1A* mutants had a reduction in the number of mitochondria per cell, with simultaneous increases in the size of individual mitochondrion in protoplasts and leaves (Scott et al., 2006). This provides further evidence for the role of FIS1A in mitochondrial fission in plants. However, the role of fission and FIS1A during salt stress is not currently known.
1.7 Thesis aims

The plant salt stress response remains an important mechanism for maintaining growth and survival under ever-changing environmental conditions, and mitochondria are known to be essential for the mediation of this response. However, how this organelle exerts its control, and what proteins are involved, is not well understood. Two mitochondrial proteins, FIS1A, and FMT, are known to be essential for mitochondrial quality control. Given the role of mitochondria in the salt stress response, and given the fact that salt stress is sensed first in the soil by the roots, we wanted to examine the effects of a knockout of either FMT or FIS1A, as well as an overexpression of FMT, in mitochondria in columella cells of the roots under both control and salt-stressed conditions. We also wanted to examine the phenotypic effects of these mutants under both control and salt-stressed conditions. Additionally, since the effects of salt stress on mitochondrial morphology in wild type plants had not previously been characterized, we wanted to examine various mitochondrial parameters in the columella cells of the roots of wild type plants exposed to salt stress. The aim is to further our understanding of the role of mitochondria in salt stress and as such add to the cannon of knowledge of the salt stress response as a whole.
2. Material and Methods

2.1 Material

2.1.1. Antibiotics

Table 1: Antibiotics used in this study

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Solvent</th>
<th>Stock concentration (mg/ml)</th>
<th>Working concentration (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gentamicin</td>
<td>H₂O</td>
<td>10</td>
<td>50</td>
</tr>
<tr>
<td>Kanamycin</td>
<td>H₂O</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Spectinomycin</td>
<td>H₂O</td>
<td>100</td>
<td>75</td>
</tr>
</tbody>
</table>

2.1.2 Bacterial strains

_E. coli_

One Shot TOP10 (Invitrogen, USA)

DH5α (Invitrogen, USA)

_Agrobacterium tumefaciens_

GV3101 (pMP90)

2.1.3 Primers for PCR-based amplification methods

All primers were purchased from the W.M. Keck Foundation (Yale School of Medicine, New Haven, CT). Primer sequences are listed in Table 2.
Table 2: Primers used in this study

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence (5’ → 3’)</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T-DNA Primers</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LBb1.3</td>
<td>ATTTTGCCGATTTTCGGAAC</td>
<td>Left border primer for T-DNA insertion</td>
</tr>
<tr>
<td>FIS1A LP</td>
<td>AAGATCCTCCTTGACACCTCGAC</td>
<td>Left primer for FIS1A (SALK_006512C)</td>
</tr>
<tr>
<td>FIS1A RP</td>
<td>GCTGATTGAGACAAGCTTTTG</td>
<td>Right primer for FIS1A (SALK_006512C)</td>
</tr>
<tr>
<td>FMT LP</td>
<td>ATACCTGCAGCAGTTTGCAAC</td>
<td>Left primer for FMT (SALK_046271C)</td>
</tr>
<tr>
<td>FMT RP</td>
<td>CTAGCGCCAACAGCTCTACTG</td>
<td>Right primer for FMT (SALK_046271C)</td>
</tr>
<tr>
<td><strong>Gateway Primers</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>attB1 FP FMT</td>
<td>GGGGACAAGTTTGTACAAAAAAGCAG GTTCATGGCTGGGAAGTCGAAC</td>
<td>attB1 Forward primer for FMT</td>
</tr>
<tr>
<td>attB1 RP FMT</td>
<td>GGGGACCACTTTGTACAAAGAAAGCTG GGCTTTTTTGGCTTTTTGCTTTT</td>
<td>attB1 Reverse primer for FMT</td>
</tr>
<tr>
<td><strong>Sequencing Primers</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M13 FP</td>
<td>GTAAAACGACGGCCAG</td>
<td>Forward sequencing primer for pDONR 221</td>
</tr>
<tr>
<td>M13 RP</td>
<td>CAGGAAACAGCTATGAC</td>
<td>Reverse sequencing primer for pDONR 221</td>
</tr>
<tr>
<td>FMT Seq1</td>
<td>ATGGCTGGGAAAGTCGAAC</td>
<td>FMT Sequencing Primer 1</td>
</tr>
<tr>
<td>FMT Seq2</td>
<td>ATCTATCAGAGCGCATGTCA</td>
<td>FMT Sequencing Primer 2</td>
</tr>
<tr>
<td>FMT Seq3</td>
<td>GAGCAGAAGAAGCcACTTACCA</td>
<td>FMT Sequencing Primer 3</td>
</tr>
<tr>
<td>FMT Seq4</td>
<td>GCCATAGGGTTTGTGCTCACG</td>
<td>FMT Sequencing Primer 4</td>
</tr>
<tr>
<td>FMT Seq5</td>
<td>AAGAGGAGATAGCTGCTGATG</td>
<td>FMT Sequencing Primer 5</td>
</tr>
<tr>
<td>FMT Seq6</td>
<td>TAATCTTTTGCCAAGGTTGCTG</td>
<td>FMT Sequencing Primer 6</td>
</tr>
<tr>
<td>FMT Seq7</td>
<td>AAAATGAGAGACTTCTTGGGTCTC</td>
<td>FMT Sequencing Primer 7</td>
</tr>
<tr>
<td>FMT Seq8</td>
<td>AACAGAAAACCTGGCTCCTG</td>
<td>FMT Sequencing Primer 8</td>
</tr>
</tbody>
</table>

2.1.4 Cloning vectors

The pDONR 207 (Invitrogen) and pFASTG02 (p*7FWG2, Plant Systems Biology) vectors were used for cloning in this study.
2.1.5 Plant lines

All experiments were performed using Arabidopsis thaliana Col-0 wild type plants or mutants in the Col-0 background. fmt homozygous mutants (SALK_046271C), and fis1A homozygous mutants (SALK_006512C) were obtained from ABRC (Arabidopsis Biological Resource Center, Ohio State University, Columbia, OH, USA). All homozygous mutant lines were confirmed by PCR.

2.1.6 Media, buffers, solutions

Media

LB Media

25g LB Broth

ddH₂O to 1L

For solid medium, 2% Agar was added to the above medium.

After autoclaving at 121°C for 20 mins and cooling to 55°C, antibiotics were added.

MS-Agar Media (pH 5.7)

4.3g MS Salts

0.5g MES

10g Agarose

ddH₂O to 1L
After autoclaving at 121°C for 20 mins and cooling to 55°C, media was poured into plates.

125mM NaCl Stress Media

800 ml MS-Agar Media
7.3g NaCl
ddH$_2$O to 1L

After autoclaving at 121°C for 20 mins and cooling to 55°C, media was poured into plates.

