The functional significance of the alternative first exons of the *Arabidopsis thaliana* APT1 gene

by

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Author's Declaration

I hereby declare that I am the sole author of this thesis. This is a true copy of the thesis, including any required final revisions, as accepted by my examiners.

I understand that my thesis may be made electronically available to the public.
Abstract

In plants, adenine nucleotides are essential for nucleic acid synthesis, as cofactors for biochemical reactions, and as precursors for the biosynthesis of the phytohormone cytokinin. Adenine nucleotides, such as AMP, are synthesized by the de novo synthesis pathway involving 10 sequential enzymatic reactions and 7 ATP/GTP molecules, and by the less energy-demanding salvage reactions. The enzymes catalyzing the latter route collectively make up the purine salvage pathway.

The predominant enzyme in purine salvage is adenine phosphoribosyltransferase (APT), which catalyzes the one-step conversion of adenine to AMP. Plant APTs are also thought to contribute to the metabolism of cytokinins by interconverting the physiologically active base forms into their inactive nucleotide forms. In Arabidopsis thaliana APT is encoded by a five-member gene family, where APT1 contributes 99% of the APT activity. The APT1 gene has a six-exon gene structure that is present in APT genes of other organisms. Recently, expressed sequence tag (EST) collections have shown that the APT1 locus gives rise to a second, longer transcript variant that contains an additional 5′ terminal exon. For the purposes of this research the longer isoform has been designated APT1.1 and the shorter version, APT1.2. The aim of the first part of my thesis research was to determine the functional significance of these two transcript variants.

Semi-quantitative RT-PCR showed that APT1.1 and APT1.2 transcripts are equally abundant in leaf and floral organs. However, immunoblot analysis revealed that APT1.2 is constitutively expressed, while APT1.1 is subject to post-transcriptional regulation in both organs. Using recombinant APT1.1 protein, preliminary enzymatic assays showed this protein to
be very active on adenine substrates. Transgenic lines were constructed that either constitutively expressed the APT1.1 ORF fused to GFP protein, or that only expressed an Exon 1-GFP fusion protein from the native APT1 promoter. In both plant lines GFP localization was observed in stationary punctate bodies in mesophyll cells, whereas constitutive expression of the APT1.1 ORF also showed co-localization of GFP with chlorophyll autofluorescence. Although the identity of these punctate-bodies is currently inconclusive, preliminary data using transmission electron microscopy show that these bodies may be localized in areas of the chloroplast devoid of chlorophyll. Finally, bioinformatic analysis of available plant ESTs predicts that many other plants species may contain a similar gene structure to APT1 of Arabidopsis that gives rise to a cytosolic APT, and a chloroplast-localized APT.

Interestingly, the only phenotype previously identified as arising from APT1-deficiency was aborted pollen development in a point mutant annotated as apt1-3. Recently, a novel APT1-deficient mutant was recovered during a screen for mutants that were tolerant to chemically induced oxidative stress. The mutant, oxt1, contains a T-DNA insertion element between the first and second exon of APT1 and surprisingly is as fertile as WT. Thus, the second part of this research was to determine the basis of fertility in the oxt1 mutant. A genetic cross between these two mutants resulted in an F1 plant that is a developmental chimera of both mutants. The initial siliques on the plant were fertile like oxt1, whereas the later siliques on the plant were sterile like apt1-3. Measurements of adenine and cytokinin levels showed substantial alterations in these metabolite levels. Thus, these metabolites may be potential contributors to the diverse phenotypes observed in each mutant. The discovery and characterization of the unusual phenotype of oxt1apt1-3 hybrids will lead the way in providing a model system to study changes in metabolite levels that contribute to the shift from fertility to sterility.
Acknowledgements

Surprisingly, this section is more difficult to write than any other section of this thesis, because it is hard to find the right words to capture my sincere gratitude.

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A BIG, HUGE thank you to my lab mates (in no particular order): Katja, Sarah, Ishari, Sang, Yong, Makoto, and Terry. You all are truly great friends and have been great to work with, share ideas, and have been there for me when I needed it. A special thanks to Andrew Doxey for all his help with the bioinformatics aspects of this project.

Last, but not least, I want to thank my family especially my parents for their constant support throughout all my years in university (both those that have past, and those yet to come). I don’t want to leave out the help of my brothers, sister-in-law, aunts, uncles, cousins (especially Stella for helping with my references and abbreviations), and my two special nieces that have done everything in their power to distract me when I was trying to focus.
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<th>Description</th>
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<tbody>
<tr>
<td>AAH</td>
<td>allantoate amidohydrolase</td>
</tr>
<tr>
<td>ADK</td>
<td>adenosine kinase</td>
</tr>
<tr>
<td>ADP</td>
<td>adenosine diphosphate</td>
</tr>
<tr>
<td>amiRNA</td>
<td>artificial microRNA</td>
</tr>
<tr>
<td>AMP</td>
<td>adenosine monophosphate</td>
</tr>
<tr>
<td>APT</td>
<td>adenine phosphoribosyltransferase</td>
</tr>
<tr>
<td>AT</td>
<td>3-amino-1,2,4-triazole</td>
</tr>
<tr>
<td>ATP</td>
<td>adenosine triphosphate</td>
</tr>
<tr>
<td>ATase</td>
<td>amidophosphoribosyltransferase</td>
</tr>
<tr>
<td>BA</td>
<td>benzyladenine</td>
</tr>
<tr>
<td>BSA</td>
<td>bovine serum albumin</td>
</tr>
<tr>
<td>BSO</td>
<td>buthionine sulfoximne</td>
</tr>
<tr>
<td>DEPC</td>
<td>dimethylpyrocarbonate</td>
</tr>
<tr>
<td>DTT</td>
<td>dithiothreitol</td>
</tr>
<tr>
<td>EDTA</td>
<td>ethylene diamine tetraacetic acid</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Definition</td>
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<td>--------------</td>
<td>------------</td>
</tr>
<tr>
<td>EMS</td>
<td>ethyl methane sulphonate</td>
</tr>
<tr>
<td>EST</td>
<td>Expressed Sequence Tag</td>
</tr>
<tr>
<td>FSD3</td>
<td>Iron Superoxide Dismutase 3</td>
</tr>
<tr>
<td>GUS</td>
<td>β-glucuronidase</td>
</tr>
<tr>
<td>HEPES</td>
<td>4-(2-hydroxyethyl)-1-piperazinepropane-sulfonic acid monohydrate</td>
</tr>
<tr>
<td>HPLC</td>
<td>high performance liquid chromatography</td>
</tr>
<tr>
<td>IMP</td>
<td>inosine monophosphate</td>
</tr>
<tr>
<td>iP</td>
<td>isopenetyl-adenine</td>
</tr>
<tr>
<td>IPTG</td>
<td>isopropyl-D-thiogalactopyranoside</td>
</tr>
<tr>
<td>K_M</td>
<td>Michaelis-Menten constant</td>
</tr>
<tr>
<td>KOH</td>
<td>potassium hydroxide</td>
</tr>
<tr>
<td>LB</td>
<td>Luria-Bertani</td>
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<td>MEP</td>
<td>methlyerythriol phosphate</td>
</tr>
<tr>
<td>MES</td>
<td>4-morpholineethanesulfonic acid monohydrate</td>
</tr>
<tr>
<td>MS</td>
<td>Murashige and Skoog</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>MVA</td>
<td>mevalonate</td>
</tr>
<tr>
<td>PBS</td>
<td>phosphate buffered saline</td>
</tr>
<tr>
<td>PCD</td>
<td>programmed cell death</td>
</tr>
<tr>
<td>PMSF</td>
<td>phenylmethyl sulfonyl fluoride</td>
</tr>
<tr>
<td>PRPP</td>
<td>phosphoribosyl pyrophosphate</td>
</tr>
<tr>
<td>RT-PCR</td>
<td>reverse transcription Polymerase chain reaction</td>
</tr>
<tr>
<td>PVDF</td>
<td>polyvinylidene fluoride</td>
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<tr>
<td>Q-TOF</td>
<td>quadrupole time of flight</td>
</tr>
<tr>
<td>SDS</td>
<td>sodium dodecyl sulphate</td>
</tr>
<tr>
<td>TAE</td>
<td>tris-acetate-EDTA</td>
</tr>
<tr>
<td>TPP</td>
<td>thiamine pyrophosphate</td>
</tr>
<tr>
<td>TSS</td>
<td>transcription start sites</td>
</tr>
<tr>
<td>tZ</td>
<td>trans-zeatin</td>
</tr>
<tr>
<td>$V_{\text{max}}$</td>
<td>maximal velocity</td>
</tr>
</tbody>
</table>
Introduction

Arabidopsis as a model organism

*Arabidopsis thaliana* is a small, flowering plant belonging to the Brassicaceae family (Figure 1). This plant has become the most widely used model system to study many aspects of plant biology including genetics, molecular and cellular biology, plant anatomy, and plant physiology. Arabidopsis offers many advantages for molecular genetic research because of its completely sequenced diploid genome of 125 Mbp that has a low percentage of repetitive DNA, short exonic and intronic regions, and relatively small intergenic distances indicative of its compact, enriched regulatory regions (The Arabidopsis Genome Initiative, 2000). This small plant is inexpensive to grow and has a short generation time of approximately 6 weeks resulting in seed yields of approximately 10,000 seeds per plant. The ease by which a variety of mutant types can be generated is a particular strength: ethyl methane sulphonate (EMS) is commonly used to create single nucleotide polymorphisms, *Agrobacterium tumefaciens*-mediated T-DNA insertion for gene disruption, or transgenic silencing to reduce the expression of single or multiple genes. Furthermore, protoplast isolation protocols have been developed for transient gene expression studies. These advantages have drawn a large community of scientists to focus their research on Arabidopsis. The research outlined in this thesis examines the functional significance of alternative first exons in an adenine salvage gene involved in purine nucleotide biosynthesis.
Figure 1. 5-week-old wild-type *Arabidopsis thaliana* plant. Image provided by Ishari Waduwara.

**Biosynthesis of purine nucleotides**

Purine nucleotides are essential to all cells as they serve as building blocks for nucleic acid synthesis, provide energy for biochemical reactions, and are precursors for a variety of cofactors (flavin adenine dinucleotide, adenine dinucleotide, and s-adenosylmethionine), B-class
vitamins, and in plants, the phytohormone cytokinin. The adenine nucleotides are of particular importance in that they are the major energy donors for innumerable biochemical reactions. The distribution of adenine nucleotides has been determined in wheat leaf protoplasts where 45% of cellular ATP is localized in chloroplasts with an equal proportion existing in the cytosol and the remaining 5% in the mitochondria (Stitt et al., 1982). The synthesis of adenine nucleotides is most relevant to the research described in this thesis.

**de novo synthesis of adenine nucleotides**

The enzymes involved in the *de novo* synthesis of purine nucleotides were first studied in *Escherichia coli*. Mutant strains deficient in each of the ten enzymatic reactions were recovered and used to identify purine biosynthetic enzymes in other species. The *de novo* synthesis enzymes from plants were identified by functional complementation of *E.coli* mutants (Senecoff and Meagher, 1993; Schnorr et al., 1994).

Purine nucleotides are synthesized from the amino acids glycine or glutamate, phosphoribosyl pyrophosphate (PRPP), and carbon dioxide (Zrenner et al., 2006) via the *de novo* synthesis pathway (Figure 2). This pathway involves a series of ten enzymatic reactions that lead to the synthesis of inosine monophosphate (IMP). Four additional reactions are required to produce ATP, which is then exported to the cytosol via an adenine nucleotide uniporter (Leroch et al., 2005). All the *de novo* enzymes in *E.coli* and plants are monofunctional, with the exception of one bifunctional enzyme in *E.coli*, whereas in most other eukaroytes (e.g. yeast, Drosophila, humans) these enzymes are present as mono-, bi- or trifunctional proteins (Smith and Atkin, 2002).
Figure 2. General schematic of purine nucleotide biosynthesis in plants. Adenine nucleotides are synthesized via two predominant routes, *de novo* synthesis and purine salvage pathways. The *de novo* pathway occurs in plastids and uses 7 ATPs and requires 10 enzymatic reactions to synthesize ATP. The salvage pathway exists in the cytosol and can be subdivided into two pathways. The predominant pathway is the one-step salvage of adenine to AMP; the secondary pathway involves the salvage of adenine to adenosine followed by phosphorylation to AMP. Purine nucleotides are subsequently used as energy donors for biochemical reactions, nucleic acid synthesis, synthesis of guanylate nucleotides, and as precursors for the plant phytohormone cytokinin. Image adapted from Zrenner *et al.* (2006).

Early studies of *de novo* synthesis focused on the subcellular localization of this pathway in root nodules of legume species. Initial studies pointed to plastid localization by the demonstration that PRPP amidotransferase activity (the first enzyme in *de novo* synthesis pathway) exists in plastid fractions of soybean nodules (Boland *et al.*, 1982). To identify if the full complement of salvage enzymes were located within one organelle, cowpea and soybean nodules were subjected to subcellular fractionation on sucrose density gradients (Shelp *et al.*, 1983; Boland and Schubert, 1983). These results showed that plastid fractions fed the purine precursor [14C] glycine produced [14C] IMP. In subsequent years, the subcellular localization of *de novo* synthesis in cowpea nodules was revisited (Atkins *et al.*, 1997). Using subcellular fractionation on sucrose and percoll density gradients, plastid and mitochondrial fractions both
synthesized \([1-^{14}C]\) IMP from \([1-^{14}C]\) glycine. These results provided the first evidence that \textit{de novo} synthesis may also reside in mitochondria of plant cells.

Further evidence to support compartmentalization of the \textit{de novo} synthesis pathway came from the recovery of full-length cDNAs for \textit{de novo} enzymes, coupled with bioinformatic analysis. Smith and Atkins (2002) showed that all ten Arabidopsis \textit{de novo} cDNAs contain N-terminal plastid transit peptides. The sequences of \textit{de novo} purine synthesis enzymes from other non-plant eukaryotic species do not contain apparent N-terminal leader sequences. Although, biochemical and bioinformatic analyses support the plastid localization, only two of the \textit{de novo} purine enzymes have been localized by immunolocalization (Hung \textit{et al.}, 2004; Goggin \textit{et al.}, 2003). Interestingly, the fifth enzyme in the \textit{de novo} pathway was shown to be dual localized to plastids and mitochondria, in cowpea nodules (Smith \textit{et al.}, 1998; Goggin \textit{et al.}, 2003). Future studies will have to experimentally identify the localization of all purine biosynthetic enzymes to verify the extent to which each purine biosynthesis enzyme occurs in each organelle or whether any of these particular enzymes possesses a dual function. Based on these early studies the general agreement is that non-legume plants compartmentalize the \textit{de novo} pathway in plastids, whereas legume species have dual compartmentalization in plastids and mitochondria (Zrenner \textit{et al.}, 2006; Smith and Atkins, 2002).

Consistent with the bioinformatic analyses, non-plant eukaryotic \textit{de novo} purine synthesis enzymes are localized in the cytosol. Gooljarsingh \textit{et al} (2003) monitored the localization of human and \textit{E.coli} GAR transformylase-GFP fusion proteins (the third enzyme in the \textit{de novo} pathway) using confocal microscopy. In both \textit{E.coli} and Cos-7 mammalian cells, GAR transformylase is present throughout the cytosol. A second study focused on the localization of
all six human *de novo* enzymes in HeLa cells (An *et al*., 2008). Interestingly, all six enzymes co-localized with each other in the cytosol forming a “purinosome” complex in response to low purine levels. This co-localization of *de novo* enzymes has yet to be demonstrated for any other organism.

**Synthesis of adenine nucleotides by the salvage pathway**

The synthesis of AMP by *de novo* synthesis is energetically expensive requiring ten enzymatic reactions and 7 ATP/GTP molecules. To overcome this high-energy demand, plant cells actively reuse purine bases produced from the degradation of nucleic acids, turnover of nucleotide cofactors or byproducts of metabolic reactions. The reuse of adenine bases is known as adenine salvage and can be further subdivided into two pathways (Figure 2). The first pathway involves the one-step conversion of adenine to AMP by adenine phosphoribosyltransferase (APT). The second is a two-step pathway in which adenine is converted to adenosine by adenine phosphorylase followed by phosphorylation to AMP by adenosine kinase (ADK).

APT’s prevalence in purine salvage was first shown in *Acer pseudoplatanus* cell cultures where [8-14C] adenine was recovered in the adenylate fraction with very little [8-14C] adenine recovered as adenosine (Doree, 1973). The predominance of the one-step pathway is further supported in that APT and ADK activity are high in plants, yet adenine phosphorylase activity is low (Lee and Moffatt 1994; Moffatt *et al*., 2000; Ross, 1981). This evidence suggests that APT catalyzes the major route for adenine salvage, where ADK catalyzes the major route for adenosine salvage.