Buffers

CTAB Buffer (100mL, pH 5.0)

2 g CTAB (Hexadecyltrimethylammonium bromide)
10 ml 1 M Tris pH 8.0
4 ml 0.5 M EDTA pH 8.0
28 ml 5 M NaCl
40 ml ddH$_2$O

Phosphate Buffer Stock A

27.6g NaH$_2$PO$_4$·H$_2$O
ddH$_2$O to 1L
Phosphate Buffer Stock B

28.4g/L NaHPO₄·H₂O
ddH₂O to 1L

0.2M Phosphate Buffer (pH 6.8)

51% Phosphate Buffer Stock A
49% Phosphate Buffer Stock B

Solutions

Fixative

0.5% (wt/vol) formaldehyde
3% (wt/vol) gluteraldehyde
0.1M Phosphate Buffer (pH 6.8)

1% osmium tetroxide fixative

1ml OsO₄ (4%)
3ml 0.1M Phosphate Buffer (pH 6.8)

2% uranyl acetate staining solution

0.4g uranyl acetate
20ml H₂O
2.2 Methods

2.2.1 Plant growth conditions and seed sterilization

Arabidopsis seeds were sown directly on Fafard #2 soil mixture (Sun Gro Horticulture) and were grown under 16-hr light/8-hr dark (long-day) photoperiods at 22°C +/-1°C under cool-white light at 100 µmol m⁻² s⁻¹. For experiments done on sterilized MS (Murashige Skoog)-Agar media, seeds were first surface sterilized by washing in 70% (v/v) ethanol for 5 sec, and this wash was replaced by 0.1% triton X-100 in 50% bleach (v/v) for 5 sec before five rinses in autoclaved ddH₂O. Seeds were then plated on 100 x 100 x 15 mm square petri dishes (Ted Pella), and plates were stratified at 4°C for three days in the dark to synchronize germination. Plates were then moved to long-day photoperiod conditions at 22°C +/-1°C under cool-white light at 100 µmol m⁻² s⁻¹. Plates were placed at an angle to allow for root growth along the surface of the agar. For salt-stressed growth conditions on plates, seeds were plated on MS-Agar plates supplemented with 125mM NaCl. For salt-stressed conditions in the soil, the following NaCl concentrations were added when the plants were watered, beginning one week after germination and increasing every week: 50mM NaCl, 75mM NaCl, 100mM NaCl, 125mM NaCl, 140mM NaCl.

2.2.2 Genomic DNA extraction from plant material

Genomic plant DNA was extracted using the CTAB method. 200 mg plant leaf tissue was ground in eppendorf tubes and 500µl CTAB Buffer was added. The mixture was incubated for 15 minutes at 55°C and tubes were then centrifuged at 13,000 rpm for 5
minutes. The supernatant was transferred to a new eppendorf tube and 250 µl of 24:1 chloroform:isopropanol was added and mixed by inversion. The tubes were spun at 13,000 rpm for 1 minute. The upper aqueous phase was transferred to a new eppendorf tube and 50 µl of 7.5M ammonium acetate and 500 µl of ice-cold absolute ethanol were added. The tubes were mixed slowly by inversion and placed at -20°C for 1 hour to precipitate the DNA. Tubes were then spun at 13,000 rpm for 1 minute to form a pellet. The supernatant was removed and the pellets were dried for 15 minutes at room temperature. The pellet was resuspended in 50 µl DNase-free H₂O and stored at 4°C.

2.2.3 PCR reaction

All PCR reactions were done using a Bio Rad C1000 Touch Thermal Cycler. For genotyping, a standard PCR reaction mix was used, using PCR Supermix (Invitrogen). The standard PCR reaction mix (Table 3) and standard PCR thermal profile (Table 4) are shown below.

<table>
<thead>
<tr>
<th>Table 3: Standard PCR reaction mix</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reagent</td>
</tr>
<tr>
<td>PCR Supermix</td>
</tr>
<tr>
<td>Forward Primer</td>
</tr>
<tr>
<td>Reverse Primer</td>
</tr>
<tr>
<td>DNA Template</td>
</tr>
<tr>
<td>H₂O</td>
</tr>
<tr>
<td><strong>25 µl</strong></td>
</tr>
</tbody>
</table>
Table 4: Standard PCR thermal profile

<table>
<thead>
<tr>
<th>Step</th>
<th>Temperature</th>
<th>Time</th>
<th>Cycles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial denaturation</td>
<td>95°C</td>
<td>3 min.</td>
<td></td>
</tr>
<tr>
<td>Denaturation</td>
<td>95°C</td>
<td>30 sec.</td>
<td>20-35</td>
</tr>
<tr>
<td>Annealing</td>
<td>55°C</td>
<td>30 sec.</td>
<td>20-35</td>
</tr>
<tr>
<td>Extension</td>
<td>72°C</td>
<td>20-300 sec.</td>
<td>20-35</td>
</tr>
<tr>
<td>Final extension</td>
<td>72°C</td>
<td>3 min.</td>
<td></td>
</tr>
<tr>
<td>Hold</td>
<td>4°C</td>
<td>∞</td>
<td></td>
</tr>
</tbody>
</table>

2.2.4 Mutant screen

Both FMT and FIS1A were screened for available T-DNA insertion lines on TAIR (The Arabidopsis Information Resource, http://www.arabidopsis.org/). PCR was used to test whether the T-DNA was inserted at the predicted insertion site. All T-DNA insertions were confirmed via PCR using left and right primers flanking the genomic sequence, and a border primer located within the T-DNA sequence (see Table 2 for primer list).

2.2.5 pFASTG02-FMT reporter construct

pFASTG02-FMT was constructed by subcloning a FMT cDNA fragment of the expected size into the pDONR 221 vector (Gateway, Invitrogen). In order to PCR amplify the cDNA fragment, the following primers were used: forward (attB1 FP FMT), 5’-GGGGACAAAGTTTGTACAAAAAAGCAGGCTTCATGGCTGGGAAGTCG AAC-3’, and reverse (attB1 RP FMT), 5’-GGGACCACTTTTGTACAAGAAAGCTGGGTCTTTTTTGCTTTTGGCTTCTT-3’. The fragment was subsequently cloned into pFASTG02 (Shimada et al., 2010) according to the manufacturer’s protocol (Invitrogen). This vector construct, pFASTG02-FMT, allowed for overexpression of the FMT gene under the control of the cauliflower mosaic
virus (CaMV) 35S promoter. The construct was sequenced to identify an error-free clone and subsequently transformed into wild type Col-0 plants by means of *Agrobacterium*-mediated transformation using the *Agrobacterium tumefaciens* strain GV3101 (pMP90).

2.2.6 Sequencing

All sequencing reactions were done by the W.M. Keck Foundation (Yale School of Medicine, New Haven, CT).

2.2.7 *Agrobacterium*-mediated transformation of *Arabidopsis thaliana*

The pFASTG02-FMT vector was transformed into the *Agrobacterium tumefaciens* strain GV3101 via electroporation and colonies were selected on LB media plates supplemented with 50 µg/ml spectinomycin. Single colonies were picked and cultured in LB media supplemented with 50 µg/ml spectinomycin and grown to OD$_{600} = 0.6$. The cultures were then centrifuged at 13,000 rpm for 30 minutes and the pellets were resuspended in a 5% sucrose solution. Plants were dipped according to Clough and Bent (1998) with the following modifications: Silwet L-77 was added to the sucrose solution to a concentration of 0.05% and *Arabidopsis* plants with emerging flower stems were dipped in the solution for five seconds. The plants were then kept under long-day photoperiod conditions under transparent covers at 22°C +/- 1°C under cool-white light at 100 µmol m$^{-2}$ s$^{-1}$ for three days. Covers were removed and plants were grown until seed was mature. Mature seeds were collected and screened to identify transgenic seed expressing the pFASTG02-FMT vector.
2.2.8 Screening and identification of transgenic *Arabidopsis* seed

Seeds from transformed plants were collected and screened for transgenic individuals containing the pFASTG02-FMT vector by the expression of green fluorescence in the seed coat by fluorescence microscopy under 4X magnification with the Zeiss Axioplan 2 fluorescence microscope (Carl-Zeiss, Germany). Transgenic seeds were then sown to produce T₂ seeds. Lines with a single transgene insertion were identified by an ~3:1 segregation ratio of GFP to no-GFP, respectively. Seeds from this line were sown to identify a homozygous plant (FMT-3), which was identified by T₃ seed that exhibited 100% GFP fluorescence. Seeds from line FMT-3 were collected and used in subsequent experiments.

2.2.9 Phenotypic quantification and statistical analysis

*Arabidopsis* plants were grown to three weeks old in the soil under control or salt-stressed conditions described above. For leaf length quantification, three leaves were selected and measured from the base of each leaf to the tip using a ruler. For quantification of root length under control conditions, *Arabidopsis* plants were grown on control MS-Agar plates and roots were measured using a ruler at days seven and fourteen. For quantification of root length under salt-stressed conditions, *Arabidopsis* plants were grown on control MS-Agar plates for one week, and then transplanted to either control or 125mM NaCl MS-Agar plates for one week and roots were measured at day fourteen. For MS-Agar plate experiments, at least 20 replicates were used for each experiment, and each experiment was repeated three times. For soil experiments, at least 10 replicates were used for each experiment, and each experiment was repeated three times. Mutant
genotypes were compared to the wild type under both control and salt-stressed conditions. Statistical differences were determined using Student’s two tailed t test, one-way analysis of variance (ANOVA), or two-way ANOVA, where appropriate.

2.2.10 Transmission electron microscopy

Arabidopsis seedlings were grown on MS-Agar plates for one week, and were then transferred to either 125mM NaCl MS-Agar plates or MS-Agar plates without the addition of NaCl for an additional week. Fixation and embedding of 14-day-old root samples was done according to Wu et al. (2012) with the following modifications: Durcupan epoxy resin (Sigma) was used for infiltration, tissue was collected on single slot copper grids (EMS) coated with formvar, and no post-sectioning heavy metal staining was used. Transverse sections were cut ~30 μM deep into the columella of the root and subsequently viewed under a Tecnai 12 Transmission Electron Microscope (FEI, USA). At least ten cells from four biological samples each of WT, fis1A, fmt, and FMT-OE roots were examined for control experiments, and at least ten cells from two biological samples each of WT, fis1A, or fmt roots were examined for salt-stressed experiments. Due to the difficulty in preserving salt-stressed FMT-OE mutants during the fixation and embedding process, these mutants were not observed for TEM.

2.2.11 Quantification and analysis of Arabidopsis root cells from transmission electron microscopy

Using Fiji, an individual cell, nucleus, vacuole, and mitochondria were traced and measured to give the following parameters: area, aspect ratio (AR), number of
mitochondria per cell, and centroid XY coordinates of each individual mitochondrion. Mitochondrial coverage was calculated as a percent using the following formula: 

\[(\text{Cytoplasmic area} - \text{mitochondrial area}) \times 100\], where cytoplasmic area = (Cell area – nuclear area – vacuole area), and mitochondrial area is the sum of all the areas of the individual mitochondria within the cell.

Mitochondrial clustering was calculated using the Nearest Neighbor Distances (NND) tool in the BioVoxxel toolbox plugin in Fiji (http://imagej.net/BioVoxxel_Toolbox). The NND tool measures the average nearest neighbor ratio (ANN), which is calculated as the distance from the center of a particular particle (in this case a mitochondrion) to the center of its nearest neighbor. The average of all the nearest neighbor distances are then taken. If the average distance is less than the average of a hypothetical random distribution, the mitochondria are considered clustered. If the average distance is greater than a hypothetical random distribution, the mitochondria are considered dispersed. The average nearest neighbor ratio (ANN) is given as:

\[ANN = \frac{\overline{D}_O}{\overline{D}_E}\]

where \(\overline{D}_O\) is the observed mean distance between each feature and its nearest neighbour:

\[\overline{D}_O = \frac{\sum_{i=1}^{n} d_i}{n}\]

and \(\overline{D}_E\) is the expected mean distance for the features given in a random pattern:

\[\overline{D}_E = \frac{0.5}{\sqrt{n/4}}\]
In the above equations, $d_i$ equals the distance between feature $i$ and its nearest neighboring feature, $n$ corresponds to the total number of features, and $A$ is the area of a minimum enclosing rectangle around all features.

The average nearest neighbor $z$-score for the statistic is calculated as:

$$z = \frac{D_o - D_E}{SE}$$

where:

$$SE = \frac{0.26136}{\sqrt{n/A}}$$

If the ANN is less than 1, then the pattern exhibits clustering. If the ANN is greater than 1, the pattern trends towards dispersion. If the ANN is exactly 1, the pattern is considered to be random (Clark & Evans, 1954, Mitchell, 2005). Mitochondria were first analyzed using the Analyze Particles command in Fiji to analyze and measure the individual mitochondria of a single cell. The NND plugin was then used to calculate the ANN of each individual mitochondrion.
3. Results

*Arabidopsis* plants lacking a functional *FMT* gene show severe defects in mitochondrial distribution and movement, as well as deficits in root growth (El Zawily et al., 2014). Plants lacking a functional *FIS1A* gene have a reduction in the number of mitochondria per cell, as well as an increase in the size of individual mitochondria in protoplasts and leaves (Scott et al., 2006). How exactly FMT and FIS1A mediate these changes in mitochondria is still unclear. Given the role of mitochondria in the salt stress response, and given the fact that salt stress is sensed first in the soil by the roots, we wanted to examine the effects of a knockout of either *FMT* or *FIS1A*, as well as an overexpression of *FMT*, in mitochondria in the roots under both control and salt-stressed conditions. Additionally, since the effects of salt stress on mitochondrial morphology in wild type plants has not previously been characterized, we wanted to examine various mitochondrial parameters in the columella cells of the roots of wild type plants exposed to salt stress. These findings will further our understanding of the roles of FMT and FIS1A in mitochondrial morphology, as well as their role(s) in the salt stress response. In addition, it will provide insight into the effects of salt stress on mitochondrial morphology in wild type plants.

3.1 Identification of the *FMT* and *FIS1A* genes

The *FMT* gene was originally identified by Logan et al. (2003) and *fmt* mutants in *Arabidopsis* were shown to have an increased number of clustered mitochondria in the leaves. These mutants were further characterized by El Zawily et al. (2014), and were
found to have shorter roots, as well an increase in the association time between mitochondria, as well as an increase in mitochondrial matrix mixing. The FIS1A gene was originally characterized by Scott et al. (2006), and fis1A mutants in Arabidopsis were found to have a reduced number of mitochondria per cell, but an increase in the size of individual mitochondria in protoplasts and leaves. However, the role of these genes with regards to whole-plant morphology, as well as their role in salt stress, has yet to be explored.

3.1.1 Expression levels of FMT and FIS1A in response to salt stress

Expression levels of FMT and FIS1A under salt stress in the Arabidopsis root were examined using the Arabidopsis Spatio-Temporal Root Stress eFP Browser (http://dinnenylab.info/browser/query). This browser examines the expression levels of ~5 day old seedlings exposed to 140mM NaCl from 1 to 48 hours, compared to exposure on MS-Agar for 1 and 48 hours. A comparison of the expression levels of FIS1A and FMT to expression levels of the Salt Overly Sensitive genes (SOS1, SOS2, and SOS3), which are known to be induced by salt stress, is shown in Figure 2. It is clear that FMT gene expression increases sharply in as little as three hours in the epidermis, with a moderate increase in expression in the stele and cortex from 1-48 hours following NaCl exposure. FIS1A is only mildly upregulated in the epidermis in response to salt stress, similar to SOS1 expression. SOS2 has a moderate decrease in expression in the epidermis, stele, and cortex, while SOS3 is initially highly upregulated from 1-3 hours, with an eventual decrease in expression after 48 hours. While this eFP Browser is informative for short-term exposure at 140mM NaCl, it does not provide information for
long-term exposure to NaCl stress at different concentrations. An in-depth phenotypic and functional analysis of both FMT and FIS1A under different salt-stressed conditions is therefore essential in furthering our understanding of the role of these genes during salt stress.

![Figure 2: Salt-regulated spatio-temporal expression in the Arabidopsis root. A) FIS1A (left) and FMT (right). B) (left to right) SOS1, SOS2, SOS3.](image)

### 3.2 Overexpression of the FMT gene in Arabidopsis thaliana

In order to further our understanding of the role of FMT in whole-plant and mitochondrial morphology, we created a transgenic Arabidopsis plant line (FMT-OE) overexpressing FMT under the control of the cauliflower mosaic virus (CaMV) 35S promoter. Arabidopsis plants were transformed with Agrobacterium containing the pFASTG02-FMT overexpression vector. The pFASTG02 vector carries a screenable marker that produces a GFP signal visible in the mature seed coat of transformed plants. Transgenic seeds from these plants were then sown to obtain T2 seeds. Lines with a single transgene insertion were identified by an ~3:1 segregation ratio of GFP to no-GFP, respectively. Transgenic GFP seeds from these lines were sown to identify a
homozygous plant (FMT-3), which was identified by T₃ seed that exhibited 100% GFP fluorescence compared to WT seed (Fig. 3). Seeds from line FMT-3 were collected and used in subsequent experiments.