Purine salvage enzymes in plants contribute to both purine and cytokinin metabolism. The role of these enzymes is two-fold. They function in reducing intracellular accumulation of
adenine as well as the more obvious role in synthesizing adenine nucleotides. Much of the research on these enzymes has been focused on the importance of purine salvage activities in the maintenance of nucleotide pools during times of high demand such as seed germination or programmed cell death (Ashihara 1983; Stasolla et al., 2005). In these and other tracer experiments the majority of [8-14C] adenine was recovered in nucleotide fractions.

Adenine salvage enzymes are also thought to contribute to cytokinin interconversion. Cytokinins are plant growth regulators implicated in many aspects of growth and development (reviewed in Hirose et al., 2008). Structurally, cytokinins consist of an adenine moiety with either an isoprenoid or aromatic side chain conjugated at the N6 position. Each of these two side chains further contains a number of derivatives adding to the diversity and modifying the functionality of these phytohormones. In Arabidopsis, the isoprenoid cytokinins are the major forms, in particular isopenetyl-adenine (iP) and its hydroxylated derivative trans-zeatin (tZ) (Sakakibara, 2006).

Cytokinins are further interconverted between bases, riboside and nucleotide forms. Although, only one enzyme in rice unequivocally catalyzes the conversion of the cytokinin nucleotide to its base form, there is growing evidence that purine salvage enzymes catalyze these interconversion reactions (Kurakawa et al., 2007). These cytokinin isoforms possess different binding affinities for cytokinin receptors indicating that the bases are the most physiologically active, with some residual activity in nucleosides, and very low levels of activity for nucleotides (Spichal et al., 2004). Therefore, APT and ADK activities theoretically contribute to reducing the abundance of active cytokinin bases and ribosides to the less active nucleotide form. In fact, it has been shown that APT purified to various degrees of homogeneity from wheat, peach, and
Arabidopsis can catalyze the interconversion of cytokinin bases to nucleotides (Chen et al., 1981; Lecomte and Le Floc’h, 1998; Moffatt et al., 1991). Interestingly, APT and ADK may also contribute to maintaining active cytokinin levels. As both purine enzymes synthesize AMP, the phosphorylated derivatives of this adenylate, may contribute the necessary precursors (ADP/ATP) for cytokinin biosynthesis (Kakimoto, 2001). The relative contributions of each purine salvage enzyme to cytokinin metabolism remains to be determined.

Purine salvage enzymes have long been assumed to reside in the cytosol. This localization is supported by a molecular analysis of these enzymes in Arabidopsis and the absence of apparent N-terminal leader sequences in recovered full-length cDNAs (Allen et al., 2002, Moffatt et al., 1994; Moffatt et al., 2000). To date, only two groups have localized purine salvage enzymatic activity outside the cytosol, and both instances are in plant species. Ashihara and Ukaji (1985) used subcellular fractionation to purify chloroplasts and mitochondria from spinach leaves. They showed that both fractions were able to catalyze the synthesis of [8-14C]-AMP from [8-14C] adenine and [8-14C] adenosine. A similar approach was used to localize APT activity in Jerusalem artichokes to mitochondria (Le Floc’h and Lafleuriel, 1983). No further studies have since shown non-cytosolic protein localization of these enzymes.

**Adenine salvage by APT in Arabidopsis**

Adenine salvage has been studied extensively in Arabidopsis since Moffatt and Somerville (1988) recovered the first APT-deficient plants based on their resistance to 2,6-diaminopurine. They isolated three point mutants, apt1-1, apt1-2, and apt1-3, which had 15%, 2%, and 1% residual APT activity, respectively. Surprisingly, the fertility of these plants correlates with the decrease in APT activity, where apt1-1 has 2% of the seed production of wild
type and *apt1*-2 and *apt1*-3 are male sterile (Moffatt and Somerville, 1988). The male sterility in *apt1*-3 is due to aborted pollen development shortly after meiosis (Moffatt and Somerville, 1988; Regan and Moffatt, 1990; Zhang *et al.*, 2002). Three possible explanations were proposed for why APT deficiency leads to male sterility: insufficient ATP during microsporogenesis, accumulation of adenine leading to inhibition of methylthioadenosine nucleosidase or altered cytokinin interconversion during pollen development.

The effect of APT activity on cytokinin metabolism was tested in the most severe APT1-deficient mutant, *apt1*-3. Four-day old seedlings of WT and *apt1*-3 were grown in liquid medium and fed the cytokinin benzyladenine [³H]-BA. *apt1*-3 plantlets metabolize 8-fold less BA than did WT plantlets (Moffatt *et al.*, 1991). The apparent Kₘ of APT purified from Arabidopsis leaves was 4.5 μM for adenine versus 730 μM for the benzyladenine (BA) cytokinin, revealing its stronger binding affinity for adenine as compared to this particular cytokinin (Lee and Moffatt, 1993). Since this was above the physiological concentrations of cytokinins, it was difficult to conclude the extent to which APT acts on cytokinins *in vivo*. Despite conclusive evidence to support the role of reduced cytokinin or adenylate metabolism in male sterility, subsequent research focused on the *APTI* locus to gain further insight on its function in pollen development.

The full-length cDNA of *APT* was first recovered by Moffatt *et al* in 1992. Alignment of its predicted amino acid sequence with those of APTs from *E.coli*, Drosophila, hamster, human, and mouse showed these proteins to be highly conserved in both in their size and amino acid sequence. A promoter fusion with the β-glucuronidase-encoding (GUS) gene using the 950 base pair region upstream of the predicted transcriptional start site indicated that APT1 was expressed
constitutively in Arabidopsis plants (Moffatt et al., 1994). The constitutive expression of the APT promoter was consistent with previous APT enzymatic assays that showed APT activity in leaves, flowers, stems, and buds (Lee and Moffatt, 1993). When expressed on a protein basis differences in expression are seen: relative to leaves APT activity is higher in buds (2.7-fold), stems (2.2-fold), and roots (3.9-fold).

With the increase in Expressed Sequence Tag (EST) collections and the availability of the complete Arabidopsis genome sequence, four additional APT genes were identified (APT2, APT3, APT4, and APT5). The amino acid sequence similarity among the five APTs is 89% (Allen et al., 2002). It is likely that these APT genes arose from genome duplication events as described by Blanc et al (2000). To date, APT4 and APT5 have not been investigated due to the minimal expression data available for these genes. As such, only the APT1, APT2, and APT3 isoforms will be discussed further.

To explore the kinetic properties of these APT isoforms Schnorr et al. (1996) recombinantly expressed APT1 and APT2 in E.coli. Based on their analysis APT1 and APT2 have similar apparent $K_M$ values for adenine, while APT2 has a $K_M$ 167-fold lower than APT1 for the cytokinin BA; the catalytic efficiency of APT1 is 180-fold higher than that of APT2. When BA is the substrate, APT2 was 250-fold more efficient than APT1. These results pointed towards a role of APT1 in adenine metabolism, and APT2 in the metabolism of cytokinins.

Allen et al. (2002) further compared the kinetic activities of these enzymes, this time including APT3. APT1, APT2, and APT3 have similar adenine $K_M$ values but APT1 had a 31-fold higher adenine $V_{max}$ than APT2, and 53-fold higher adenine $V_{max}$ than APT3. This evidence
supports the model that APT1 is the predominant isoform acting on adenine in Arabidopsis; APT2 and APT3 had greater activity on cytokinins than did APT1.

The predominance of APT1 in adenine metabolism is consistent with the male sterility observed in the most APT-deficient mutant, apt1-3, that contains a point mutation at the splice donor site downstream of Exon 4 in APT1 (Gaillard et al., 1998). Although the plant has two other functional APTs, APT2 and APT3, these isoforms are clearly not able to compensate for the loss of APT1 activity in apt1-3.

The subcellular localization of these three isoforms was also determined by Allen et al. (2002). Based on subcellular fractionation and immunoblotting analysis with isoform-specific peptide antibodies, APT1 and APT3 were localized to the cytosol; the results for APT2 were less conclusive but is likely cytosolic since it lacks an N-terminal transit peptide for chloroplast import.

In situ hybridization was used to address why the other APTs did not complement APT1 deficiency in pollen (Zhang, 2000). Consistent with previous data, APT1 was constitutively expressed with the highest level of expression in young buds. APT2 showed the strongest transcript levels in floral receptacles although the amount of signal was near background levels. This result is consistent with that obtained by Schnorr et al. (1996) who found that APT2 could only be detected by primer extension assay in floral organs but not in leaves. APT3 transcripts were highest in ovaries. Interestingly, the transcript levels of all three APTs increase in apt1-3 mutants flowers; the molecular mechanism of this increase remains unknown (Zhang, 2000).

Thus, based on these studies, APT1 appears to be the only isoform expressed in pollen. The timing of pollen abortion correlates with APT1’s expression in both tapetal cells and
microsporocytes (Smith, 1997). Before meiosis APT1 expression increases in both these cell types, but immediately decreases soon thereafter. This correlation provides strong support for the importance of APT1 activity in pollen development.

**Additional gene models for Arabidopsis APT1**

Transcriptional profiling of Arabidopsis organs by EST analysis has revealed that a second, longer APT1 mRNA arises from this locus (Figure 3). For the purpose of discussion, these two mRNAs and their corresponding protein products have been named APT1.1 to identify the longer isoform, and APT1.2 to identify the shorter isoform; APT1.2 is the form previously isolated by Moffatt et al (1994). When APT1.2 was first reported it was shown to have a six-exon gene structure with a high sequence similarity to all other known APTs. The APT1.1 mRNA differs from APT1.2 by the presence of an additional 5' terminal exon. This phenomenon of two transcripts arising from the same gene, yet containing different initiating exons is referred to as alternative first exons. For the purposes of clarity in discussing the alternative first exons of this gene, the first exon of APT1.1 is designated as Exon 1 and the first exon of APT1.2 is Exon 2. As seen in Figure 3, Exons 2 through 7 are common to both gene models and transcripts.
Figure 3. Predicted gene models of APT1.1 and APT1.2 based on EST collections. Two EST clusters are shown in green. The predicted gene models derived from each EST cluster is shown at the top with the blue boxes representing the exons. APT1.1 has a seven-exon gene structure; APT1.2 has a six-exon gene structure as identified by Moffatt et al (1992). The ESTs show that APT1.1 and APT1.2 have unique 5’UTR regions where that of APT1.2 is located in the first intron. Image is modified from The Arabidopsis Information Resource website (http://www.arabidopsis.org).

Aside from their unique first exons, a second distinguishing feature between the APT1.1 and APT1.2 mRNAs is that each has a unique 5’ UTR region. As seen in Figure 4, the 5’ UTR of APT1.2 resides in the first Intron where the 5’ UTR of APT1.1 is upstream of the methionine codon of Exon 1.
Figure 4. Partial genomic sequence of the *APT1* gene. Two mRNAs arise from the *APT1* gene in Arabidopsis. The *APT1.1* primary transcript initiates at the first red arrow and is spliced at the splice acceptor and splice donor sites underlined in blue, to remove the first Intron. The translation of this mRNA begins at Exon 1 (highlighted in yellow) and the resulting polypeptide includes the 6 amino acids adjacent to Exon 2 (highlighted in red). The *APT1.2* primary transcript initiates at the second red arrow with part of Intron 1 forming the 5’UTR. The translation of this mRNA initiates at Exon 2. Note that the APT1.2 polypeptide does not contain the 6 amino acids adjacent to Exon 2; these are only present in an APT1.1 polypeptide.
Interestingly, the distance between the transcriptional start sites (TSS) of \textit{APT1.1} and \textit{APT1.2} is approximately 478 base pairs. Bioinformatic analysis in humans shows that alternative first exons with TSS greater than 500 base pairs apart are unlikely to have one promoter direct the expression of both mRNAs (Kimura \textit{et al.}, 2006). This conclusion is consistent with the results of studies of the expression of alternative first exons in genes from humans (Kimura \textit{et al.}, 2006), rice, and Arabidopsis (Chen \textit{et al.}, 2007). Both studies found that the shorter mRNA is normally constitutively expressed, whereas the longer mRNA shows a spatial/temporal-specific expression in 52% of the human cases, 20-66% of the rice cases, and 5-18% of the Arabidopsis cases. The most plausible explanation for this occurrence is that the two transcripts are produced from two promoters. This would predict that \textit{APT1.2} gene is constitutively expressed and \textit{APT1.1} might exhibit spatial/temporal-specific expression. If this is true, the \textit{APT1.1} promoter is presumed to be located upstream of its transcription start site (Figure 4), whereas the \textit{APT1.2} promoter resides in the first intron and potentially further upstream.

The recovery of the \textit{APT1.1} cDNA from both the RIKEN and CERES full-length cDNA clone collections indicates this is a functional mRNA. Both of these cDNA libraries were created by selecting for transcripts containing a 5´-guanidine cap, a hallmark of the 5´ end of mRNAs. This evidence initiated part of the research outlined in thesis: to identify if \textit{APT1.1} produces a polypeptide and if so, what is its functional significance.

Chen \textit{et al} (2007) highlight various alternative first exon genes in rice and Arabidopsis documenting the various roles of the first exon including being a transit peptide, adding or altering enzymatic activity, or controlling the spatial/developmental/temporal expression of the associated gene. Results of various subcellular localization prediction programs differ in their
proposed localization of APT1.1, although a chloroplast localization appears more frequently than others (Table 1). All the programs predict APT1.2 to be situated in the cytosol. As for differences in enzymatic activities, this cannot be predicted using bioinformatics thus this role for Exon 1 must be determined experimentally. Finally, the tissue sources for the ESTs were investigated to see if there was a bias in the tissues each mRNA was recovered from. Of the eighteen APT1.1 ESTs, ten were obtained from floral organs, and the remaining were obtained from leaf tissue of young plantlets.
Table 1. Predictions of the subcellular localization of APT1.1 using bioinformatic tools. MitoProtII and TargetP are the only programs to predict both length and cleavage site of the proposed transit peptide (57 amino acids predicted by MitoProtII and 52 amino acids by TargetP). Date of access August 2009.

<table>
<thead>
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<td>Hoeglund et al., 2006</td>
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</table>

**Novel apt1-deficient mutant is not male sterile**

Concurrent with the discovery of the APT1.1 isoform a novel APT1-deficient mutant was identified. This mutant was recovered by D. Falcone at the University of Massachusetts and named *oxidative stress tolerant 1 (oxt1)*. The oxt1 mutant arose from a population of T-DNA mutagenized seeds that were plated on solid plant growth media supplemented with 3-amino-
1,2,4-triazole (AT) and buthionine sulfoximine (BSO) (Zhang et al., 2008). These compounds work synergistically to induce oxidative stress by inhibiting catalase activity and suppressing glutathione biosynthesis, respectively (Zhang et al., 2008). WT seedlings exposed to these conditions initially show inhibited root growth; prolonged exposure results in chlorosis and death. Under these conditions oxt1 exhibited 75% increased tolerance relative to WT (i.e. it shows symptoms on 75% higher concentration).

oxt1 was also tested for increased tolerance to other abiotic stresses such as temperature and drought. When oxt1 mutants were grown at 28°C their dry mass was 1.5-fold greater than WT, indicating a higher rate of growth in these mutants. Moreover, oxt1 mutants grown at 38°C for 5-days showed less anthocyanin production relative to WT plants. Increased drought resistance in WT and oxt1 was tested by withholding water from 3-week-old plants. After 2 weeks 50% of WT plants died while all oxt1 mutants survived (Sukrong et al., manuscript in preparation).

Since oxt1 exhibited greater stress tolerance, RT-PCR analysis was used to investigate the expression of select antioxidant defense genes. oxt1 mutants have a higher expression of antioxidant defense genes in both non-stress and stress-induced conditions. During non-stress conditions three genes are up-regulated relative to WT: stromal ascorbate peroxidase, thylakoid ascorbate peroxidase, and catalase 3. Under stress-conditions these genes were further upregulated above the levels found in WT (Sukrong et al., manuscript in preparation).

Subsequent map-based cloning revealed that oxt1 carries an uncharacterized T-DNA insertion immediately upstream of Exon 2 in APT1 (D. Falcone, personal communication). APT activity was undetectable in oxt1 based on a spectrophotometric assay (D. Falcone, personal
communication). Despite their reduced APT activity, oxt1 is completely fertile, unlike all previously characterized APT mutants. Dr. Falcone contacted Dr. Moffatt about collaborating on the analysis of the oxt1 mutant’s APT activity and stress tolerance.