![Image of Arabidopsis seeds with GFP fluorescence](image)

**Figure 3**: Seeds of *Arabidopsis* plants transformed with pFASTG02-FMT give green fluorescence.

A) (right) GFP-expressing T₃ generation seeds obtained from FMT-3, a homozygous *Arabidopsis* plant overexpressing the FMT gene in the pFASTG02 vector, (left) non-transformed WT seeds do not give green fluorescence. B) The same field view as in (A), but viewed under bright field light. Scale bar = 100 µm.

### 3.3 Phenotypic analyses of the fis1A, fmt, and FMT-OE mutants

#### 3.3.1 fis1A mutants have shorter roots and slightly shorter leaves

As described above, a homozygous T-DNA insertion line for *FIS1A* (SALK_006512C) was found within the SALK collection. When using primers spanning the insertion site, no transcript could be detected via PCR. The T-DNA insertion hypothetically leads to a block of transcription, rendering a truncated or non-functional FIS1A protein.

Under control conditions, *fis1A* mutant plants did not have a significantly different germination rate compared to WT plants (98.6% ±3.26% versus 99.2% ±1.88%, respectively) (Fig. 4A), nor did they take longer to germinate than WT plants (1.5 days
±0.51 days versus 1.24 days ±0.43 days, respectively) (Fig. 4B). Leaf length was not significantly (p=0.0775) shorter in fis1A mutants compared to the WT (1.03 cm ±0.21 cm versus 1.23 ±0.25 cm, respectively) (Figs. 4C, 5N). These mutants also did not display differences in days to flowering (29.4 days ±1.99 days) compared to the WT (28.89 days ± 1.28 days) (Fig. 4D). However, fis1A mutants did display significantly shorter roots at both 7 (0.59 cm ± 0.17 cm) and 14 (0.85 cm ± 0.43 cm) days old under control conditions compared to the WT (0.86 cm ±0.27 cm and 1.56 cm ±0.48 cm, respectively) (Figs. 4E,F, 5B,F).

3.3.2 fmt mutants have shorter roots and leaves and take longer to flower

As described above, a homozygous T-DNA insertion line for FMT (SALK_046271C) was found within the SALK collection. When using primers spanning the insertion site, no transcript could be detected via PCR. The T-DNA insertion hypothetically leads to a block of transcription, rendering a truncated or non-functional FMT protein. Under control conditions, fmt mutant plants did not have a significantly different germination rate (98.6% ±3.13%) (Fig. 4A) nor did they take longer to germinate (1.33 days ±0.48 days) compared to WT plants (Fig. 4B). However, leaf length was significantly shorter in fmt mutants (0.93 cm ±0.20 cm) (Fig. 4C, 5O), and these mutants also took significantly longer to flower (33.57 days ±2.82 days) compared to the WT (Fig. 4D). fmt mutants also displayed significantly shorter roots at both 7 (0.54 cm ± 0.18 cm) and 14 (0.85 cm ± 0.33 cm) days old under control conditions compared to the WT (Fig. 4E,F, 5C,G).
3.3.3 *FMT-OE* mutants have a lower rate of germination, take longer to germinate, and have shorter roots and leaves

Under control conditions, *FMT-OE* mutant plants had a much lower rate of germination, at only 82.33% ± 9.77% (Fig. 4A). These plants also took longer to germinate (3.12 days ±0.64 days) compared to WT, *fis1A*, and *fmt* plants (Fig. 4B). Despite this delayed germination, these mutants did not take longer to flower (29.83 days ±1.16 days) compared to the WT (Fig. 4D). Similar to *fmt* mutants, leaf length was significantly shorter in *FMT-OE* mutants (0.90 cm ±0.26 cm) versus the WT (Fig. 4C, 5P). Interestingly, *FMT-OE* mutants displayed significantly shorter roots at 7 days old (0.21 cm ± 0.10 cm), but not at 14 days old (1.25 cm ±0.51 cm) under control conditions compared to the WT (Fig. 4E,F, 5D, H).
Figure 4: Phenotypic differences between WT, fis1A, fmt, and FMT-OE lines under control conditions. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 7 days old, F) Root length at 14 days old. Statistical analysis indicates significant differences (****, P ≤ 0.0001, *** P ≤ 0.001, **, P ≤ 0.01, ns = not significant, P > 0.05) compared with controls using one-way ANOVA.
Figure 5: Visualization of phenotypic differences between WT, fis1A fmt, and FMT-OE lines under control and salt-stressed conditions. A-D) Arabidopsis seedlings at 7 days old under control conditions on MS-Agar plates. A) WT, B) fis1A, C) fmt, D) FMT-OE. E-H) Arabidopsis seedlings at 14 days old under control conditions on MS-Agar plates. E) WT, F) fis1A, G) fmt, H) FMT-OE. I-L) Arabidopsis seedlings at 14 days old under salt-stressed conditions on 125mM NaCl MS-Agar plates. I) WT, J) fis1A, K) fmt, L) FMT-OE. M-P) Arabidopsis seedlings at 7 days old under control conditions in the soil. M) WT, N) fis1A, O) fmt, P) FMT-OE. Scale bar = 1 cm.
3.4 Effects of salt stress on fis1A, fmt, and FMT-OE mutants and WT plants