**Objectives of this Research**

The research outlined in this thesis has three principal objectives:

1. The only evidence for the existence of APT1.1 is based on EST collections. This research set out to initially address the following questions: Is the APT1.1 transcript a functional mRNA; how does APT1.1 transcript abundance compare to that of APT1.2, and does APT1.1 give rise to a polypeptide?

2. The second objectives are to investigate whether APT1.1 is a functional enzyme and if so, to determine the functional significance of Exon 1 in APT1.1.

3. The oxt1 mutant is APT1-deficient yet it is not male sterile. I did comparative analyses of oxt1 and apt1-3 to look for an explanation for their differing phenotypes.
Materials and Methods

Source of Chemicals and Reagents

All chemicals and reagents used were purchased from Sigma-Aldrich, Bioshop, or BioBasic, unless otherwise noted. Restriction enzymes were purchased from New England Biolabs or Fermentas.

Seed germination and plant growth

Seeds were sterilized in a 1.5 mL microfuge tube containing a maximum of 100 mg of seeds. The 1.5 mL microfuge tube was placed in a glass dessicator beside a 400 mL beaker containing 100 mL of Javex and 3 mL of concentrated HCl. The seeds were sterilized by chlorine gas for 2 hours. Sterilized seeds were stored afterward at room temperature away from direct light.

Sterilized seeds were grown on Murashige and Skoog (MS) solid medium in Petri plates. The medium was prepared using MS Salts and Gamborg’s vitamins (Sigma), 2.56 mM 4-morpholine-ethanesulfonic acid monohydrate, 30 g/L sucrose, 0.8 g/L agar; the pH was adjusted to 5.8 using 1M potassium hydroxide (KOH). After stratification for 48 hours at 4°C in the dark, seeds were transferred to a 21°C growth chamber under 50 µEinsteins m⁻² s⁻¹ continuous fluorescent light. After 10-14 days the seedlings were transplanted into 4-inch deep plastic pots containing a 50:50 mix of “Sunshine LC1 mix” and “Sunshine LG3 Germination mix” soil (JVK, St Catharines, ON). Potted plants were grown in a 21°C growth chamber under 16-hour days and fluorescent lighting providing 120 µEinsteins m⁻² s⁻¹ fluorescent light.
Radiochemical assay of Adenine phosphoribosyltransferase enzyme activity

This assay was performed as described by Allen et al (2002). Crude protein extracts used for this assay were prepared on ice by grinding leaf tissue (50-100 mg) or flower tissue (30-70 mg) in 50 mM Tris-HCl buffer pH 8.8 at a ratio of 3 µL of buffer to 1 mg of tissue in glass homogenizers. The extracts were centrifuged at 13,200 rpm (18,500 x g) for 3 minutes to remove insoluble material and debris. The supernatant was assayed for enzyme activity. The general protein concentration recovered was 1-2 mg/mL for leaf extracts, 2-3 mg/mL for flower extracts. Note that desalting the protein extract through Sephadex G-25 beads (GE Healthcare) results in a 50% decrease in APT activity, and is not recommended.

The assay which measures the conversion of $^3$H-adenine to $^3$H-AMP was carried out in a final reaction volume of 100 µL containing 50 mM Tris-HCl pH 8.8, 15 mM NaF, 2.4 mg/mL bovine serum albumin (BSA), 10 mM NaN$_3$, 5 mM MgCl$_2$, 19 µM 5-phosphorylribose 1-pyrophosphate (PRPP), and 100 µM [2,8-$^3$H] adenine (40 Ci mmol$^{-1}$; MP Biomedicals Inc.). The 100 µM of adenine consisted of 0.2 µL of [2,8-$^3$H] adenine (initial concentration 25 µM) and 12.2 µL of 819 µM non-radioactive adenine resulting in 10 nmol of adenine per reaction. The PRPP had to be prepared fresh; all the other reagents were used from frozen stocks stored at −20°C.

The reaction mixture was pre-warmed for 5 minutes at 37°C before plant crude protein extract was added and allowed to incubate at 37°C for 5 minutes. The reaction was stopped by adding 1 mL of ice-cold stop buffer (0.05 M NaOAc pH 5.0, 2.0 mM K$_2$HPO$_4$) followed by precipitation of $^3$H-AMP with 200 µL of 0.5 M LaCl$_3$. Nucleotides were precipitated at 4°C for a minimum of 1 hour before being collected on a glass-fiber filter with pore size of 1.2 µM.
(Fisherbrand, Cat.No. 09-804-24C), by vacuum filtration. The filter was placed in a scintillation vial containing 4 mL of Cytoscint (MP Biomedicals Inc.) and incubated for at least 1 hour before liquid scintillation counting (Model LS 1701, Beckman). Each sample was assayed in triplicate and in triplicate volumes (i.e. 2, 4, and 6 µL) to ensure the linearity of the assay. Activity was expressed as nmol of AMP mg of protein$^{-1}$ min$^{-1}$. It should be noted that the assays performed in this thesis research were not linear at the highest volume. This issue was resolved for a short time by using a newly prepared stock of NaF, but later became an issue once again.

**ADK Enzyme Assay**

This assay was performed as described by Moffatt *et al.* (2000). Plant crude protein extracts for this assay were prepared on ice by grinding leaf tissue (80-100 mg) in 100 mM HEPES buffer pH 7.2 at a ratio of 3 µL of buffer to 1 mg of tissue. The extracts were centrifuged at 13,200 rpm for 2 minutes. The supernatant was then desalted through Sephadex G-25 beads (GE Healthcare) that had been resuspended in 50 mM HEPES pH 7.2 (1 gm of beads per 5 mL of buffer). The concentration of the crude protein extract was generally 1.0-1.5 mg/mL as measured using Bio-Rad protein assay reagent (Bio-Rad). This extract was subsequently diluted to 0.1 mg/mL. Extracts of ADK-deficient mutants were not diluted.

This assay measures the conversion of $^3$H-adenosine to $^3$H-AMP. It was carried out in a final reaction volume of 50 µL containing 50 mM HEPES buffer pH 7.2, 4 mM ATP, 1 mM MgCl$_2$, 30 mM NaF, 2.5 µM deoxycoformycin (Warner Lambert), 1 mg/mL BSA, and 2 µM [2,8-$^3$H] adenosine (31.4 Ci mmol$^{-1}$; Sigma). The samples were processed as described for the APT assay with the exception that the reactions were carried out at 30°C. Activity was expressed as nmol of AMP mg of protein$^{-1}$ min$^{-1}$. 
RNA isolation, cDNA synthesis, and RT-PCR of *APT1, APT2, and APT3*

Plant tissue was homogenized using a mortar and pestle containing liquid nitrogen and quickly transferred to 1 mL TriPure Isolation Reagent (Roche Applied Science) in a 1.5 ml microfuge tube. After a five min incubation at room temperature, 200 µL of chloroform was added, the mixture was vortexed for 15 seconds, incubated for 15 minutes at room temperature and centrifuged at 13,200 rpm for 15 minutes at 4°C. To the aqueous phase an equal volume of isopropanol was added; the mixture was vortexed thoroughly and centrifuged at 13,200 rpm (18,000 rcf) for 15 minutes. The pellet was then washed with 1 mL of 75:25 ethanol:water and allowed to air dry for 10-15 minutes. The pellet was resuspended in 30 µL of dimethylpyrocarbonate (DEPC) treated water by heating at 60°C for 15 minutes. The RNA was stored at -80°C. Generally the yield was 30µg per 50 mg of leaves; 60µg per 30 mg of inflorescences.

Before cDNA synthesis 5 µg of RNA was treated with Turbo DNase I (Ambion) according to the manufacturer’s instructions. All cDNA was synthesized with Invitrogen’s SuperScript II reverse transcriptase using an anchor-dT primer at 42°C for 1 hour. The quality of cDNA synthesis and genomic DNA contamination was assessed by PCR amplification of 1 µL of cDNA with actin 2 intron-spanning primers (actin2F and actin2R) using the following cycle conditions: initial denaturation at 94°C for 3 min, 22 cycles of 94°C 30 seconds, 55°C 30 seconds, 72°C 1 min, and final elongation at 72°C for 10 minutes.

To measure the transcript abundance of the two *APT1* mRNA isoforms forward primers were designed to anneal within Exon 1 of *APT1.1* (AP186) and the 5’ UTR of *APT1.2* (187), which are unique to each gene (Appendix 3). The reverse primer (AB19754) annealed within
Exon 7, which is common to both isoforms. To ensure that each primer set was amplifying within its linear range a standard curve was developed by plotting the amount of cDNA amplified (determined by spot density analysis of the digitized image) against the number of cycles (20, 24, 28, 32, and 36) on a semi-log graph. The number of cycles to amplify was chosen by selecting a value midway in the linear range. Using these data, RT-PCR conditions for *APT1.1* and *APT1.2* were carried out as follows: 94°C 3 min, 30 cycles of 94°C 30 seconds, 55°C 30 seconds, 72°C 1 min, and final elongation at 72°C for 10 minutes. An equal volume of each PCR product was electrophoresed on a 1% Tris-acetate-EDTA (TAE) agarose gel at 100V for 16 minutes. The amount of PCR product in nanograms was measured using AlphaImager HP (Alpha Innotech) spot density software.

To measure the transcript abundance of *APT2* primers APT2L and APT2R were used; for APT3 the primers used were APT3RTFwd1 and AB11935. A standard curve was developed to determine the linear range for both primer sets as described above. The following cycle conditions were used for both primer sets: 94°C for 3 min, 30 cycles of 94°C 30 seconds, 55°C 30 seconds, 72°C 1 min, and final elongation at 72°C 10 minutes. The resulting PCR product was analyzed as outlined in the previous paragraph.

**Western Blotting**

Protein extracts were prepared by homogenizing 50-100 mg of leaf tissue or 50 mg of inflorescences in 100 µL 50 mM Tris-HCl pH 8.0 buffer (2 µL buffer per mg of fresh weight). The extracts were centrifuged at 13,200 rpm for 2 min to remove insoluble plant material and the supernatant saved. Protein quantification was determined by the Bradford protein assay (Bio-Rad) using BSA as the standard. These extracts were stored at -20°C.
After quantification of the protein sample, 20 µg of crude protein extract in 1X SDS-loading dye was electrophoresed on a 12.5% acrylamide separating gel layered with a 5% stacking gel. The samples and pre-stained protein ladder (Fermentas, PageRuler SM0671) were initially electrophoresed at 90 V for 20 minutes, after which the voltage was increased to 150 until the 18 kDa pre-stained marker ran off the gel. After electrophoresis, the gel was washed twice for 10 minutes in freshly prepared Bjerrum and Schafer-Nielsen transfer buffer (25 mM Tris, 190 mM glycine, 20% methanol, 0.05% (v/v) SDS). The polyvinylidene fluoride (PVDF) membrane was first soaked in methanol for 30 seconds then pre-soaked, along with the filter papers, in transfer buffer.

Electrophoretic transfer was carried out using Bio-Rad Trans-Blot SD at 20 V for 45 minutes followed by staining with Ponceau s (3% TCA, 0.2% w/v Ponceau s). This stain allowed for reversible detection of proteins on the membrane in order to estimate equal loading of each sample. After removing the stain by rinsing in water the membrane was blocked in polyvinyl alcohol (5 mg per 200 mL; MW 30-70,000) for 30 seconds followed by a quick wash in water (Miranda et al., 1993).

All further washes and antibody incubations were carried out in Phosphate-Buffered Saline solution with 1% (w/v) skim milk powder (Carnation) and 0.3% (v/v) Tween-20 (PBS/M/T). Incubation in primary antibody was for 2 hours at room temperature or overnight at 4°C using the following dilutions: anti-APT1 1:10 000, anti-ADK 1:3500, anti-SAHH 1:5000, anti-GFP 1:1000 (Invitrogen, Cat. No. A6455). APT1 antibody generation will be addressed in the following sections. Sang Lee generated the ADK antibody using full-length ADK1; SAHH
antibody was generated by Shira Davis (Biol 499 Senior Honours Project, University of Waterloo, 2009).

Washes after primary and secondary antibody consisted of two rinses with 10 mL of PBS/M/T followed by three, 10 minute washes in 20 mL PBS/M/T. Incubation in secondary antibody was for 1 hour at room temperature using a horseradish peroxidase-conjugated anti-rabbit IgG (1:10000; Jackson ImmunoResearch). Bound antibody was detected by incubating the membrane in 1 mL of homemade ECL detection reagent (1.25 mM luminol, 2 mM 4-IPBA, 100 mM Tris-HCl pH 8.8, 0.02% H₂O₂) (Haan and Behmann, 2007). The signal was imaged using a chemiluminescent gel documentation system (Model BIS303PC, DNR Bio-Imaging Systems). The system preset “ECL medium size blots” was selected, the detection period set between 0:20 sec to 1:00 min, and gain adjusted to be between 100 and 200.

**Construction of transgenic lines**

Unless otherwise noted, all the construction of all transgenic plants involved the following steps and cloning into the pSAT series of vectors (Tzfira et al., 2005). The ligation mixture was introduced into DH5α for plasmid propagation and colony screening. Plasmid isolated from the DH5α host was sequenced around the insert and junction regions. The *promoter-insert-epitope tag-transcriptional terminator* fragment was subcloned into the pPZP-RCS2-bar binary vector and re-introduced into DH5α for plasmid propagation. This plasmid was transformed into the GV3101 *Agrobacterium tumefaciens* used to mediate stable transformation of the transgene into the Arabidopsis genome. As the pPZP-RCS2-bar binary vector contains the herbicide resistant gene *bar*, all transgenic lines are resistant to BASTA (Sigma) and Wipeout (Wilson Laboratories Inc.). Table 2 summarizes the transgenic lines constructed for this research.
Further details on the generation of each construct will follow; primer sequences used in the cloning process can be found in Table 3.

Table 2. List of transgenic lines generated throughout this study.

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<th>Name</th>
<th>Genetic background (transgene)</th>
<th>Purpose</th>
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<td>APT1::Ex1:GFP</td>
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<td>Subcellular localization. Expression of Exon 1 from APT1.1 promoter</td>
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<td>ami1.2</td>
<td>WT (35S::ami1.2)</td>
<td>Silencing of APT1.2 transcript by targeting its 5’UTR</td>
</tr>
<tr>
<td>APT1.1:StrepII</td>
<td>WT (35S::APT1.1:HA:StrepII)</td>
<td>APT1.1 fused to StrepII tag to purify interacting proteins</td>
</tr>
<tr>
<td>APT1.2:StrepII</td>
<td>WT (35S::APT1.2:HA:StrepII)</td>
<td>APT1.2 fused to StrepII tag to purify interacting proteins</td>
</tr>
<tr>
<td>actP::APT1.1</td>
<td>WT, oxt1, apt1-3 (actin promoter::APT1.1 ORF) oxt1 (see Appendix 4)</td>
<td>Complementation of apt1-3 with APT1.1 isoform</td>
</tr>
<tr>
<td>actP::APT1.2</td>
<td>WT, oxt1, apt1-3 (actin promoter::APT1.2 ORF) oxt1 (see Appendix 4)</td>
<td>Complementation of apt1-3 with APT1.2 isoform</td>
</tr>
</tbody>
</table>
### Cloning of APT1-GFP fusion proteins

The $APT1::gAPT1$ clone was constructed by amplifying the segment of genomic DNA extending from 800 bp upstream of the transcriptional start site of the $APT1.1$ promoter, through the 5’ UTR of $APT1.1$, and terminating at the end of Exon 1 using the primers AP185 and APex1R. This fragment was cloned into the AgeI/BamHI restriction sites of pSAT6 in-frame with eGFP sequence (strain #800, Tzfira et al., 2005). The resulting fusion gene was subcloned into the PI-PspI restriction site of the pPZP-RCS2-bar binary vector.

The $APT1::gAPT1\Delta\text{Intron 1}:GFP$ clone was constructed by amplifying the same fragment that was used in the $APT1::gAPT1$ clone, with the addition of Intron 1 and Exon 2, using the primers AP185 and APex2R. This fragment was cloned into the AgeI/BamHI restriction sites of the pSAT6 vector in-frame with eGFP (strain #800, Tzfira et al., 2005) then further subcloned into the PI-PspI restriction site of pPZP-RCS2-bar.

Construction of the $actP::APT1.1:2GFP$ clone was a two-step process that involved cloning the actin promoter into a previously modified pSAT6-2eGFP vector (Sanghyun Lee, strain #1047) followed by the addition of the APT1.1 ORF. First, the actin promoter was subcloned from the pSAT7-actP-MCS-agSt vector (strain #1376) into the AgeI/NcoI sites of the
pSAT6-2eGFP vector. In the second step, the *APT1.1* ORF (lacking the 5’ UTR) was amplified from WT cDNA using the primers AP181 and AB19750. This fragment was cloned into the *Nco*I site of the modified *actP*-pSAT6-2eGFP vector. This construct was further subcloned into the PI-*PspI* restriction site of pPZP-RCS2-bar.

Creating the *actP::APT1.2::GFP* clone was also a two-step cloning process. The actin promoter was first subcloned from the pSAT7-actMCS-agST vector into the *AgeI/NcoI* sites of pSAT6-eGFP vector (strain #800). The *APT1.2* ORF (lacking its 5’ UTR) was amplified from WT cDNA using primers APT cDNA and AB19750. The *APT1.2* ORF was cloned into the *NcoI* restriction site of the modified *actP*-pSAT6-eGFP vector and the resulting fragment was subcloned into the PI-*PspI* restriction site of pPZP-RCS2-bar.

**Cloning of amiRNA constructs**

Artificial microRNAs (amiRNAs) were designed using the online tool *WMD 2 – Web MicroRNA Designer* created by Schwab *et al.*, 2005. The complete protocol can be found online; here I will briefly describe the design and cloning steps only. The online tool requires the user to enter a sequence of interest that is to be the target of the amiRNA. For this project, the 5’ untranslated regions for *APT1.1* and *APT1.2* were used, as they are unique to each transcript. The program outputs a ranked list of amiRNAs and their specific target regions. There is no guarantee regarding which amiRNA will be most efficient, although it is recommended to select the amiRNAs ranked at the top of the list. Once an amiRNA is selected the program designs a set of four overlapping oligonucleotides that are used in site-directed mutagenesis using the pRS300 plasmid as the initial template (primers are listed in Table 3). In this step the user is essentially replacing an amiRNA in the pRS300. The resulting amiRNA-producing fragment was cloned
into the EcoRI/BamHI restriction sites of the pSAT4 vector under control of the constitutive 35S promoter. The fusion gene was subsequently subcloned into either the I-SceI or PI-PspI restriction site of pPZP-RCS2-bar; ami1.1 was cloned into I-SceI, ami1.2 was cloned into PI-PspI.

**Cloning of APT1.1 and APT1.2 with StrepII epitope tag**

The ORFs of *APT1.1* and *APT1.2* were amplified from WT cDNA using primer sets pXCS-Ex1F/pXCS-APT1R and pXCS-Ex2F/pXCS-APT1R, respectively. The PCR products were subsequently cloned into the *HindIII/PstI* restriction sites of pXCS-HAStrep (Witte *et al.*, 2004) in-frame with the StrepII tag. This design fuses the StrepII tag to the carboxy terminus of each ORF. The resulting plasmids were transformed into DH5α for plasmid propagation and screening. The fusion protein sequence was verified by DNA sequencing and the plasmid was transformed into the *A. tumefaciens* vector GV3101 MP90RK. It was essential to use this Agrobacterium strain as the oriV on the pXCS vector is dependent on the *trf/tra* genes provided by GV3101 MP90RK to be functional (Koncz and Schell, 1986).

**Cloning of APT1.1 and APT1.2 ORFs**

The *APT1.1* coding region including its stop codon was amplified using primers AP181 and AB12380 and cloned into the *NcoI/HindIII* restriction site of the actP-pSAT4 vector (strain #1500). This places the ORF under the control of the actin promoter. The fusion gene was subcloned into the PI-PspI restriction site of pPZP-RCS2-bar.

The APT1.2 coding region including its stop codon was subcloned from a pET30a plasmid (strain #592) into the *NcoI/HindIII* restriction sites of the actP-pSAT6 vector (strain #1501). The fusion gene was subcloned into the I-SceI restriction site of pPZP-RCS2-bar.
Table 3. DNA sequence of primers used in cloning and RT-PCR experiments outlined in this thesis.

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence (5’ to 3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AP185</td>
<td>5’-CAGACCGGTGAGTTTACTTCG -3’</td>
</tr>
<tr>
<td>APex1R</td>
<td>5’- TCAGGATCCAGGCGAGGAG -3’</td>
</tr>
<tr>
<td>APex2R</td>
<td>5’- TCAGGATCCATGGTTTAGGAAG -3’</td>
</tr>
<tr>
<td>AP181</td>
<td>5’- CTGCCATGGGAATGCAAACAATAATAATTTCCTCC -3’</td>
</tr>
<tr>
<td>AB19750</td>
<td>5’- TAGCCATGGCAGCAGCCGACTTTACAAG -3’</td>
</tr>
<tr>
<td>APT cDNA</td>
<td>5’-TGCCATGGCGACTGAGATGTGC-3’</td>
</tr>
<tr>
<td>miRAPT1Ex1A-I</td>
<td>5’-GATAATAATGGACAAAGCCGTCGTCCTCTCTTTTTGTATTCC-3’</td>
</tr>
<tr>
<td>miRAPT1Ex1A-II</td>
<td>5’-GAGACGGCTTTTGTCCATTATTATCAAAGAGAATCAATGA-3’</td>
</tr>
<tr>
<td>miRAPT1Ex1A-III A</td>
<td>5’-GAGAAGGCTTTTGTCCATTATTTCACAGGTCGTGATGATG-3’</td>
</tr>
<tr>
<td>miRAPT1Ex1A-IV</td>
<td>5’-GAAAAATAATCGACAAGGACCTTCTCTCATATATATTTCC-3’</td>
</tr>
<tr>
<td>miRAPT1Ex2A-I</td>
<td>5’-GATTTAGCAAAAAATCTCGGCCGTGCTCTCTTTTTGTATTCC-3’</td>
</tr>
<tr>
<td>miRAPT1Ex2A-II</td>
<td>5’-GACACGGCGAGTATTTTGCTAAATCAAAGAGAATCAATGA-3’</td>
</tr>
<tr>
<td>miRAPT1Ex2A-III</td>
<td>5’-GACAAGCCGAGTATTATGCTAATTCACAGGTCGTGATGATG-3’</td>
</tr>
<tr>
<td>miRAPT1Ex2A-IV</td>
<td>5’-GAATTAGCATAATATCGGCTTTGCTACATATATATTTCC-3’</td>
</tr>
<tr>
<td>pXCS-Ex1F</td>
<td>5’- TCGAAGCTTATGCAAAACAATAATAATTTCCTCC -3’</td>
</tr>
<tr>
<td>pXCS-APT1R</td>
<td>5’- AGCTGCAGAGCAGCCGACTTTACAAGAC -3’</td>
</tr>
<tr>
<td>pXCS-Ex2F</td>
<td>5’- TCG AAGCTT ATG GCG ACT GAA GAT GTG C -3’</td>
</tr>
<tr>
<td>AB12380</td>
<td>5’- CCTAAGCTTTTTCTCCAGTTTC -3’</td>
</tr>
</tbody>
</table>
Stable transformation of Arabidopsis using the floral dip method

The floral dip protocol is a simple method to utilize Agrobacterium tumefaciens to mediate the transformation of a transgene of interest into the Arabidopsis genome. The protocol outlined here is adapted described by Clough and Bent (1998). Briefly, the Agrobacterium GV3101 strain carrying the transgene of interest was used to inoculate 5 mL of Luria-Bertani (LB) Media supplemented with rifamycin (25 µg/mL), gentamicin (25 µg/mL), spectinomycin (100 µg/mL), and streptomycin (300 µg/mL). After incubation overnight at 28°C, this culture was used to inoculate 100 mL of LB media supplemented with the same antibiotics and
incubated for a second overnight at 28°C. The culture was centrifuged at 5500 rpm for 10 minutes, resuspended at an OD$_{600}$ of 0.8 in 5% (w/v) sucrose, 0.05% (v/v) Silwett-L77 (Lehle Seeds, Cat. No. VIS-01) and placed in a tin foil baking dish (diameter 20 cm, depth 5 cm).

Approximately 100 seedlings were grown in 16 cm x 12 cm x 5.5 cm containers, for transformation. At approximately 4 weeks of age, when the plants had just developed a primary shoot with a few flowers, the flowers were submerged in Agrobacterium solutions prepared as described above; the plants were gently agitated for 5 seconds. After dipping the plants were placed in a growth tray, covered with plastic dome to maintain high humidity, and kept at room temperature away from direct sunlight for 24 hours. After this time, the dome was removed and the plants were transferred back to the growth chambers until seed set. After 2 to 3 weeks the plants were allowed to dry out for harvesting of seeds.

The resulting T1 seeds were sown on soil at a high density and maintained in growth chambers. T1 transformants carrying a Basta resistance gene were selected based on their ability to grow following foliar application of the commercial herbicide “Wipeout” (0.067% (v/v); Wilson Laboratories Inc). At 14 days, plants were sprayed with Wipeout once a day, for five continuous days. Transformants make healthy true leaves in these conditions while non-transformed seedlings turn yellow and die. The Basta-resistant T1 plants were transplanted into individual pots for seed set.

**Genomic DNA extraction and genotyping of the apt1-3 allele**

Leaf tissue was first homogenized in 50 µL extraction buffer (0.2 M Tris-HCl pH 7.2, 0.25 M NaCl, 0.025 M EDTA, 0.5% (w/v) SDS). After homogenization 350 µL of extraction buffer was added before vortexing for 5-10 sec. The homogenate was centrifuged for 1 min at
14,000 rpm. The resulting supernatant, 300 µL, was transferred to a new tube and an equal volume of isopropanol was added to precipitate DNA by inverting the mixture multiple times. After a 2 min incubation period at room temperature the DNA was pelleted by centrifugation at 14,000 rpm for 5 min. The pellet was allowed to air dry before resuspension in 50 µL of TE buffer (10 mM Tris-HCl pH 7.5, 1mM EDTA).

The WT APT1 gene contains three BstNI restriction enzyme sites. The mutation of a single nucleotide within the BstNI site in the apt1-3 allele does not allow the BstNI enzyme to recognize this site thus yielding restriction fragments that are different from those of the WT gene (Gaillard et al., 1998). The appropriate region of the APT1 gene was PCR amplified using the APT Forward and AB19754 primers using the cycle conditions: 94°C 4:00 min, 36 cycles of 94°C 30 sec, 52°C 30 sec, 72°C 1:15 min, and final elongation of 72°C 10 min. One half of the PCR reaction was used for the restriction enzyme digest using BstNI for 2h at 60°C. BstNI digestion of the WT PCR product produces fragments of 80, 343, 461 and 479 bp; similar digests of the corresponding PCR fragment from an apt1-3 mutant produces fragments of 80, 363, and 940 bp. These were most easily detected by electrophoresis of the complete digest on a 1% TAE agarose gel.

**Affinity purification of StrepII-tagged proteins**

StrepII-tagged proteins and their associated protein interactors were isolated using the following protocol that was adapted from Witte et al., 2004. One gram of leaf tissue was homogenized in liquid nitrogen and resuspended in 2 mL Ex-Strep buffer: 100 mM Tris-HCl pH 8.0, 5 mM EDTA, 200 mM NaCl, 10 mM dithiothreitol (DTT; from a 100 mM stock solution), 0.5% (v/v) Triton X-100, 10% (v/v) glycerol, 1 mM phenylmethyl sulfonyl fluoride (PMSF;
prepared fresh as a 100 mM stock in methanol), and 200 µL of protease inhibitor cocktail (Roche). The protease inhibitor cocktail is provided as a tablet; 1 tablet was dissolved in 1 mL of sterile water and 100 µL was added per mL of Ex-Strep buffer.

The homogenate was centrifuged at 4°C for 20 minutes, at 20,000 g. Before incubation with the StrepTactin Macroprep beads (Novagen), 40 µL of resin per purification was centrifuged at 700 g for 30 seconds to pellet the beads and allow removal of storage buffer in the supernatant fraction. The beads were washed once with 1X PBS and 2X with Ex-Strep buffer. The beads where incubated with the supernatant by end-over-end mixing for 15 minutes at 4°C. The mixture was centrifuged at 700 g for 30 seconds and the resin washed five times with 0.8 mL of W-Strep buffer: 100 mM Tris-HCl pH 8.0, 0.5 mM EDTA, 200 mM NaCl, 2 mM DTT 0.5% (v/v) Triton X-100, 10% (v/v) glycerol, 1 mM PMSF, and protease inhibitor cocktail (100 µL per mL of W-Strep buffer; Roche). After the final wash the resin was resuspended in 20 µL 2X SDS loading buffer and boiled for 6 minutes to aid the dissociation of StrepII tag from the strepavidin beads. The boiled mixture was placed on ice for 10 minutes to allow the beads to settle and facilitate the removal of the supernatant.

Protein concentration was not determined for any of the extractions as the protein was eluted directly into SDS loading dye. However, over-expressed StrepII fusion protein is easily detected by less sensitive staining methods such as Coomassie staining. If the intended purpose is to excise polypeptides for mass spectrometry analysis it is recommended to use either silver staining or Sypro Ruby Gel Protein Stain (Invitrogen). Polyacrylamide gels were stained with Sypro Ruby according to the manufacturer’s instructions. Briefly, the gel was incubated twice in 100 mL fixative solution (50% methanol, 7% acetic acid) for 30 min. The gel was moved into a clean container and stained with 60 mL of Sypro Ruby gel stain overnight. The next day, the gel
was placed in a new container and washed with 100 mL Wash solution (10% methanol, 7% acetic acid) for 30 min. This was followed with two washes in MilliQ water for 5 min after which the gel was viewed and imaged using a 300 nm UV transilluminator.

**Cloning and over-expression of recombinant APT1.1 and APT1.2 proteins**

Recombinant APT1.1 and APT1.2 were expressed in *E. coli* using Novagen’s pET28 and pET30 vectors. These vectors fuse six histidine residues (6xHIS) at the carboxy terminus of the protein product providing a small and highly specific epitope tag for protein purification. The small size of the 6xHIS helps alleviate concerns of the tag interfering with protein folding and protein activity. A second feature of the pET system is the ability to control expression of the recombinant protein using isopropyl-D-thiogalactopyranoside (IPTG). IPTG induces expression of the T7 RNA polymerase gene contained in DE3 lysogens, leading to synthesis of the fusion protein.

The *APT1.2* ORF was cloned into the *NcoI/HindIII* restriction sites of pET30a (Antonio Facciulo, Biol 499 Senior Honours Project, University of Waterloo, 2007). The ligation mix was transformed into DH5α (strain #591) and once the desired clone was identified it was introduced into the expression host BL21(DE3)pLysS (strain #601).

The *APT1.1* ORF was amplified from WT cDNA using the primers apt1-exon1 and AB12380. This fragment was cloned into the *NdeI/HindIII* restriction sites of the pET28a vector. The plasmid was transformed into DH5α (strain #1304) and the BL21(DE3)pLysS expression host (strain #1305). The *APT1.1* ORF was also amplified from WT cDNA using the primers apt1-exon1 blunt and AB12380. This fragment was subsequently cloned into the *EcoRI/HindIII* sites of pET30a. The recombinant APT1.1 was recovered from the expression hosts.
BL21(DE3)pLysS and CodonPlus BL21(DE3)RIPL creating strains #601 and #1305. APT enzymatic assays on the APT1.1-6xHIS recombinant protein presented in this thesis were derived from strain #601.

The APT1.1-6xHIS and APT1.2-6xHIS recombinant proteins were over-expressed and purified under the same conditions. First, each strain was used to inoculate a 5-10 mL overnight LB starter culture supplemented with kanamycin (50 µg/mL) and chloramphenicol (25 µg/mL). The starter culture was used to inoculate 250 mL LB media supplemented with the same antibiotics as above and incubated at 37°C until the OD$_{600}$ reached 0.8. At this point T7 expression was induced by adding 0.2 mM IPTG; the culture was incubated at 28°C for 5 h. The cells were collected by centrifugation at 4°C at 13,200 rpm for 15 min. The supernatant was discarded and the pellet flash frozen on dry ice for 10 min before being stored at -20°C.

Freshly prepared cells or frozen cell paste were placed on ice or kept at 4°C for the remainder of the purification protocol. The pellet was resuspended in 20 mL Binding Buffer (Table 4) plus 0.1% (v/v) Tween 20 to reduce the viscosity of the crude cell extract. The resuspended cells were lysed by sonicing 4X for 30 sec, with a 1 min rest period on ice between each sonication (Sonic Dismembrator 100, Fisher Scientific). This extract was centrifuged at 4°C 13,200 rpm for 30 min and the supernatant filtered through a 0.45 µm filter to remove cellular debris.

To capture 6xHIS-tagged proteins Novagen’s Ni-NTA resin was used for column chromatography. The composition of each buffer can be found in Table 4. One mL of Ni-NTA resin was prepared by loading the column bed with 3 volumes of strip buffer followed by 3 volumes of water, 5 volumes of charge buffer, and 3 volumes of bind buffer at a flow rate of 0.5
mL/min. The protein extract was loaded on the column resin and the flow through passed over the column to maximize capture of 6xHIS protein. The column bed was washed with 10 volumes of bind buffer followed by 12 volumes of wash buffer, and eluted using 3 volumes of elution buffer. The purified protein was concentrated and dialyzed using a Centricon Centrifugal Filter Device with 10,000 MW membrane cut-off (Millipore). For enzymatic assays the protein was dialyzed in 50 mM Tris-HCl pH 7.5; for antibody generation the protein was dialyzed in sterile PBS.