In order to examine the role of the FIS1A and FMT mitochondrial proteins in response to salt stress, fis1A, fmt, and FMT-OE mutants were exposed to either 125 mM NaCl stress on MS-Agar media, or gradual salt stress ranging from 50-140 mM NaCl in the soil. Additionally, to examine the effects of salt stress on WT Arabidopsis plants, and to serve as a control, these plants were also exposed to salt stress on plates and in the soil. On 125 mM NaCl MS-Agar plates, WT plants did not have a significant difference in percent germination (Fig. 6A), but took longer to germinate compared to control conditions (3.64 days ±0.43 days compared to 1.24 days ±0.43 days, respectively) (Fig. 6B). These plants also had significantly shorter roots compared to WT plants under control conditions (0.99 cm ±0.33 cm compared to 1.56 cm ±0.48 cm, respectively) (Fig. 6E, 5I). Under salt-stressed conditions in the soil, WT leaves were significantly shorter compared to the WT control (0.90 cm ±0.15 cm compared to 1.23 cm ±0.25 cm, respectively) (Fig. 6C), and these plants took significantly longer to flower compared to WT plants under control conditions (31.37 days ±2.21 days compared to 28.89 days ±1.28 days, respectively) (Fig. 6D).
Figure 6: Salt stress affects days to germination and flowering, as well as leaf and root length in wild type *Arabidopsis*. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 14 days old. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (***, P ≤ 0.001, ***, P ≤ 0.001) compared with controls using two-tailed Student’s t test.
On 125mM NaCl MS-Agar plates, *fis1A* mutants germinated at relatively the same rate compared to control conditions (96.5% ±4.94% compared to 98.66% ±3.26%, respectively) (Fig. 7A), however these plants took longer to germinate compared to control conditions (3.84 days ±0.80 days compared to 1.5 days ±0.51 days, respectively) (Fig. 7B). These plants also had significantly shorter roots compared to *fis1A* plants under control conditions (0.64 cm ±0.25 cm compared to 0.85 cm ±0.43 cm, respectively) (Fig. 7E, 5J). Under salt stress conditions in the soil, *fis1A* leaves were significantly shorter compared to controls (0.82 cm ±0.13 cm compared to 1.03 cm ±0.21 cm, respectively) (Fig. 7C), and these plants took significantly longer to flower compared to controls (34.31 days ±2.96 days compared to 29.4 days ±1.99 days, respectively) (Fig. 7D).
Figure 7: Salt stress affects days to germination and flowering, as well as leaf and root length in fis1A mutants. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 14 days old. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (****, \( P \leq 0.0001 \), **, \( P \leq 0.01 \)) compared with controls using two-tailed Student’s \( t \) test.
Compared to WT plants under salt-stressed conditions, salt-stressed fis1A mutant plants did not have a significantly different germination rate (Fig. 8A), nor did they take longer to germinate (Fig. 8B). These mutants also did not have significantly shorter roots at 14 days old compared to the WT under the same conditions (Fig. 8E). Under salt-stressed conditions in the soil, the leaf length of fis1A mutants was not significantly shorter compared to WT plants under the same conditions (Fig. 8C), although the number of days to flowering was significantly increased (Fig. 8D).
Figure 8: Days to flowering is increased in fis1A mutants compared to the WT under salt-stressed conditions. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 14 days old. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (**, P ≤ 0.01, ns = not significant, P > 0.05) compared with controls using two-tailed Student’s t test (A, B, C) or two-way ANOVA (C, E).
On 125mM NaCl MS-Agar plates, *fnt* mutants germinated at the same rate compared to control conditions (98.25% ±2.36% compared to 98.6% ±3.13%, respectively) (Fig. 9A), however these plants took much longer to germinate compared to controls (3.84 days ±0.98 days compared to 1.3 days ±0.48 days, respectively) (Fig. 9B). These plants did not have significantly shorter roots compared to *fnt* plants under control conditions (0.75 cm ±0.28 cm compared to 0.85 cm ±0.33 cm, respectively) (Fig. 9E, 7K). Under salt stress conditions in the soil, the leaves of *fnt* mutants were not significantly shorter compared to controls (0.83 cm ±0.16 cm compared to 0.93 cm ±0.20 cm, respectively) (Fig. 9C), nor did these plants take longer to flower compared to controls (33.42 days ±3.24 days compared to 33.57 days ±2.82 days, respectively) (Fig. 9D).
Figure 9: Salt stress affects days to germination in fmt mutants. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 14 days old. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (****, P ≤ 0.0001) compared with controls using two-tailed Student’s t test.
Compared to WT plants under salt-stressed conditions, *fmt* mutant plants did not have a significantly different germination rate, nor did they take longer to germinate (Fig. 10A,B). These mutants also had shorter roots, although it was not statistically significant (P= 0.0605) (Fig. 10E). However, under salt-stressed conditions in the soil, leaves of *fmt* mutants were significantly shorter and days to flowering was significantly longer compared to WT plants under the same conditions (Fig. 10C,D).
Figure 10: Days to flowering is increased and root and leaf length are decreased in fmt mutants compared to the WT under salt-stressed conditions. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 14 days old. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (*, P ≤ 0.05) compared with controls using two-tailed Student’s t test (A, B) or two-way ANOVA (C, D, E).
On 125mM NaCl MS-Agar plates, the percent germination of *FMT-OE* mutants decreased further compared to *FMT-OE* seedlings under control conditions (57.5% ±3.53% compared to 82.33% ± 9.77%, respectively) (Fig. 11A). Days to germination increased significantly compared to controls (4.46 days ±1.12 days compared to 2.71 days ±0.82 days, respectively) (Fig. 11B). These plants also had significantly shorter roots compared to *FMT-OE* plants under control conditions (0.53 cm ±0.11 cm compared to 1.25 cm ±0.51 cm, respectively) (Fig. 11E, 5L). Under salt-stressed conditions in the soil, the leaves of *FMT-OE* mutants were significantly shorter compared to *FMT-OE* plants under control conditions (0.65 cm ±0.14 cm compared to 0.90 cm ±0.26 cm, respectively) (Fig. 11C). However, these plants did not take longer to flower compared to controls (31.9 days ±3.03 days compared to 29.83 days ±1.16 days, respectively) (Fig. 11D).
Figure 11: Salt stress affects percent germination, days to germination, and leaf and root length in *FMT-OE* mutants. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (****, $P \leq 0.0001$, **, $P \leq 0.01$, *, $P \leq 0.05$, ns = not significant, $P > 0.05$) compared with controls using two-tailed Student’s *t* test.
Compared to WT plants under salt-stressed conditions, salt-stressed *FMT-OE* mutants had a significantly decreased germination rate (Fig. 12A), and a difference in days to germination approached significance (p=0.0739) (Fig. 12B). Root length was also significantly shorter (Fig. 12E). Under salt-stressed conditions in the soil, neither the length of the leaves nor the days to flowering were significantly different in *FMT-OE* mutants compared to WT plants under the same conditions (Fig. 12D).
Figure 12: Germination percentage and root length are decreased in FMT-OE mutants compared to the WT under salt-stressed conditions. A) % Germination, B) Days to germination, C) Leaf length, D) Days to flowering, E) Root length at 14 days old. Black bars indicate control conditions, grey bars indicate salt-stressed conditions. Statistical analysis indicates significant differences (**, P ≤ 0.01, *, P ≤ 0.05, ns = not significant, P > 0.05) compared with controls using two-tailed Student’s t test (D,E) or two-way ANOVA (A,B,C).
3.5 Electron microscopy analysis of WT, fis1A, fmt, and FMT-OE plants

Mitochondria are known to play an integral role in the salt stress response. One response to salt stress is salt-avoidance gravitropism, in which amyloplasts from the root cap are degraded, and auxin is asymmetrically distributed via PIN2 transport, in order to induce root bending. The mitochondrial FMT protein is known to be upregulated during salt stress, and fmt mutants were shown to have significantly decreased root cap/meristematic zones in their roots (El Zawily et al., 2014). Thus, an analysis of the mitochondria in the root cap of fmt mutants would provide insight into the role of this gene in root cap morphology and the salt stress response. Furthermore, an analysis of the root cells of another mitochondrial mutant, fis1A, as well as FMT-OE mutants, would provide further insight into the role of mitochondria in the root. Thus, we performed an electron microscopy analysis of mitochondria in columella cells in fis1A, fmt, and FMT-OE mutants and compared these results to WT plants under both control and salt-stressed conditions.

Electron microscopy analysis of the columella cells in the roots of WT, fis1A, fmt, and FMT-OE plants under control conditions revealed major differences in the number of mitochondria per cell, mitochondrial coverage, and mitochondrial clustering (Fig. 13). The wide variation in mitochondrial area (<0.1 µm$^2$ to >0.6 µm$^2$) did not reveal any significant differences between the three mutant plants compared to the WT (Fig. 14A). WT plants had an average mitochondrial area of 0.276 µm$^2$ ±0.15 µm$^2$, fis1A mutants had an average mitochondrial area of 0.279 µm$^2$ ±0.16 µm$^2$, fmt mutants had an average mitochondrial area of 0.279 µm$^2$ ±0.14 µm$^2$, and FMT-OE mutants had an average mitochondrial area of 0.298 µm$^2$ ±0.15 µm$^2$. fis1A mutants did not display a significant
difference in mitochondrial AR (average 1.70 ±0.05), nor did fmt mutant plants (average 1.56 ±0.13), or FMT-OE mutants (average 1.85 ±0.13) compared to WT plants (average 1.71 ±0.05) (Fig. 14B). fmt mutant plants had a significantly increased average number of mitochondria per cell (20.86 ±13.19) compared to the WT (9.21 ± 3.21), fis1A (13.06 ±4.28), and FMT-OE plants (14.23 ±8.15) (Fig. 14C). Both fis1A and fmt plants had increased mitochondrial coverage (15.92% ±8.15% and 16.34% ±9.31%, respectively), compared to the WT (7.36% ±2.33%) and FMT-OE plants (11.13% ±5.49%) (Fig. 14D).
Figure 13: Differences in mitochondria between WT, fis1A, fmt, and FMT-OE lines under control conditions. A) WT, B) fis1A, C) fmt, D) FMT-OE. Each square represents a 4 µm x 4 µm area selected from a representative cell from each genotype. Scale bar = 2 µm.
3.5.1 Salt stress affects various mitochondrial parameters in WT, fis1A, and fmt