Table 4. Buffers used in purifying 6xHIS recombinant protein by affinity chromatography

<table>
<thead>
<tr>
<th>Buffer</th>
<th>Composition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charge Buffer</td>
<td>50 mM NiSO₄</td>
</tr>
<tr>
<td>Strip Buffer</td>
<td>100 mM EDTA, 0.5 M NaCl, 20 mM Tris-HCl pH 7.9,</td>
</tr>
<tr>
<td>Bind Buffer</td>
<td>5 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl pH 7.9</td>
</tr>
<tr>
<td>Wash Buffer</td>
<td>60 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl pH 7.9</td>
</tr>
<tr>
<td>Elution Buffer</td>
<td>400 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl pH 7.9</td>
</tr>
</tbody>
</table>

**APT1 polyclonal antibody generation**

APT1.2-6xHIS recombinant protein was generated, purified, and dialyzed in sterile PBS buffer as outlined in the previous section. A white New Zealand rabbit was given four booster injections. The first booster consisted of 500 µg of recombinant protein in PBS mixed with an equal volume of Freund’s complete adjuvant. The remaining three boosters were administered every 3 weeks and consisted of 300 µg of recombinant protein mixed with an equal volume of Freund’s incomplete adjuvant.
Three weeks after the last booster injection the rabbit was bled and approximately 150 mL of blood was collected. To separate the blood from the antiserum, it was placed at 37°C for 2 hours to allow the blood to naturally clot. The clot was pelleted by centrifugation for 15 min at 1500 x g and the supernatant transferred to a new tube. The antiserum was aliquoted, flash frozen in liquid nitrogen, and stored at -80°C.

The APT1-antibody was tested for sensitivity and specificity by immunoblotting. To test for sensitivity, 10 ng, 1 ng, and 500 pg of APT1.2-6xHIS recombinant protein was probed with the APT1 antibody at a dilution of 1:5000 and 1:10,000 (Appendix 1). These blots showed that the antibody could detect 500 pg of protein at a dilution of 1:10,000. Next, specificity was tested by probing WT and an APT1-deficient plant extract (oxt1). Using a dilution of 1:10,000, an abundant polypeptide was detected in WT at 24 kDa representing APT1.2; a faint band was detected in the APT1-deficient mutant.
Chapter 1 - Results

Recovering APT1.1 and APT1.2 full-length cDNAs

The first objective of this research was to determine if APT1.1 encodes a functional mRNA. The fact that full-length cDNAs for APT1.1 have been previously recovered by selecting for mRNA species containing a 5’-guanidine cap indicates that it is a functional mRNA. Capped cDNAs can be found in the Ceres cDNA clone collection (NCBI AY084300) and the RIKEN Arabidopsis full-length cDNA clone collection (NCBI BP837943). I attempted to recover these full-length cDNAs from WT leaves using Ambion’s RLM-RACE kit. The initial step in this technique involves treatment of total RNA with calf intestinal phosphatase. This step removes 5’ phosphate groups from all RNA species except for mRNAs, which have their 5’phosphate group protected from enzymatic cleavage by the 5’-guanidine cap. The second step uses tobacco acid pyrophosphatase to remove the 5’-guanidine cap exposing a 5’-phosphate group, allowing for a suitable substrate for ligation to a DNA adaptor, in a reaction mediated by RNA ligase.

The subsequent cDNA synthesis initiates from primers binding to the polyA tail and the adaptor; amplification by PCR provides sufficient template for DNA sequencing of specific cDNAs. RLM-RACE was successful in recovering the APT1.2 full-length cDNA previously reported by Moffatt et al., 1994. No full-length cDNA was recovered for APT1.1.

It was possible that RNA secondary structure, particularly in the 5’ end of the mRNA limited recovery of the full-length transcript. The methods used by RIKEN involved the use of trehalose, an agent known to stabilize reverse transcriptase at high temperatures (Carninci et al., 1998; Seki et al.,1998) . Thus a higher temperature cDNA synthesis reaction was carried out at
and 42°C and 55°C using Roche’s Transcriptor High Fidelity cDNA synthesis kit. Several different forward primers were used for the PCR amplification step, to determine the 5’ end point of the cDNA product. AP183 anneals to the reported 5’ terminal end of the transcript; AP184 anneals 100 base pairs into the 5’ UTR, and AP187 anneals 30 base pairs upstream of the start codon (Figure 5, Appendix 3). PCR amplification using AP183 was unsuccessful for cDNA synthesized at both temperatures; PCR amplification using AP184 and AP187 consistently yielded DNA fragments of the expected sizes (787 bp for AP184 and 737 bp for AP187). The identity of the smaller fragment in the AP184 reaction at 55°C was not determined. Based on the terminal end of the 5’ UTR of APT1.1 may contain significant secondary structure that blocked my attempts to recover the full-length cDNA of APT1.1 as reported by Ceres and RIKEN. However I was confident that their data were reliable and did not invest further effort in duplicating their results.
Figure 5. PCR primer walking along the *APT1.1* 5′UTR. RNA was extracted from leaf tissue and cDNA synthesized at 42°C and 55°C using Roche Transcriptor High Fidelity cDNA synthesis kit. Three forward primers were designed: AP183 to anneal to the 5′ end of the cDNA, AP184 to anneal 100 bp into the 5′ UTR, and AP187 to anneal 30 bp upstream of Exon 1. The reverse primer AB19754, anneals within the terminal exon, was used with all three forward primers.

**Expression analysis of APT1.1 and APT1.2**

The identification of two *APT1* transcripts in numerous expressed sequence tag collections along with the recovery of apparently full-length cDNAs lead to the second objective of this research: comparing the expression profiles of *APT1.1* and *APT1.2* in WT and the APT1-deficient mutants *oxt1* and *apt1-3*. The *oxt1* mutant contains a T-DNA insertion element between Exon 1 and Exon 2; *apt1-3* contains a point mutation in the Exon 4 splice junction. RT-PCR analysis was used to evaluate the impact of each of these mutations on the *APT1.1* and *APT1.2* transcript levels, as compared to the corresponding transcript levels in WT.

Semi-quantitative RT-PCR was carried out using forward primers specific to the 5′ UTRs of each transcript and a reverse primer, located in the terminal exon which is common to both *APT1.1* and *APT1.2* transcripts. Each primer’s linear range was determined using WT cDNA as a template. Parallel reactions were initiated and starting at the 20th cycle, one reaction was stopped every 4 cycles, until the 44th cycle was reached. The amount of amplified product in each case
was quantified in nanograms using spot density analysis of the digitized image and the molecular
weight markers as standards; the results were plotted against cycle number, on semi-log graph
(Figure 6). Based on these data, I concluded that PCRs of 30 cycles were within the linear range
for both reactions and would provide a meaningful relative abundance estimate.
Figure 6. Linear range estimate of APT1.1 and APT1.2 primer sets for RT-PCR analysis. WT cDNA template was amplified by PCR for 20 through 44 cycles and electrophoresed on a 1% (w/v) TAE agarose gel. The gel image was digitized and each DNA fragment quantified, in nanograms, using spot density analysis. The nanograms of cDNA synthesized were plotted against the cycle number on semi-log graph paper. Thirty cycles were used for subsequent RT-PCRs using these primer pairs.
In WT, \textit{APT1.1} and \textit{APT1.2} transcripts were equally abundant in both expanded leaves (Figure 7A) and WT inflorescences (Figure 7B). \textit{oxtl}, which contains a T-DNA insertion within Intron 1, lacked the expected \textit{APT1.1} transcript in both leaves and inflorescences. A fore-shortened \textit{APT1.2} transcript was present in both tissues of \textit{oxtl}, in the same abundance as in WT.

Three distinct \textit{APT1.1} and \textit{APT1.2} related RT-PCR products were detected when mRNA from \textit{apt1-3} leaves or inflorescences was used as a template. This was somewhat unexpected since \textit{apt1-3} contains a point mutation in the splice junction of Exon 4 which was anticipated to trigger rapid turnover of both \textit{APT1.1} and \textit{APT1.2} transcripts. To identify the mRNA template of these products, each was gel purified and sent for sequencing. The results showed that the largest PCR product was amplified from an \textit{APT1} transcript that had retained the third intron, consistent with the site of the \textit{apt1-3} mutation. The second largest PCR product recovered from the \textit{apt1-3} reaction was identical to the WT \textit{APT1} product, consistent with there being some normal splicing of the primary transcript. The smallest PCR product was copied from an mRNA that was missing Exon 4. Thus, these RT-PCR results confirmed that Intron 3 splicing was abnormal in \textit{apt1-3} resulting in the synthesis of aberrant transcripts. More importantly the analysis showed that \textit{APT1.1} and \textit{APT1.2} transcripts were present in approximately equal abundance in WT leaves and inflorescences. These transcripts were missing in the APT1-deficient mutants \textit{oxtl} and \textit{apt1-3}, consistent with the mutation each contains.
Figure 7. RT-PCR analysis of *APT1.1* and *APT1.2* in leaves and inflorescences. Transcript abundance was measured using forward primers specific to each transcript variant’s 5’ UTR. The reverse primer, common to both transcripts, anneals with the terminal exon. WT organs showed equal expression of *APT1* isoforms in both leaves (A) and inflorescences (B). The oxt1 mutants had no *APT1.1* product but did have an equal expression of *APT1.2* as in WT. apt1-3 lines show three aberrant *APT1* transcripts. Actin was used as control to test that the same amount of cDNA was added to each PCR.

The next questions addressed were: are both APT1.1 and APT 1.2 transcripts translated in WT plants and if so, how do the protein and transcript abundances compare? This analysis was carried out by immunoblotting of crude extracts of leaves and inflorescences of four-week-old plants. *APT1.2* was easily detected in WT leaves while *APT1.1* was present at approximately one-tenth the level of *APT1.2* (Figure 8, compare Lanes 5 and 6). *APT1.2* was slightly more
abundant in inflorescences, than in leaves, while APT1.1 was undetectable (Figure 8, Lanes 7 and 8). Thus, although the *APT1.1* and *APT1.2* transcripts levels were similar in leaves, the immunoblots clearly showed that either the two transcripts were not translated with equal efficiency or that the APT1.1 protein is turned over more rapidly relative to APT1.2. APT1.1 transcripts were subject to post-transcriptional regulation that appeared to differ in extent, in leaves and inflorescences.

![Image](image_url)

Figure 8. APT1.1 and APT1.2 protein abundance in WT leaves and inflorescences. Leaf and inflorescence protein extracts of 10µg and 20µg (with two biological replicates of each) were probed with rabbit anti-APT1 antibody; bound antibody was detected by anti-rabbit antibody conjugated to HRP. Polypeptides in lanes 1, 2, 3, 4, 7, and 8 are approximately 24 kDa. Polypeptides in lanes 5 and 6 are 24 kDa and 26 kDa, respectively.

A similar immunoblot analysis was carried out on leaf and floral extracts of the APT1-deficient mutants, *apt1*-3 and *oxt1*. In *apt1*-3, expression of APT1.1 and APT1.2 was below the level of detection, which is consistent with the 1-2% residual APT activity in this mutant (Figure 9). In *oxt1*, a faint APT1.2 polypeptide was visible after a normal detection period (20 sec); this polypeptide became more apparent after a longer exposure (1 min 10 sec). Based on these data it was estimated that *oxt1* has about 1% of the APT1.2 present in WT leaves. No APT1 polypeptides were detected in *apt1*-3 leaf and floral extracts even after long detection periods of the immunoblots (data not shown).
Figure 9. Abundance of APT1.1 and APT1.2 in WT and APT1-deficient mutants. Leaf protein extracts (20µg each lane) were probed with anti-APT1 antibody and bound antibody detected using a rabbit secondary antibody conjugated to HRP. The immunoblot was visualized for two different detection periods; only the longer detection period revealed APT1 in oxt1. The membrane was stained with Ponceau S to test for equal loading of samples.

The absence of an APT1.1 polypeptide in oxt1 and apt1-3 indirectly confirmed that the faint APT1.1 detected in WT leaves is indeed an APT1 protein. Its weak detection on initial immunoblots was not convincing that this polypeptide was APT1.1. Rather it could be regarded as a nonspecific target. To address this concern a direct approach was used to confirm that translation initiation occurs at Exon 1 in vivo.

**APT1.1 coding capacity and regulation**

Two transgenes were constructed to test whether the APT1 Exon 1 is translated in vivo and thus prove that APT1.1 is a functional mRNA (Figure 10A). The first construct contained 800 base pairs upstream of the APT1.1 transcriptional start site, plus the 5’ UTR and the first Exon, fused in-frame with GFP (APT1::Ex1:GFP). The second construct contained the same elements (APT1 promoter region, 5’UTR , Exon 1) in addition to Intron 1, and Exon 2, again fused in-frame with GFP (APT1::Ex1-i-Ex2:GFP). The premise behind the design of these
constructs was that if translation initiation occurs at Exon 1 then a polypeptide of 32 kDa would be produced in APT1:Ex1:GFP plants (26 kDa GFP +7 kDa Exon 1) whereas translation of the second transgene would produce both a 36 kDa polypeptide (26 kDa GFP + 7 kDa Exon 1 + 3 kDa Exon 2) and a 29 kDa polypeptide (26 kDa GFP + 3 kDa Exon 2).

Each transgene was introduced into WT Arabidopsis and four stable T1 transformants for each construct were analyzed by immunoblotting. Crude leaf extracts were separated by electrophoresis through SDS-PAGE, blotted to membranes and probed for the presence of GFP. APT1:Ex1:GFP plants synthesized a polypeptide of 28 kDa, slightly smaller than the expected 32 kDa but larger than GFP alone (Figure 10B). The APT1::Ex1-i-Ex2:GFP plants were predicted to express two GFP-fusion polypeptides: the 36 kDa polypeptide initiated at Exon 1 and a 29 kDa polypeptide representing translation initiation at Exon 2. Interestingly, only the 29 kDa was detected in these plants.

Although the results were not exactly as anticipated, these immunoblotting data confirmed that translation initiated at Exon 1 and an APT1.1 polypeptide is synthesized in vivo as observed in APT1::Ex1:GFP plants. In APT1::Ex1-i-Ex2:GFP plants only a polypeptide of 29 kDa was detected with the anti-APT1 antibody representing translation initiation occurring at Exon 2. In these same plants there was no detectable translation initiation of Exon 1.

These results showed that translation initiation at Exon 1 is greatly reduced by the presence of Intron 1 and Exon 2. Since these transgenes were expressed in a WT background, which has functional APT1 genes, it is evident that the inhibition of APT1.1 synthesis by Intron 1-Exon 2 occurred only when these elements are immediately downstream of Exon 1, as in the APT1::Ex1-i-Ex2:GFP transgene; this transgene did not affect expression of the WT gene.
Thus the $\textit{APTI}::\text{Ex1-i-Ex2:GFP}$ transgene mimicked expression of the WT $\textit{APTI}$ gene in leaves and inflorescences in that a polypeptide was initiated at Exon 2 (Ex2:GFP fusion) while the corresponding APT1.1 polypeptide was undetectable. To confirm this post-transcriptional effect I needed evidence that $\textit{APTI}.1$ and $\textit{APTI}.2$ transcripts are produced from the $\textit{APTI}::\text{Ex1-i-Ex2:GFP}$ transgene, and that the absence of APT1.1 protein was not an artifact of this particular transgenic construct.

RT-PCR was used to investigate the transcriptional products arising from the $\textit{APTI}::\text{Ex1-i-Ex2:GFP}$ transgene. RNA was extracted from the leaves of two independent homozygous $\textit{APTI}::\text{Ex1-i-Ex2:GFP}$ plant lines (2-2 and 2-4). Two forward primers were used for RT-PCRs specific to the 5’ UTR of $\textit{APTI}.1$ and $\textit{APTI}.2$, respectively; the reverse primer was specific for the GFP open reading frame, which was located at the 3’ end of both transgenes. The selection of this reverse primer ensured that only $\textit{APTI}$ transcripts arising from the transgenes and not from the WT allele would be amplified (Figure 10C). This RT-PCR was not done semi-quantitatively as this was a test for the presence or absence of two transcriptional products; 36 PCR cycles were arbitrarily chosen. Amplification of the $\textit{APTI}.1$ transcript using these primers was expected to yield a product of 428 bp whereas amplification of the $\textit{APTI}.2$ transcript would yield a 360 bp fragment. Both of these products were detected in the $\textit{APTI}::\text{Ex1-i-Ex2:GFP}$ plants.
Figure 10. Transcript and protein products arising from *APT1*:Ex1:GFP transgenes. A, Schematic of *APT1*:Ex1:GFP and *APT1*:Ex1-i-Ex2:GFP transgenes. Arrows represent primers used for RT-PCR. B, Immunoblot analysis of GFP protein synthesized from the 1-6 line of *APT1*:Ex1:GFP and the 2-4 line of *APT1*:Ex1-i-Ex2:GFP. The left blot was probed with anti-GFP antibody; the right with anti-APT1 antibody, and the top blot with anti-SAHH antibody. In all cases bound antibody was detected using anti-rabbit antibody conjugated to HRP. SAHH was used to test for equal loading. C, RT-PCR analysis of *APT1*:Ex1-i-Ex2:GFP transgene. RNA used for cDNA synthesis was extracted from leaves of two independent lines (2-2 and 2-4). Forward primers used for RT-PCR were specific to each isoform’s 5’ UTR. The reverse primer anneals within the GFP ORF and is common to both transcripts.
The RT-PCR detected a second, larger APT1.1 product arising from the APT1::Ex1-i-Ex2:GFP transgene. The size of this fragment, approximately 600 to 700 bp, was larger than the expected 428 bp fragment, but consistent with the size of the primary APT1.1 mRNA (i.e. mRNA in which the intron was not spliced out; 648 bp). This fragment was not sequenced so this conclusion has yet to be verified unequivocally.

Taken together with the immunoblot analysis these data showed that the APT1.1 transcript is post-transcriptionally regulated while the APT1.2 transcript is not. Assuming that the APT1.1 and APT1.2 transcripts arising from the APT1::Ex1-i-Ex2:GFP transgenes are of similar abundance as occurs from the WT gene, then the reduced APT1.1 protein levels must be a post-transcriptional effect. This post-transcriptional regulation could be acting on either the primary transcript or the processed mRNA. If the regulation is exerted on the primary transcripts then the targets of this process could be the 5′ UTR/Intron 1; if the regulation acts on the mRNA then the target must be the APT1.1 5′ UTR as this is the only difference between the two transcripts.

**APT1.1 encodes an active enzyme**

Having concluded that APT1.1 is a functional mRNA whereby translation initiation can occur at Exon 1 the next question addressed whether the APT1.1 protein produced from this transcript is an active enzyme. The APT1.1 and APT1.2 ORFs were previously cloned into the pET28a and pET30a vectors in-frame with 6 histidine residues (APT1.1-6xHIS, APT1.2-6xHIS). The recombinant proteins were over-expressed in *E. coli* BL21(DE3)pLysS cells and purified by affinity chromatography using a Ni-NTA resin (Novagen). Purified recombinant protein was than dialyzed in 50 mM Tris-HCl, pH 8.8. The enzymatic activity of APT1.1-6xHIS and APT1.2-6xHIS was measured using a radiochemical assay that monitored the amount of $^3$H-
adenine converted into $^{3}$H-AMP (Table 5). Although the kinetic properties of these recombinant proteins were not determined, the results showed that APT1.1-6xHIS and APT1.2-6xHIS recombinant proteins were very active enzymes. APT1.1-6xHIS had 805-fold greater specific activity than WT extracts; APT1.2-6xHIS had 41-fold greater specific activity than WT extracts.

Table 5. Enzymatic activity for APT1.1-6xHIS and APT1.2-6xHIS recombinant proteins. Recombinant proteins were over-expressed in *E.coli* BL21(DE3)pLysS cells and affinity purified using Ni-NTA resin. Recombinant protein was dialyzed in 50 mM Tris-HCl pH 8.8 and APT activity measured using a radiochemical assay. Results are the average of two individual over-expression and purification experiments.

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**Subcellular localization of APT1.1 and the function of Exon 1**

Given that both APT1 isoforms were very active enzymes, the question arose as to whether the proteins resided in the same subcellular compartments. One possibility was that Exon 1 encodes a transit peptide that compartmentalizes APT1 activity in a cellular location other than the cytosol. Bioinformatic analysis using TargetP predicts Exon 1 is a cleavable transit peptide that localizes APT1.1 to chloroplasts (Emanulesson *et al.*, 2000; Nielsen *et al.*, 1997). A
recent proteomic study of the plastid proteome supports this bioinformatic analysis as APT1.1 was identified in the chloroplast stroma (Sun et al., 2009).

To test this hypothesis 800 base pairs of the presumptive APT1.1 promoter were used to direct the expression of an Exon 1 GFP fusion protein (APT1::Ex1:GFP). This transgene was introduced into WT Arabidopsis and four stable transformants were recovered. The localization of the GFP in leaves of four homozygous lines was observed using confocal microscopy. In leaves of 3-4 week-old plants, the GFP signals were strongest in epidermal cells and were localized in the cytosol. The GFP signals in the underlying mesophyll cells were contained in stationary punctate bodies of approximately 2 µm (Figure 11). GFP localization was also observed in nuclei, as expected, since this fusion protein is below the exclusion limit of the nuclear pore (28 kDa versus 70 kDa exclusion limit) (D’Angelo et al, 2009).
The identity of the punctate signals in mesophyll cells was investigated next. The punctate bodies did not consistently colocalize with chlorophyll autofluorescence, which would have indicated chloroplast localization. This might have been due to weak GFP expression from the native APT1.1 promoter. A count of 20 punctate bodies in the APT1::Ex1::GFP plants showed that these punctate bodies were between 2 to 3 µm in diameter. In addition, these punctate bodies were stationary. This observation rules out the likelihood that these punctate bodies represent mitochondria or peroxisomes since those organelles are smaller (0.5µm to 1µm in diameter), and more dynamic in their movements (Logan and Leaver, 2000; Titorenko and Mullen, 2006).
With the hope that higher expression of GFP might give a more definitive signal a new construct was made using the actin promoter in place of the native APT1 promoter. The open reading frame of APT1.1 was cloned in-frame with two tandem C-terminal GFP ORFs (actP::APT1.1:2GFP). The 5´ UTR of APT1.1 was not included in this construct. The presence of the second GFP ORF increased the size of the fusion protein to 78 kDa so that diffusion into the nucleus should be eliminated (D’Angelo et al., 2009). The actP::APT1.1:2GFP transgene was introduced into WT Arabidopsis and five stable transformants were examined by confocal microscopy. In all T1 lines there was a clear overlap of the GFP signal with chlorophyll autofluorescence (Figure 12). Once again, the stationary punctate bodies of 2 to 3 µm in diameter were observed at the mesophyll layer cells. Interestingly, these signals did not overlap with chlorophyll autofluorescence. As expected, no nuclear GFP signals were detected in these lines indicating that the nuclear localization seen in the APT1::Ex1:GFP plants was due to diffusion.
Figure 12. Confocal imaging of actP::APT1.1:2GFP translational fusions in Arabidopsis leaves. Top panel represents 3.5-week-old leaves, bottom panel represents 4.5 week old leaves of stable transformants. Confocal laser scanning microscope was used to separately capture GFP signals (left) and chlorophyll autofluorescence (middle). A merged image of both fluorescent signals is shown in the right panels.

Overall, these observations suggested that APT1.1 under the expression of the actin promoter is localized to chloroplasts with a secondary undefined punctate-like location. Furthermore, in APT1::Ex1:GFP-containing plant lines, Exon 1 alone was sufficient for targeting to the punctate bodies.

If Exon 1 is a transit peptide for chloroplast import, it might be expected to be proteolytically cleaved once it has accomplished its role in targeting and directing translocation of APT1.1 across the chloroplast envelope. Bioinformatic analysis of APT1.1 identifies Exon 1
as a chloroplast transit peptide that cleaves away all but eight amino acids. To address this issue, the size of the GFP fusion proteins in *APT1::Ex1:GFP* and *actP::APT1.1:2GFP* plants was estimated by immunoblotting. In the *APT1::Ex1:GFP* lines a polypeptide of 33 kDa was expected if Exon 1 was not cleaved (26 kDa GFP + 7 kDa Exon 1); cleavage of Exon 1 would release a 26 kDa GFP polypeptide. *actP::APT1.1:2GFP* plants should synthesize a 78 kDa polypeptide in the absence of cleavage (52 kDa GFP + 26 kDa APT1.1); if Exon 1 was cleaved, a 71 kDa protein would remain.

Immunoblot analysis was carried out on leaf extracts of these transgenic lines. In the four *APT1::Ex1:GFP* lines examined a polypeptide of 28 kDa was detected with the anti-GFP antibody. This is smaller than the complete Exon1-GFP fusion protein and is thus suggestive that APT1.1 is subject to proteolytic cleavage (Figure 10B). However, in the five *actP::APT1.1:2GFP* lines that were similarly analyzed, a polypeptide of 78 kDa was detected, which is consistent with no cleavage (Figure 13). Thus, these lines provided conflicting data regarding Exon 1 being subject to proteolytic cleavage. The fusion of Exon 1 alone with GFP showed that this protein was cleaved. However, when only the ORF was expressed, Exon 1 was not proteolytically cleaved.
Figure 13. Immunoblot analysis of actP::APT1.1:2GFP containing plant lines. Crude leaf extracts (20 ug) of stable transformant Line #1 and WT inflorescences were probed with anti-APT1 antibody (right panels) and anti-GFP antibody (left panels). APT1.1:2GFP detected in lanes 2 and 4 migrated at 78 kDa, endogenous APT1.2 in lane 3 migrated at 24 kDa, and endogenous APT1.1 and APT1.2 in lane 4 migrated at 26 kDa and 24 kDa, respectively. Ponceau S staining shows equal loading of total protein.

Searching for protein-protein interactors of APT1.1 and APT1.2

Knowing that APT1.1 and APT1.2 were localized to different subcellular compartments raised questions regarding the function of each isoform. For example, are APT1.1 and APT1.2 differently localized because they function in metabolic pathways specific for distinct subcellular locations? To address this question, an attempt was made to identify potential protein-protein interacting partners for each isoform in the hopes that these might provide insight into their in vivo roles. Of the numerous methods available to detect protein-protein interactors, I chose to
express epitope-tagged versions of each APT isoform in planta and recover the tagged APT, along with its interactors, by affinity chromatography. To this end, APT1.1 and APT1.2 were cloned under the control of the Cauliflower Mosaic Virus 35S promoter with a C-terminal StrepII epitope tag (Witte et al., 2004). The StrepII tag is a small, eight amino acid peptide that is structurally similar to biotin. Assuming the StrepII tag does not interfere with APT1 protein interactions, the APT1-StrepII fusion can be captured along with its interactors, by chromatography on a strepavidin resin. The interactors can be separated by SDS-PAGE, and the identity of unique polypeptides identified by mass spectrometry.

Four stable APT1-StrepII transformants were used for protein extraction and purification along with a control line expressing an allantoate amido hydrolase (AAH)-StrepII fusion protein (Werner et al., 2008). The polypeptides recovered from APT1.1-StrepII and APT1.2-StrepII expressing lines were separated by SDS-PAGE, and visualized by Sypro Ruby Protein Gel Staining. Only one polypeptide was identified that interacted with APT1.1-StrepII, and was distinct from those in the control sample (Figure 14). This polypeptide was approximately 3 kDa larger than APT1.1-StrepII.
Figure 14. Affinity purification of APT1.1:StrepII and APT1.2:StrepII protein from 4-week-old Arabidopsis leaves. *APT1.1* and *APT1.2* ORFs were cloned into the pXCS-StrepII vector containing the constitutive 35S CaMV promoter and C-terminal StrepII tag (Witte *et al*., 2004). One gram of crude leaf protein was incubated with StrepTactin Macroprep resin (Novagen) to pull down StrepII tagged protein complexes as described in Witte *et al*., 2004. Purified extracts were electrophoresed on a 12.5% SDS polyacrylamide gel and stained using Sypro Ruby Protein Gel Stain (Invitrogen). Lane 1, APT1.1-StrepII purification; Lane 2, APT1.2-StrepII purification; Lane 3, AAH-StrepII purification used as a control for non-specific protein interactions. Black arrows represent StrepII tagged APT1.1, APT1.2 and AAH; white arrow represents unique polypeptide.

To identify the unknown protein, the polypeptide was excised from the polyacrylamide gel. Z. Cheng performed sample preparation, mass spectrometry, and analysis as described in Cheng *et al*., 2009. Briefly, sample preparation involved trypic digestion that cleaves proteins after every arginine residue resulting in a number of smaller peptide fragments. The mass of these fragments was analyzed by mass spectrometry using a Waters Micromass quadrupole time of flight (Q-TOF) Ultima. PEAKS software 3.1 was used to predict the amino acid sequence of these fragments based on their molecular weight and additionally used to search databases for the
identity of the unknown proteins. Using this approach, the unknown polypeptide recovered from SDS PAGE analysis was determined to be the APT1.1-StrepII fusion protein.

To understand the basis of recovering the 3 kDa larger APT1.1-StrepII fusion protein the molecular weights empirically determined by mass spectrometry were compared to the expected molecular weights; no differences were identified. This result signifies that no post-translational modifications occurred to the APT1.1-StrepII fusion protein to account for its 3 kDa shift in size.

Interestingly, the mass spectrometry analysis recovered peptide fragments that covered the entire sequence of APT1.1 except for those amino acids from Exon 1. This could be explained by the amino acid sequence of Exon 1, which is rich in arginine residues. As arginines are the target sites for trypsin digestion, Exon 1 is cleaved in multiple locations producing a number of smaller peptides that are not detected in the mass spectrometry analysis. However, a peptide fragment arising from Exon 2 was identified that contained the amino acid residues DSE at its N-terminus; Exon 1 codes for all three of these amino acids. Despite this unusual anomaly, the initial objective of this experiment was not obtained as no protein interacting partners were identified for APT1.1 and APT1.2 StrepII fusion proteins.

**Complementation of the male sterile phenotype of apt1-3**

Since the investigation of APT1.1 and APT1.2 protein interactors did not reveal any insights into the *in vivo* functional significance of the APT1 isoforms, another means was sought to test their metabolic potential. I tested whether the constitutive expression of either APT1.1 or APT1.2 could rescue the male sterility of *apt1-3*. Previous work by Gaillard *et al.* (1998) showed that expression of the APT1.2 ORF from the 35S promoter could complement the sterility of *apt1-3*. The plan was to do the same for the APT1.1 isoform. As *apt1-3* mutants are male sterile
and not easily transformed by the commonly used floral dip method (Clough & Bent, 1998) the APT1 transgene was introduced into the apt1-3 background by genetic crossing.

The APT1.1 cDNA was cloned under the expression of the actin promoter in a vector expressing glufosinate ammonium resistance (which confers BASTA resistance). The construct was introduced into WT Arabidopsis by the floral dip method. Four stable transformants were recovered and immunoblot analysis was used to confirm APT1.1 over-expression in them (data not shown).

These plants, which were assumed to be carrying one copy of the APT1 transgene, were then crossed to apt1-3 lines. The resulting progeny were all heterozygous for the apt1-3 allele, however, only half were resistant to BASTA treatment. Four resistant F1 plants were allowed to self fertilize to ideally produce a F2 population segregating 9:3:3:1 (9 WT containing the transgene, 3 WT, 3 apt1-3 containing the transgene, and 1 apt1-3). Approximately 50 F2 progeny were germinated on soil and sprayed with BASTA to eliminate all individuals not containing the transgene. The remaining plants were genotyped as described by Gaillard et al 1998 to identify the apt1-3 plants containing a transgene (Figure 15). The apt1-3 allele causes a mutation in a BstNI restriction site associated with the 4th exon. Thus PCR amplification of this region of the APT1 gene followed by digestion with BstNI reveals whether plants carry the WT or apt1-3 allele. PCR was also used to verify the presence of the transgene in the apt1-3 plants.
Figure 15. Restriction mapping of the *APT1* locus to identify homozygous *apt1-3* lines. *APT1* was PCR amplified from genomic DNA for each individual (bottom panel). The PCR product was digested with *Bst*NI (top panel) as the *apt1-3* allele has a single point mutation within this recognition site found endogenously within the *APT1* gene. The following is the restriction fragments expected for each allele: WT 479 bp, 461 bp, 344 bp, 144 bp (lane 2); heterozygotes 940 bp, 479 bp, 461 bp, 344 bp, 144 bp (lanes 1, 3, 4, 5, 6, 7, 8); *apt1-3* 940 bp, 344 bp, and 144 bp (lane 9).