Electron microscopy analysis of the columella cells in the roots of WT, fis1A, and fmt plants under salt-stressed conditions revealed major changes in mitochondrial area, number of mitochondria per cell, mitochondrial coverage, and mitochondrial clustering (Figs. 15, 16). Under 125mM NaCl salt-stressed conditions, the average mitochondrial area (0.177 \( \mu m^2 \pm 0.09 \mu m^2 \)) decreased significantly in the WT compared to control conditions (Fig. 17A,B), although AR did not differ significantly (1.56 \( \pm 0.09 \)). Both the average number of mitochondria per cell (39.83 \( \pm 14.82 \)) and mitochondrial coverage
(12.88% ±0.98%), increased significantly (Fig. 17C,D). In fis1A mutants, the average mitochondrial area (0.169 \( \mu m^2 \pm 0.08 \mu m^2 \)) decreased significantly, although AR did not (1.59 ±0.05) (Fig. 18A,B). The average number of mitochondria per cell (24.33 ±12.29) was significantly increased, while mitochondrial coverage was significantly decreased (6.52% ±1.36%) (Fig. 18C,D). Mitochondria in fmt plants were significantly smaller (0.219 \( \mu m^2 \pm 0.07 \mu m^2 \)), however, aspect ratio (1.48 ±0.03), the average number of mitochondria per cell (22.85 ±7.35), and total mitochondrial coverage (17.34% ±6.45%) did not change significantly compared to control conditions (Fig. 19 A,B,C,D).
Figure 15: Differences in mitochondria between WT, fis1A, and fmt lines under salt stress. A) WT. B) fis1A. C) fmt. Each square represents a 4 µm x 4 µm area selected from a representative cell from each genotype. Scale bar = 2 µm.
Figure 16: Visualization of the effects of salt stress on WT, fis1A, and fmt lines. A) WT under control conditions, B) WT under salt-stressed conditions, C) fis1A under control conditions, D) fis1A under salt-stressed conditions, E) fmt under control conditions, F) fmt under salt-stressed conditions. Mitochondria are shown filled in white. Scale bar = 2 μm.
Figure 17: Comparison of various mitochondrial parameters between WT plants under control and salt-stressed conditions. A) Mitochondrial area, B) Mitochondrial aspect ratio, C) # of mitochondria per cell, D) Mitochondrial coverage as a % of cytoplasm. Statistical analysis indicates significant differences (***, P ≤ 0.0001, ***, P ≤ 0.001) compared with controls using one-way ANOVA.
Figure 18: Comparison of various mitochondrial parameters between *fis1A* mutants under control and salt-stressed conditions. A) Mitochondrial area, B) Mitochondrial aspect ratio, C) # of mitochondria per cell, D) Mitochondrial coverage as a % of cytoplasm. Statistical analysis indicates significant differences (****, *P* ≤ 0.0001, **, *P* ≤ 0.01, *, *P* ≤ 0.05) compared with controls using one-way ANOVA.
Figure 19: Comparison of various mitochondrial parameters between fmt mutants under control and salt-stressed conditions. A) Mitochondrial area, B) Mitochondrial aspect ratio, C) # of mitochondria per cell, D) Mitochondrial coverage as a % of cytoplasm. Statistical analysis indicates significant differences (****, P ≤ 0.0001, ns = not significant, P > 0.05) compared with controls using one-way ANOVA.

3.5.2 Clustering is regulated by FMT and is sensitive to salt stress

Mutants with a non-functional CLU gene show a significant increase in mitochondrial clustering in a variety of species, including slime mold, yeast, and fruit flies (Fields et al., 1998, Cox & Spradling, 2009, Gao et al., 2014). In Arabidopsis, it was previously shown via electron microscopy that fmt mutants had increased mitochondrial clustering in their leaves (El Zawily et al., 2014), but an analysis of clustering in the roots of fmt mutants had yet to be done. Additionally, a similar analysis in fis1A and FMT-OE plants, as well as an examination of the effects of clustering under salt-stressed conditions was lacking.
We therefore examined mitochondrial clustering in the roots of WT, fis1A, fmt, and FMT-OE plants under control conditions. Due to difficulties in fixing and embedding salt-stressed FMT-OE roots, we were only able to analyze the roots of WT, fis1A, and fmt plants under salt-stressed conditions. We examined mitochondrial clustering using the average nearest neighbor ratio (ANN) (see Materials and Methods). If the average ANN is less than 1, the mitochondria trend towards clustering. If the ANN is greater than 1, the mitochondria trend towards dispersion. We found that indeed, mitochondrial clustering was evident in fmt mutants (average 0.734 ±0.617) compared to WT plants (1.95 ±1.85). Interestingly, both fis1A and FMT-OE mutants exhibited similar random clustering patterns, with an ANN around 1 (1.08 ±0.75, and 1.06 ±0.752, respectively) (Fig. 20A). An analysis of mitochondrial clustering under salt-stressed conditions revealed that WT plants had a decreased ANN (0.71 ±0.41) compared to WT plants under control conditions, indicating a trend towards mitochondrial clustering under salt stress (Fig. 20B). However, there were no differences in ANN observed in fis1A (0.95 ±0.70) or fmt mutant plants (0.72 ±0.44) (Fig. 20C,D) under salt stress conditions compared to controls.
Figure 20: Clustering patterns in WT, fis1A, fmt, and FMT-OE lines under control and salt-stressed conditions. A) Clustering as measured by Average Nearest Neighbor ratio (ANN), B) ANN comparison between WT under control and salt-stressed conditions, C) ANN comparison between fis1A under control and salt-stressed conditions, D) ANN comparison between fmt under control and salt-stressed conditions. Statistical analysis indicates significant differences (****, P ≤ 0.0001, ns = not significant, P > 0.05) compared with controls using one-way ANOVA.
4. Discussion

Mitochondria are dynamic organelles that orchestrate a variety of cellular functions in all eukaryotic organisms. Although mitochondria from plants and animals have evolved to suit the particular needs of the organism, some mitochondrial genes remain highly conserved, including $CLU/FMT$, and $FIS1A$. The high degree of conservation between these genes across organisms indicates highly conserved and fundamentally important functions. Investigation of these genes in lower organisms such as $Arabidopsis$ can help to serve as a simplified model of gene function, which can help to more quickly elucidate function in higher organisms. The function of $CLU$ remains unknown, and while the basic function of $FIS1A$ in mitochondrial fission is known, it is not well understood how these genes function in response to stress. Thus, in order to gain further insight into the function of these genes and how they regulate mitochondrial and whole-plant morphology in response to stress, we performed phenotypic and cellular analyses of $FMT$ and $FIS1A$ in $Arabidopsis$ under both control and salt-stressed conditions.