Immunoblot analysis confirmed over-expression of the APT1.1 isoform in positively identified *apt1-3/actinp::APT1.1* plants (Figure 16). The degree of over-expression varied amongst these lines, where approximately 10-fold greater APT1.1 protein relative to WT was observed in the 6-4 line and 100-fold greater expression was observed in the 7-9 line. In all three independent lines the expression of APT1.1 from the actin promoter provided sufficient APT activity for normal development indicating it must function on all substrates that APT1.2 does. This was quite remarkable considering the plastid subcellular localization of APT1.1.
Figure 16. Immunoblot analysis of APT1.1 over-expression in WT and apt1-3. Leaf protein extracts (20µg) were probed with anti-APT1 antibody and bound antibody detected using an anti-rabbit antibody conjugated to HRP. Four independent lines were analyzed in the WT background (6-2, 12-3, 9-3, and 7-2), three independent lines were analyzed in the apt1-3 background (6-4, 9-2, and 7-9).

**Using an amiRNA-based approach for isoform specific gene silencing**

Since constitutive expression of APT1.1 and APT1.2 both restore male fertility to apt1-3 this provided evidence that these isoforms may act on similar substrates in their respective subcellular localizations. An alternative approach to evaluate differences in the functionality of APT1.1 and APT1.2 involved analyzing the phenotypes of mutants that lacked only one isoform. The method chosen to accomplish this was the design and construction of artificial microRNA (amiRNA) lines.

AmiRNAs have become a widely used tool for gene silencing in organisms with sequenced genomes and established transformation methods. In Arabidopsis, the mechanisms of gene silencing mediated by amiRNAs have been well elucidated. The empirical data obtained from these studies has lead to the development a bioinformatic tool that attempts to design amiRNAs with high silencing efficiency for specific transcripts (Schwab et al., 2005). In the case of *APT1.1* and *APT1.2*, amirNAs were designed to target their respective 5´ UTRs (ami1.1 was designed to silence APT1.1 transcripts; ami1.2 to silence APT1.2).
The amiRNAs were designed and cloned as outlined in Schwab et al, 2005. Briefly, the region of interest for targeting was inputted into the online tool WMD2 Web MicroRNA Designer (http://wmd2.weigelworld.org). The output consisted of a list of amiRNAs that would potentially be effective in silencing the region of interest. This list was ranked according to a number of criteria that have been experimentally confirmed as being the most important in endogenous amiRNAs. There is no guarantee that any of the amiRNAs chosen will be effective, nor is there a guarantee that the amiRNA at the top of the list will be more effective than the one at the bottom. However, the creators of this tool recommend to choose the top-listed amiRNAs categorized in the green section as opposed to those in the yellow and red sections. Once the amiRNA of interest has been selected, it is synthesized in a series of three PCRs using four oligonucleotides, the sequences of which are provided by WMD2. The PCR product is designed to be cloned into pRS300, replacing an amiRNA-encoding sequence resident in this plasmid. This sequence will be transcribed from the constitutive 35S promoter when introduced into plants. Five T1 lines for ami1.1 and five T1 lines for ami1.2 were recovered. These lines were followed through to the T3 generation when homozygous lines for all lines were identified.

Immunoblot analysis of ami1.1 lines showed about a 50% reduction in APT1.1, as compared to the WT, in leaves of 4-week-old plants in the most efficient lines (Figure 17B, lines 3, 4, and 5). In line 2, APT1.1 protein levels were slightly higher than that observed in lines 3, 4, and 5. Line 1 showed APT1.1 protein levels similar to WT.

Interestingly, the amount of APT1.2 protein was also affected in the ami1.1-expressing lines. This reduction seemed to correlate with the degree to which APT1.1 was reduced. Lines 3, 4, and 5 showed half the APT1.2 protein observed in WT, while line 2 had one-quarter the
amount of APT1.2. Phenotypically, all the ami1.1 lines showed normal vegetative and reproductive growth despite these differences in APT1.1 and APT1.2 protein levels.

Of the lines expressing the ami1.2, lines 8 and 9 had the greatest reduction in APT1.2 (approximately 75%), in leaves of 4-week-old plants. A reduction of approximately 50% was observed in lines 6, 7 and 10. Unlike the ami1.1 lines where reduction of APT1.1 was associated with decreases in APT1.2, ami1.2-expressing lines did not show reduced APT1.1 protein levels (Figure 17A). Phenotypically, the growth of the ami1.2 lines was indistinguishable from that of WT and ami1.1 lines.

Immunoblot analysis was also used to examine the extent of amiRNA-induced silencing of APT1 in inflorescences. In both ami1.1 and ami1.2 lines, APT1 protein levels were similar to those in WT (Figure 17C). This may explain why all amiRNA containing plants observed were normal in their reproductive growth and as fertile as WT. Thus, these particular amiRNA constructs were partially successful in reducing APT1 protein in leaves, but unsuccessful in silencing either isoform in inflorescences.
Figure 17. Immunoblot analysis of APT1 protein in ami1.1 and ami1.2 lines. Artificial microRNA (amiRNA) transgenes were designed using WMD2 Web MicroRNA Designer (Schwab et al., 2006). Five independent homozygous lines for ami1.1 and ami1.2 were studied. Leaf protein extracts (20µg) for ami1.1 (B) and ami1.2 (A) as well as inflorescences (C) were probed with anti-APT1 antibody (top panel). Ponceau S staining of Rubisco was used to test for equal loading in A and B; anti-SAHH antibody was used in C to test for equal loading.
Chapter 2 - Results

Recently, a novel APT1-deficient mutant carrying a T-DNA insertion between Exon 1 and Exon 2 was identified during a screen for mutants with increased tolerance to chemically induced oxidative stress (D. Falcone, personal communication). This mutant is called oxt1 (oxidative stress-tolerant 1). Unlike previously described APT mutants, such as apt1-3, which have a male sterile phenotype, oxt1 is completely fertile. APT activity in leaves of oxt1 plants was undetectable using a spectrophotometric assay (D. Falcone, unpublished data).

The objective of this part of my research was to determine the basis of the oxt1 fertility. The following questions were addressed in this study: 1) To what degree is oxt1 APT1-deficient? 2) Does apt1-3 have similar stress tolerance to that of oxt1? 3) Is there a correlation between APT activity and fertility? 4) Does the expression of other APT genes change in these APT1-deficient mutants? 5) Does the activity of the two-step adenine salvage pathway change in APT1-deficient mutants?

Is oxt1 APT-deficient?

Moffatt and Somerville isolated the first APT-deficient mutants based on their resistance to 2,6-diaminopurine and found that they all carried mutations in the APT1 gene; all were male sterile to some extent. The degree of APT deficiency in these plants correlated with the severity of the fertility defect. apt1-1 has 10-15% residual APT activity and produces about 2% of the seed that WT plants do. apt1-3 has 1-2% residual APT activity and is completely sterile (Moffatt and Somerville 1988).
oxt1 was DAP resistant suggesting it was also APT deficient (D. Falcone, unpublished data). Although the spectrophotometric assay data indicated a lack of APT activity in leaf extracts, a more quantitative measurement was needed. Thus, the first step in this research was to measure APT activity in oxt1 extracts using the more sensitive radiochemical assay that detects the conversion $^3$H-adenine to $^3$H-AMP. In oxt1, the residual WT activity averaged 3.1% (0.21 ± 0.03 nmol AMP mg protein$^{-1}$ min$^{-1}$) (Figure 18). This is slightly higher than apt1-3 as the average was 2.1% (0.15 ± 0.04 nmol AMP mg protein$^{-1}$ min$^{-1}$), but not statistically significant (p=0.137, n=13, T-Test). The value for apt1-3 matches previously reported values (Allen et al., 2002).
Figure 18. APT enzymatic activity in WT, oxt1, and apt1-3 leaves of 4 week-old plants. APT activity was measured in crude protein extracts using a radiochemical assay that measures the amount of \(^3\)H-adenine converted to \(^3\)H-AMP. oxt1 activity was 3.1%, apt1-3 was 2.1%. Error bars represent standard error of the mean where n=4 for WT, n=13 for oxt1, and n=3 for apt1-3. The difference in APT activity between oxt1 and apt1-3 is not statistically significant (p=0.137, T-Test).

A qualitative means to evaluate APT deficiency involves measuring the length of the primary root of seedlings grown on solid media supplemented with DAP. WT, oxt1, and apt1-3 seeds were germinated on \(\frac{1}{2}\) MS media supplemented with 0.1 mM DAP and root length was
measured after 11 days (Figure 19). Root length measurements were lowest for WT (0.15 ± 0.01 cm, n=12) followed by oxt1 (0.50 ± 0.02 cm, n=12) and apt1-3 (3.16 cm ± 0.2, n=8). These results also indicated that oxt1 is APT-deficient, consistent with the results of the radiochemical assay. However, the APT1 deficiency in the oxt1 mutant as predicted by the DAP growth test is less profound as its tolerance to DAP resembles WT more than it does apt1-3.

Figure 19. Primary root growth analysis of WT and APT1-deficient mutants on 2,6-DAP media. Seeds were germinated on ½ MS media alone (grey bars) or ½ MS media supplemented with 0.1 mM DAP (white bars). At 11 days, the primary root length was measured. The average root length is shown along with the standard error of the mean; n=12 for WT and oxt1, n=8 for apt1-3 and oxt1apt1-3.

Is apt1-3 stress tolerant since it is also APT1-deficient?

The similarity in the APT-deficiency of oxt1 and apt1-3 led to the obvious question of whether apt1-3 was also stress tolerant. This was examined using the same growth test used for
the isolation of the *oxt1* mutant. Seedlings are grown on media supplemented with AT and BSO, which lead to the production of H$_2$O$_2$ and reduced glutathione biosynthesis, resulting in oxidative stress in seedlings (D. Falcone, personal communication). The test was slightly modified to include an initial germination on $\frac{1}{2}$ MS media (WT and *oxt1*) or $\frac{1}{2}$ MS + DAP (*apt1*-3) to identify the homozygous *apt1*-3 seedlings within the segregating F2 population.

WT, *oxt1* and *apt1*-3 seedlings were transferred to $\frac{1}{2}$ MS media supplemented with 2 µM AT and 0.4 mM BSO and their growth was evaluated after two weeks (Figure 20). The WT seedlings were stunted and had chlorotic leaves; *oxt1* seedlings showed no growth inhibition or chlorosis. *apt1*-3 seedlings had evident chlorosis, and their growth was reduced.

![Figure 20](image)

Figure 20. Growth of WT and APT1-deficient mutants on oxidative stress inducing media. WT and *oxt1* seeds were initially germinated on $\frac{1}{2}$ MS media; *apt1*-3 was germinated on $\frac{1}{2}$ MS media supplemented with 0.1 mM DAP to select for homozygous *apt1*-3 seedlings. At 2 weeks seedlings were transferred to $\frac{1}{2}$ MS media containing 2 uM AT and 0.4 mM BSO. Top panel shows representative seedlings after 4 weeks on $\frac{1}{2}$ MS media alone, bottom panel shows representative seedlings grown on AT+BSO media for 2 weeks.

The inhibitory effects of the AT/BSO media on seedling growth are quantified by measuring the lengths of their primary roots (D. Falcone, personal communication). This was not done for this analysis of *apt1*-3 because of the confounding growth effects of the initial DAP.
selection. Although, *apt1-3* mutants are more resistant than WT to the effects of DAP, these mutants are not completely APT1-deficient and as a result their growth is affected by the DAP; this results in stunted growth in comparison to their growth on ½ MS media alone. However, even visual observation of the seedlings showed that *apt1-3* seedlings did not thrive on the AT/BSO media as well as *oxt1*. Thus there is no correlation between APT-deficiency and stress tolerance. Instead there is something distinct about the APT deficiency of *oxt1* that leads to its tolerance.

**What is the correlation between APT activity and fertility?**

Although the difference in APT activity is quite marginal between *oxt1* and *apt1-3* it is quite extraordinary that one genotype is completely fertile and the other is sterile. One could postulate that there is a threshold of APT activity whereby activity above this threshold is sufficient to sustain male fertility and any lower APT level results in male sterility. If there is a fertility threshold then the APT activity of *oxt1* must lie above this level and a decrease in APT activity would diminish fertility. To test this hypothesis, *oxt1* and *apt1-3* plants were crossed to generate *oxt1apt1-3* F1 hybrids which express only one APT 1.2 allele. It was postulated that these plants would have an intermediate APT activity and perhaps be sterile.

In order to interpret the phenotype of *oxt1apt1-3* plants the level of APT activity in different organs needed to be established. A qualitative measure of APT activity was estimated by germinating *oxt1apt1-3* seedlings on ½ MS media supplemented with 0.1 mM DAP. After 11 days of growth, root length measurements were taken to establish residual APT activity (Figure 19). *oxt1apt1-3* roots were slightly longer than those of *oxt1* (0.59 cm ± 0.02 cm, n=8 versus 0.50
cm, respectively) This difference in root length between oxt1 and oxt1apt1-3 is statistically significant (p=0.002, T-Test).

Next, the DAP-resistant seedlings were transferred to soil and their growth observed for up to 7 weeks. After about 4.5 to 5.5 weeks mature siliques had developed on oxt1 and oxt1apt1-3 plants, similar to those that developed on the WT. However, after about 5.5 weeks of growth the subsequent siliques that formed on the oxt1apt1-3 plants were sterile, similar to those on apt1-3 (Figure 21). This “shift” from fertile to sterile was indeed a phenomenon of the F1 heterozygous mutant line and not a result of pest infestation or growth chamber conditions as this shift was apparent each time these plants were grown.
Figure 21. A seven-week-old oxt1apt1-3 plant. Photographed is the primary shoot of a representative oxt1-apt1-3 plant. The initial siliques have dried out at this time and are all fertile; the later siliques are all sterile.

To determine how consistent this fertility shift was in oxt1apt1-3 line thirteen individuals were analyzed; these individuals were derived from multiple crosses between at least three oxt1 plants onto at least three different apt1-3 female recipient plants. The siliques on the primary shoot of thirteen 7-week-old plants were counted and recorded as to whether each was fertile or
sterile. On the primary shoot, upwards of the first 10 siliques were fertile, or approximately 20-40% of the total number of siliques (Figure 22A). All subsequent siliques were sterile. Interestingly, these plants would produce many more sterile siliques than fertile siliques, at times numbering upwards of 60.

This fertility shift was also observed on the secondary branches. However, the timing of the shift was more variable on the secondary branches than it was on the primary shoot. For any given secondary branch, 10-60% of the siliques were sterile. Moreover, it was not uncommon to see fertile siliques intermixed amongst the sterile siliques along these branches (Figure 22B).

Together these observations are consistent with the model that there is a threshold of APT activity that is necessary for fertility in Arabidopsis and the reduction in APT activity in oxt1apt1-3 plants relative to the oxt1 homozygote mutants leads to sterility. During vegetative growth on ½ MS media supplemented with DAP the oxt1apt1-3 seedlings had root lengths that were more comparable to oxt1 than they were to apt1-3. During the early stages of reproductive growth fertile siliques were observed on oxt1apt1-3 plants like that seen on oxt1, and later a shift to sterility occurred, similar to the phenotype of the apt1-3 mutant.
Figure 22. Analysis of fertile and sterile siliques on primary and secondary shoots of oxt1apt1-3 plants. Thirteen oxt1apt1-3 individuals were used to determine the extent of fertility and sterility displayed on the primary shoot (A) and secondary shoot (B). Left panels: fertile siliques are represented as black bars, grey bars represent number of sterile siliques for each individual. Right panels: percent of fertile siliques.

The next question addressed was whether the shift to sterility seen in oxt1apt1-3 plants was accompanied by a decrease in APT activity. To test this APT activity was measured by the radiochemical assay in protein extracts of inflorescences of WT, oxt1, apt1-3 and oxt1apt1-3 plants. Extracts were prepared from the first inflorescences (those that gave rise to the early fertile siliques) and from later inflorescences (those that gave rise to sterile siliques). The early buds and late buds of WT, oxt1, and apt1-3 plants had consistent APT activities, in each case (Figure 23). Surprisingly, the late, sterile buds of the oxt1apt1-3 plants had a significant increase in APT activity as compared to the early buds (p < 0.05, T-Test). Across the two individual
experiments there was a consistent increase of 4.6% in APT activity in sterile buds over fertile buds. The APT activity in the early, fertile buds of oxt1apt1-3 is equivalent to that of oxt1, but in late, sterile buds the APT activity increases. This observation provides evidence that in the case of oxt1 and oxt1apt1-3, APT activity is not clearly correlated with sterility. Thus, the hypothesis concerning a threshold of APT1 activity can be rejected.
Figure 23. APT enzymatic activity in early and late buds of oxt1apt1-3 plants. Crude protein extracts for WT, oxt1, apt1-3, and oxt1apt1-3 buds were assayed for APT activity at two stages of development. The early stage involved collecting the first initial buds produced by the plant, the late stage defines buds collected after the first observation of sterile siliques. The two sets of data for each experiment is shown where the WT activity was 12.39 and 12.75 nmol of AMP/mg of protein/min (A); and 28.71 and 16.76 nmol of AMP/mg of protein/min (B). The asterisk marks significant changes between early and late buds where p=0.004 (A) and p=0.02 (B) using Student’s T-Test.
Does the expression of other APT genes change in APT1-deficient mutants

The unexpected increase in APT activity in the sterile buds of oxt1apt1-3 plants led to the consideration that the fertility differences in oxt1 and aptl-3 plants might be a result of how these two APT1 alleles signal another component of cellular metabolism. Perhaps oxt1 mutants activate a compensatory activity to alleviate male sterility associated with APT1-deficiency? One possibility was that oxt1 was able to activate another APT gene. Allen et al (2002) showed that APT1, APT2 and APT3 have similar apparent K_m values for adenine. However APT1 has the highest V_max of the three isoforms (APT1 V_max = 9.6; APT2 V_max 0.31; APT3 V_max 0.18; units are expressed in nmol of AMP mg of protein^{-1} min^{-1}). So the other APTs that are naturally expressed only in flowers (Zhang, 2000) might contribute to the increase in APT that is observed in the oxt1apt1-3 flowers.