4.1 FMT regulates multiple cellular processes in a variety of organisms

$FMT$ in $Arabidopsis$ belongs to the $CLU$ (CLustered mitochondria) family of genes that is highly conserved throughout eukaryotes, including yeast, fruit flies, and humans (Fields et al., 1998, Cox & Spradling, 2009, Gao et al., 2014). Knockouts of this gene consistently display a phenotype of mitochondrial clustering, leading to the hypothesis that this gene was involved in mitochondrial positioning/orientation within the cell. However, recent work by Gao et al. (2014) revealed that $CLU$ may in fact function as an
RNA-binding protein (RBP) that specifically binds mRNAs of nuclear-encoded mitochondrial proteins. CLU may also specifically bind ribosomes on the surface of the mitochondrial outer membrane, in order to help facilitate site-specific mRNA translation (Sen & Cox, 2016). It was also shown that CLU binds TOM20, one component of the TOM mitochondrial import complex that is important for the import of proteins into mitochondria. Additionally, Sen et al. (2015) found that CLU may work to negatively regulate the interaction between PINK1 and PARKIN in Drosophila, which has implications for a role in mitochondrial quality control. In this model, under control conditions, PINK1 is normally imported into the mitochondrial inner membrane where it is degraded by PARL, a rhomboid protease. Under conditions of mitochondrial membrane depolarization, mitochondrial protein import is halted, and as a result PINK1 is unable to enter mitochondria. Instead, PINK1 accumulates on the outer mitochondrial membrane, where it is hypothesized to complex with CLU and well as PARKIN. PARKIN then targets the depolarized mitochondrion for ubiquitination and subsequent degradation. This pathway is an example of a quality control mechanism employed by the cell to prevent unhealthy (i.e. depolarized) mitochondria from accumulating within the cytoplasm. Although Arabidopsis does not possess homologues for PINK1 or PARKIN, it seems likely that a type of mitochondrial quality control might exist within plants, and may be mediated by FMT. Plants also possess a TOM20 complex on the mitochondrial outer membrane, as well as ubiquitinating enzymes (Duncan et al., 2013). It is therefore not unreasonable to hypothesize that plants may also possess a mechanism for mitochondrial quality control that is mediated by FMT functioning as an RBP.
It was also recently found that the homologue of CLU in humans, CLUH, might serve a function in the localization of nuclear proteins. Ando et al. (2016) showed that CLUH can bind viral influenza M1 proteins and transport them into the nucleus, where viral progeny synthesis and subsequent nuclear export occur. This indicates that CLUH is, in certain cases, able to bring proteins into the nucleus in a manner that can alter transcription. All of these experiments indicate that CLU has evolved to regulate multiple cellular processes in a variety of organisms. However, it is currently unclear whether CLU performs a similar function in Arabidopsis, or if it evolved these functions only in higher organisms. Analysis of FMT in Arabidopsis by El Zawily et al. (2014) revealed that fmt mutants had an increased time of association between mitochondria, as well as increase in matrix mixing between mitochondria, and an increase in the number of transient changes in mitochondrial membrane potential. The authors take these data to indicate deficits in mitochondrial fusion, and imply that FMT may function as a mitochondrial fusion protein. As Arabidopsis lacks any homologues to known mitochondrial fusion proteins, it is possible that FMT may serve this function, and in higher organisms this function was eventually relegated to other proteins. Mitochondrial fusion mutants, such as those observed in yeast, worms, and flies (for review see Westermann, 2010), show an increased number of mitochondria, and an increase in numbers of fragmented mitochondria. Although an increase in fragmented mitochondria was not observed in fmt mutants in Arabidopsis, these mutants did show an increase in mitochondrial number. This supports the claim that FMT may function as a fusion protein in Arabidopsis, but more evidence is needed. It is also possible that FMT is not a fusion protein itself, but in fact regulates some as-yet unknown fusion proteins via
mRNA binding. It may also function by negatively regulating the binding of fission proteins to the mitochondrial outer membrane, thereby promoting fusion.

However, considering how similar the phenotypes are between \textit{fmt} mutants in \textit{Arabidopsis} and higher organisms, and the fact that sequence similarity is highly conserved, especially the TPR domain, which was shown to be the RNA binding domain of CLU in \textit{Drosophila}, it seems more likely that FMT and CLU possess similar functions, including RNA binding, and possible tethering to the ribosome and mediation of protein import into mitochondria. Additionally, the role of CLUH in influenza subnuclear transport indicates a role for FMT during stress. Biotic stress of plants, including infection by viruses or bacteria, can induce a variety of responses involving mitochondria (see Introduction). Thus, it is not unreasonable to posit a role for CLU in the biotic stress response in \textit{Arabidopsis}, as well as abiotic stress responses, including salt stress.

\section*{4.2 \textit{fmt} and FMT-OE mutants show deficits in root and leaf length, and delays in flowering and germination}

El Zawily et al. (2014) observed that \textit{fmt} mutants in \textit{Arabidopsis} displayed shortened roots, an observation that was corroborated by our findings. Additionally, we also found that \textit{fmt} mutants had shorter leaves, as well as increase in the number of days to flowering. Overexpression of FMT in \textit{Arabidopsis} resulted in a significant deficit in percent germination and days to germination, as well as a decrease in leaf length. Roots of FMT-OE plants were initially shorter by day 7, but by day 14 had grown to lengths comparable to that of the WT, indicating a key role of FMT in regulating root growth.
The significant delay in germination and flowering in *FMT-OE* and *fmt* mutants, respectively, also indicates a clear role for this gene in both of these processes.

### 4.2.1 Putative role of FMT in germination and flowering control in *Arabidopsis*

In plants, the switch from vegetative to reproductive growth triggers flowering, and several pathways are known to be involved in the initiation and regulation of this transition (for review see Blumel et al., 2015). Several NAC transcription factors have been shown to play a key role in flowering (Sablowski & Meyerowitz, 1998), as well as germination (Kim et al., 2008) and stress (Tran et al., 2004, He et al., 2005, Jiang & Deyholos, 2006) in *Arabidopsis*. NTL8 is a NAC transcription factor belonging to the sub-family of NTLs (*NTM1-Likes*), which are also known to function as membrane-associated transcription factors (MTFs). MTFs remain membrane-bound and dormant until they are activated and transported to the nucleus. This mechanism allows for fast and efficient regulation of gene expression in response to environmental changes. MTFs are known to be activated by regulated ubiquitin (Ub)/proteasome-dependent processing (RUP), in which MTFs are ubiquitinated and partially degraded by the 26S proteasome, and subsequently released in transcriptionally active form (Hoppe et al., 2000). NTL8 is unique among other members of the NTL family in that it is highly induced by elevated levels of salt, and an overexpression of NTL8 delays flowering in *Arabidopsis*. It was also found that NTL8 regulates expression of FLOWERING LOCUS T (FT) in salt-responsive flowering (Kim et al., 2007). Additionally, NTL8 was shown to modulate gibberellic acid (GA)-mediated salt signalling during seed germination (Kim et al., 2008). Interestingly, *fnt* mutants display a similar delay in flowering, although not as severe as
Additional flavonoids. Additionally, *FMT* was shown to be upregulated during salt exposure (Fig. 2). Furthermore, *CLU* was shown to negatively regulate the interaction between PINK1 and PARKIN, (Sen et al., 2015), and PARKIN is known to activate a ubiquitin-proteasome system which targets proteins for degradation by recruitment of the 26S proteasome (Aguileta et al., 2015, Um et al., 2010, Livnat-Levanon & Glickman, 2011).

Additionally, *CLUH* was shown to bind viral M1 proteins and transport them into the nucleus in a manner that can alter transcription. Thus, it is possible that FMT may be able to regulate germination or flowering by mediating RUP and subsequent activation of transcription factors that control these processes.

**4.3 Changes in FMT expression affect mitochondrial size, number, and clustering in columella cells**

In order to gain a more complete understanding of how FMT mediates mitochondrial morphology in the roots of *Arabidopsis*, transmission electron microscopy (TEM) analysis was performed in columella cells. Mitochondria in *fmt* mutants were not significantly different in area compared to WT plants, however they had a significantly increased number of mitochondria per cell, leading to increased mitochondrial coverage. As expected, these cells also exhibited significant mitochondrial clustering, consistent with TEM experiments performed in other species and in the leaves of *fmt Arabidopsis* mutants (El Zawily et al., 2014, Cox & Spradling, 2009). TEM analysis of mitochondrial morphology in *FMT-OE* mutants revealed phenotypes opposite to those observed in *fmt* mutants. *FMT-OE* mutants displayed a random clustering pattern and also had fewer numbers of mitochondria compared to *fmt* mutants, and as a result lower mitochondrial coverage.
coverage, although these values were not significantly different from the WT. These data provide evidence for FMT in the regulation of mitochondria size, number, and positioning of columnella cells in Arabidopsis roots. The absence of a “severe” phenotype in FMT-OE mutants compared to fmt mutants could be explained by the data suggesting that FMT functions as an RBP. It is possible that excess FMT does not have enough ribosomes to bind, and overexpression reaches a certain “threshold” after which point excess FMT has little effect.

4.4 fis1A mutants have shorter roots and may regulate mitochondrial clustering

Mitochondrial fission is essential for cell survival; if a build-up of deleterious components, such as mutated mtDNA, accumulates inside the organelle, it can be segregated to one side and the mitochondrion can divide, leaving one healthy organelle and one bound for autophagy (Youle & van der Bliek, 2012). Mammals and yeast contain only one FIS1 protein, whereas the Arabidopsis FIS1 family (AtFIS1) contains two isoforms, FIS1A and FIS1B, which are known to facilitate the division of both mitochondria and peroxisomes (Zhang & Hu, 2008). It was shown that both fis1A and fis1B single mutants displayed a decrease in the number of peroxisomes and mitochondria. The fis1A/fis1B double mutant showed a similar phenotype to the fis1A single mutant, with an increase in the incidence of clumped mitochondria and peroxisomes, as well as a significant increase in size and number of the two organelles, indicating a deficit in fission. However, the total organelle volume was slightly lower in the double mutants than in the fis1A single mutant, indicating that these two proteins may have overlapping or redundant functions. Overexpression of either FIS1A or FIS1B
increased the abundance of both mitochondria and peroxisomes, but no obvious physiological changes were observed (Zhang & Hu, 2008). While this preliminary research helped to determine that *Arabidopsis* FIS1 proteins have a similar role to their mammalian and yeast counterparts, an examination of these proteins in the roots, as well as under salt-stressed conditions, had yet to be done. Since FIS1A and FIS1B were shown to have partially redundant roles, and since the *fis1A/fis1B* double mutant was unavailable, we decided to examine the *fis1A* mutant in *Arabidopsis* roots, under both control and salt-stressed conditions. The most significant phenotype of *fis1A* mutants was significantly shorter roots, but all other phenotypic characteristics measured did not vary significantly from the WT. Thus, FIS1A may regulate root length, possibly through a control of mitochondrial fission. Interestingly, *fis1A* mutants did not display a decrease in numbers of mitochondria, or an increase in area, as was previously described (Scott et al., 2006, Zhang & Hu, 2008). This can be explained by differences in technique; previous experiments were done in leaves, while we examined mitochondria in the roots. In addition, mitochondria were visualized using a fluorescence-tagged protein under low magnification, while we used high-resolution electron microscopy. As a result, while a cluster of fluorescently tagged mitochondria may appear as a large clump and may be counted as a single mitochondrion, using our technique we were able to discern individual mitochondria within a single cell, and thus gain a more accurate representation of mitochondrial number and size, as well as various other mitochondrial parameters not possible with previous techniques. Although we did not see significant differences in mitochondrial size, AR, or number, we did see an increase in mitochondrial coverage. This could be due to a decrease in overall cell size, and thus a decrease in total
cytoplasm, although this was not quantified. It is important to note that the FIS1B isoform is still functional in fis1A mutants, meaning that mitochondrial fission is not totally abolished. While previous research suggested that fis1A/fis1B double mutants have similar phenotypes to fis1A single mutants, it appears that this is not the case on the level of individual mitochondrial morphology. However, fis1A mutants did display a tendency towards “random” clustering, as opposed to dispersion, as was observed in the WT. Thus, it appears that FIS1A may have a functional role in controlling mitochondrial clustering, likely through control of fission, in columella cells in Arabidopsis roots.

4.5 Salt stress affects various phenotypic and mitochondrial parameters

While it is well established that salt stress has adverse effects in plants, an analysis of the changes to mitochondrial morphology in response to salt stress had yet to be done. Additionally, analyses of various phenotypic and mitochondrial parameters in fmt and fis1A mutants exposed to salt-stress was lacking. Salt stress is first sensed in the root, and salt-stress-coping mechanisms, such as salt-avoidance tropism, are known to involve the root cap. Additionally, mitochondria are known to play a key role in the salt stress response. As such, we performed an analysis of phenotypic and mitochondrial morphology in the Arabidopsis root in WT, fmt, and fis1A mutants under salt-stressed conditions.

Upon exposure to salt stress, WT plants exhibited a decrease in root and leaf length, as well as an increase in days to germination and time to flowering. These same changes were observed in fis1A mutants. fmt mutants exhibited significantly shorter roots compared to WT plants under control conditions. Leaf length was shorter compared to
WT plants under salt stress, but not compared to \textit{fmt} mutants under control conditions. Interestingly, there was no increase in days to flowering, as was seen in WT, \textit{fis1A}, and \textit{FMT-OE} plants under the same salt stress conditions. In contrast, salt stress further exacerbated the decreased germination rate and time to germination of \textit{FMT-OE} plants. Additionally, these plants exhibited an increase in days to flowering and decreases in root and leaf length.

These data indicate that salt stress affects all major phenotypic parameters in \textit{Arabidopsis} wild type plants, although percent germination was unaffected. However, salt inhibition of seed germination (SSG) is a well-known phenomenon (Lee et al., 2010, Yu et al., 2016), although it is usually known to occur under conditions of high salinity (>150mM NaCl). It is therefore likely that we would see an inhibition of germination if our experiments were repeated at a higher level of salt exposure. The exacerbation of the \textit{FMT-OE} germination suppression phenotypes following salt exposure further implicates this gene in regulation of germination, especially during salt stress. Interestingly, despite the fact that \textit{fmt} mutants displayed an increase in days to flowering under control conditions, this phenotype was not exacerbated upon exposure to salt stress. In addition, root and leaf length were not significantly increased in \textit{fmt} mutants under salt-stressed conditions. These data indicate that salt stress-induced changes, such as an increase in days to flowering and a decrease in leaf and root length, as were observed in the WT, are, at least in part, under the control of \textit{FMT}.

An observation of the mitochondria of salt-stressed wild type plants under the electron microscope revealed a decrease in mitochondrial area as well as an increase in mitochondrial number and coverage. A tendency towards clustering was also observed.
*fnt* mutants displayed a decrease in mitochondrial area while AR, mitochondrial number, coverage, and clustering remained unchanged. *fis1A* mutant plants displayed a decrease in mitochondrial area and coverage, with an increase in mitochondrial number, while clustering and AR were not significantly different.

These data further implicate mitochondria in the salt stress response, as it is clear they undergo dynamic changes upon exposure to NaCl. The dynamic increase in number and decrease in size indicates an increased demand for these organelles, possibly through an upregulation of fission. Additionally, these data indicate that mitochondrial number and clustering may be regulated by FMT and FIS1A during NaCl stress.

### 4.6 Conclusion and future perspectives

This study has provided several insights into the roles of the *FMT* and *FIS1A* genes in whole-plant and mitochondrial morphology. It was found that *FMT* may work to regulate root and leaf length, as well as flowering, under both control and salt-stressed conditions. It was also found that *FMT* controls mitochondrial organization via clustering, as well as mitochondrial number under both control and salt-stressed conditions. *FIS1A* was found to potentially regulate root length as well as mitochondrial clustering. Salt stress was also found to affect various mitochondrial parameters, including mitochondrial area, number, and clustering.

More work must be done to fully characterize FMT and FIS1A under both control and salt-stressed conditions. Future research will focus on elucidating the mechanisms of fission in root length control as well as clustering. It appears that CLU may function as an RNA-binding protein in other species, and it should be seen if this is also true for FMT.
in *Arabidopsis*. The role of FMT in salt-avoidance tropism and amyloplast degradation should also be examined. Putative targets of FMT, such as NTL8 and PIN2, and potential ubiquitination of these proteins, should also be explored.
References


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