Thus, I investigated whether the oxt1 APT1-deficiency might signal a transcriptional response from APT2 or APT3. RNA was extracted from the late buds of WT, oxt1, aptl-3, and oxt1apt1-3 and APT2 and APT3 transcript abundance analyzed by semi-quantitative RT-PCR. Each primer’s linear range was determined using WT cDNA as a template. For the APT2 primer set the linear range was between 32 and 36 cycles; for the APT3 primer set it was between 28 and 36 cycles. Initially, 34 cycles was chosen for RT-PCR, however, there was concern that the signal strength of each DNA fragment may have been saturated and closing in on the higher end of the linear range (Figure 24). The cycle number was than lowered to 30 cycles for further analyses.
Figure 24. Linear range testing of APT2 and APT3 primer sets for RT-PCR analysis. WT cDNA template was PCR amplified for 20 through 36 cycles and electrophoresed on a 1% TAE agarose gel. The gel image was digitized and each DNA fragment quantified, in nanograms, using spot density analysis. The nanograms of cDNA synthesized was plotted against the cycle number on a semi-log graph. 34 cycles was chosen as the linear range for APT2; 30 cycles was chosen for APT3.
At 30 cycles, the RT-PCR results showed a modest decrease in $APT2$ transcript abundance for $oxt1$, $aptl$-3, and $oxt1aptl$-3 relative to WT, but this was not substantially different (Figure 25). APT3 transcript abundance was equal across all the samples (Figure 25). These observations suggest that APT2 and APT3 are not transcriptionally responsive to APT1-deficiency, and are not likely contributing to the diverse phenotypes seen among $oxt1$, $aptl$-3, and $oxt1aptl$-3 mutants. However, a translational activation of some sort cannot be ruled out.
Figure 25. RT-PCR analysis of APT2 and APT3 in WT and APT1-deficient mutants. RNA was extracted and cDNA synthesized from late buds of two individuals per line. F1 represents oxt1apt1-3 line. RT-PCR was performed at 34 cycles (A) and 30 cycles (B). Actin was used to test for equal loading of cDNA.

**Does the activity of the two-step adenine salvage pathway change in APT1-deficient mutants**

Given that the other APT genes of the primary salvage pathway do not appear to account for the fertility differences of oxt1 and apt1-3 mutants, I next considered the possibility that the secondary salvage pathway was more active in oxt1. This pathway involves the sequential action of adenine phosphorylase and ADK. Adenine phosphorylase converts adenine to adenosine, which is subsequently phosphorylated to AMP.

ADK activity was assayed in leaf extracts to determine if the two-step pathway is differentially active in oxt1 versus apt1-3 plants. The radiochemical ADK assay monitors the rate
of conversion of $^3$H-adenosine to $^3$H-AMP in protein extracts. Extracts were prepared from leaves of 4-week-old plants. In \textit{oxt1} the ADK activity was 2.07-fold greater than that in WT; in \textit{aptl-3} ADK activity was 1.67-fold greater (Figure 26). However, neither increase was statistically significant ($oxt1$ \(p=0.07\), \textit{aptl-3} \(p=0.05\)). These data showed that ADK activity might be slightly higher in \textit{oxt1} and \textit{aptl-3} relative to WT, but was not significantly different in \textit{oxt1} and \textit{aptl-3} (\(p=0.30\), T-Test). However, since ADK activity was similar in \textit{oxt1} and \textit{aptl-3} the two-step adenine salvage pathway cannot account for the phenotypic differences observed in these mutants.

![Figure 26](image_url)

Figure 26. ADK enzyme activity in leaves of WT and APT1-deficient mutants. ADK activity was measured in crude protein extracts using a radiochemical assay that measured the amount of $^3$H-adenosine to $^3$H-AMP. Error bars represent standard error of the mean, where \(n=3\) for WT, \textit{oxt1}, and \textit{aptl-3}. The difference between \textit{oxt1} and \textit{aptl-3} is not statistically significant (\(p=0.30\), T-Test).
Discussion

At present, Arabidopsis APT1 is unique in comparison to the APT genes of other species as it is the only APT locus for which there is experimental evidence for two gene models: one with 7 exons (APT1.1) and another with 6 (APT1.2). To date, all other annotated APTs encode only one polypeptide that has a high amino acid sequence similarity to APT1.2; the sizes and locations of the introns and exons are also conserved amongst these genes. It is possible that all organisms express an APT1.1 but the genetic resources are not yet available to detect this expression. Thus although Arabidopsis may be the only plant documented to have a dual isoform APT1 locus to date, the study of these dual gene products will further our understanding of the role of this enzyme activity in all plants.

The NCBI EST database has 18 entries for APT1.1 transcripts and 53 records for APT1.2. The APT1.2 ESTs were recovered from all tissue types whereas the APT1.1 ESTs have only been isolated from either leaves of young seedlings or floral organs. The predicted gene models derived from these ESTs suggest that the APT1.1 and APT1.2 transcripts arise from distinct promoters. This conclusion is based on two observations: both transcripts are capped and they have unique transcriptional start sites and 5’ UTR sequences. Their specific promoter regions have yet to be delineated.

Earlier studies of APT1 expression (which is now designated APT1.2) located a promoter activity in the 950 bp upstream of the first methionine in Exon2. This stretch of nucleotides directed constitutive expression of a downstream β-glucuronidase reporter gene similar to the pattern of APT1.2 (Smith, 1997). The “native APT promoter” used in the transcriptional studies outlined in this thesis are the 800 bp upstream of the initiating methionine of Exon 1.
The first objective was to address the expression of *APT1.1* and *APT1.2* transcripts. Semi-quantitative RT-PCR was used to measure transcript abundance in leaves from 4-week-old plants and older inflorescences. These data showed that *APT1.1* and *APT1.2* transcript levels are similar in both tissues. This result is quite extraordinary considering that only a few ESTs are reported for *APT1.1*, while *APT1.2* has over double the number of ESTs.

The basis for this discrepancy may be related to the difficulty to recover ESTs that include the 5´ end of this transcript. In the course of my research, I attempted to recover the full-length cDNA for *APT1.1* using RLM-RACE and high-temperature cDNA synthesis. I was successful in isolating a full-length cDNA for *APT1.2*, but not for *APT1.1* using RLM-RACE. As this method is dependent on the availability of a 5´-capped end to mediate RNA ligation of an adaptor sequence, any significant secondary structure could inhibit this reaction. I attempted to reduce potential secondary structure using high temperature cDNA synthesis at 55°C, and three primers were designed to cover the entire 5´ UTR region of *APT1.1* (Appendix 3) The only primer that did not lead to synthesis of a RT-PCR product was the one that anneals at the 5´ terminus of the UTR.

To further investigate any significant secondary structure in the 5´ UTR, computational modeling of the mRNA structure was performed. This modeling revealed a possible RNA secondary structure at the 5´ UTR of *APT1.1* that contains a region with a substantial stem-loop located at the terminal 5´ end (Appendix 2; Y.Li, personal communication).

Concurrently, a comparative analysis of the *APT1.1* 5´ UTR with characterized metabolite-binding regions was also carried out. These metabolite-binding regions are more commonly known as riboswitches and have been elucidated in prokaryotes with two cases
reported in eukaryotes (Breaker et al., 2008). Interestingly, a thiamine pyrophosphate (TPP)-sensing riboswitch has been found in bacteria, fungi, and plants yet the mechanism of gene repression is different in these species. In bacteria, TPP binds the mRNA inducing a conformational change in the mRNA that blocks the ribosome binding site, leading to reduced expression (Winkler et al., 2002). In fungi, TPP binds the mRNA in the 5’ UTR inducing an alternative splicing event that leads to an upstream open reading that acts as a decoy for translation and reduces expression of this mRNA (Cheah et al., 2007). In plants, TPP binds within the 3’ UTR of the mRNA inducing an alternative splicing event that leads to changes in the 3’ UTR length and sequence whereby mRNA stability is affected leading to reduced expression (Wachter et al., 2007).

The comparative analysis of the secondary structure in APT1.1 was performed against known riboswitches. This analysis showed no similarity between known riboswitches and the RNA structure predicted in the APT1.1 5’ UTR (Y. Li, personal communication). However, this analysis does not rule out the possibility of a riboswitch or a metabolite-binding region within the 5’ UTR. As this analysis was performed using prokaryotic sequences, and only a few riboswitches have been elucidated in eukaryotes, it could be possible that these regulators of gene expression have evolved differently in eukaryotes. Future experiments such as in-line probing assays to test conformational changes in RNA secondary structure upon binding of metabolites may provide valuable insight into whether for example, adenine or cytokinins can provoke this event (Li and Breaker, 1999).

With the evidence that both APT1.1 and APT1.2 mRNAs are synthesized in vivo, the next question addressed whether both transcripts were translated in vivo. Immunoblot analysis of leaves and inflorescences showed that APT1.2 is abundant in both tissues. In leaves, the APT1.1
level was approximately one-tenth that of APT1.2, and below the level of detection in inflorescences. Thus, the protein levels do not match transcript abundances estimated by the RT-PCR analysis suggesting that *APT1.1* may be subject to post-transcriptional regulation, while *APT1.2* is constitutively expressed. Interestingly, these expression patterns are in agreement with the analysis of alternative first exons by Chen *et al* (2007). This group showed that most alternative first exon events in rice and Arabidopsis involve a short isoform that is constitutively expressed, and a longer isoform that is spatially/temporally/developmental expressed. Although, it is unknown what induces expression of APT1.1 this model is consistent with the APT1 isoforms as APT1.2 is expressed in all tissues.

**What is the possible basis for the post-transcriptional regulation of APT1.1?**

In WT, it was found that the equal transcript abundance of *APT1.1* and *APT1.2* did not correlate with the amount of protein detected on immunoblots as APT1.2 was much more abundant than APT1.1. As the only difference between these mRNAs is their 5´ UTR regions, it was hypothesized that the 5´ UTR of *APT1.1* must be the target of post-transcriptional regulation resulting in its lower expression. The secondary structure of *APT1.1* alluded to earlier in this discussion may be involved in this regulation. While this may be true, a second target of post-transcriptional regulation was elucidated by two GFP fusion constructs.

Two constructs were initially designed to test the expression of Exon 1 *in vivo*. The first construct consisted of the first 800 base pairs of the APT1.1 promoter including the 5´ UTR and Exon 1 fused in-frame with GFP (*APT1::Ex1:GFP*). The second construct was identical to *APT1::Ex1:GFP* but included Intron 1 and Exon 2, fused in-frame with GFP (*APT1::Ex1-i-Ex2:GFP*).
Immunoblot analysis of *APT1::Ex1:GFP* detected an abundant 28 kDa Ex1-GFP fusion protein. This was not expected, as *in vivo* APT1.1 expression from this promoter and 5′ UTR produces a low level of APT1.1. These results suggested that perhaps the 5′ UTR alone was insufficient in regulating translation of the Ex1-GFP fusion protein; the same may be true for the WT gene *in vivo*.

Two polypeptides were expected to arise from the *APT1::Ex1-i-Ex2:GFP* transgene: the 36 kDa Exon1-Exon2-GFP fusion protein and the 29 kDa Exon2-GFP fusion protein. Surprisingly only the 29 kDa polypeptide was detected. Although the abundance of Exon1-Exon2-GFP and Exon2-GFP fusion proteins did not reflect that found in the corresponding proteins arising from the native gene, it can be said with some confidence that only Exon2-GFP is synthesized and Exon1-Exon2-GFP is once again subject to regulation. Both transcripts that code for these proteins were detected by RT-PCR. Preliminary semi-quantitative RT-PCR analysis indicated that the two transcripts were equally abundant, consistent with transcription from the native gene (Y. Li, personal communication). Thus, APT1.1 is subject to post-transcriptional regulation and the cis elements required for this regulation are present in the *APT1::Ex1-i-Ex2:GFP* transgene.

These results suggest that Exon 1 must be upstream and colinear with Intron 1 and Exon 2 for the post-transcriptional regulation of APT1.1 (i.e. this regulation affected transcripts arising from *APT1::Ex1-i-Ex2:GFP* and WT *APT1* genes but not from the *APT1::Ex1:GFP* transgene). The commonality between *APT1::Ex1-i-Ex2:GFP* plants and WT is the co-linearity of Exon1, Intron 1, and Exon 2. The cis nature of the factors controlling this regulation is further supported by the fact that the *APT1::Ex1:GFP* construct was expressed in a WT background that contains two functional copies of the APT1 gene, yet no effect on translation of the WT alleles from Exon
APT1.1 was detected in these plant lines. If this is the case, then the target of post-transcriptional regulation could be the primary mRNA. Future work will address the effect of removing Intron 1, to further distinguish the contribution of this intron and the 5’ UTR of APT1.1 in its post-transcriptional regulation. As such, work has begun to clone the genomic fragment of APT1 beginning with the 800 base pair promoter region and ending with the terminal exon fused to GFP (APT1::Ex1-Ex7:GFP). The expression of the APT1.1 and APT1.2 polypeptides arising from this transgene will be monitored by immunoblotting with an anti-GFP antibody. These APT1::Ex1-Ex7:GFP transgenic plants will be a control for another set of transgenic plants that do not contain the first intron (APT1::Ex1-ΔIntron1-Ex7-GFP).

**APT1.1 is an active enzyme localized to chloroplasts**

It was next addressed whether the APT1.1 polypeptide is an active enzyme. Recombinant APT1.1 and APT1.2 fused with a 6xHIS tag were over-expressed in *E.coli*, purified, and dialyzed into Tris-HCl buffer. Using a radiochemical assay their activities on adenine substrates were tested relative to the APT activity in WT crude extracts (expressed as nmol AMP mg\(^{-1}\) min\(^{-1}\)). In a preliminary kinetic analysis APT1.1 had 805-fold greater APT activity; APT1.2 showed 41-fold greater activity. As these data are preliminary, it can only be concluded that APT1.1 is an active enzyme. Future work will determine the apparent binding affinities of each isoform to adenine and cytokinins, in order to provide a meaningful idea of what substrate each prefers and its efficiency in metabolizing each of these.

The subcellular localization of each APT1 isoform was investigated to discern if Exon 1 functions as a transit peptide as predicted by multiple subcellular localization programs. To test this two different transgenic plant lines were studied: those expressing the APT1::Ex1:GFP transgene and those expressing actP::APT1.1:2xGFP. The GFP signal from APT1::Ex1:GFP
transformants was observed in the cytosol of leaf epidermal cells, as well as in punctate bodies and nuclei of mesophyll cells.

Two issues arose from these results. First, the nuclear GFP signal is likely due to diffusion since the size of the GPF fusion protein is below the exclusion limit of the nuclear pore (28 kDa versus 70 kDa; D’Angelo et al., 2009). The second issue was the overall low expression of GFP in these lines, which made it difficult to tell if it co-localized with the chlorophyll autofluorescence of the chloroplast. To overcome this problem, the native APT1 promoter was substituted with the constitutive actin promoter to drive the expression of an APT1.1 ORF with two tandem GFP ORFs (actP::APT1.1:2xGFP). These lines eliminated the nuclear localization, as this fusion protein was larger than the size exclusion limit (78 kDa versus 70 kDa). The stronger expression from the actin promoter clearly showed co-localization of GFP with chlorophyll autofluorescence. Additionally, the localization of APT1.1 to punctate-bodies was observed again. Taken together these observations implicated Exon 1 as a transit peptide. However, it remained unclear as to whether Exon 1 was proteolytically cleaved in this process.

Immunoblots of leaf extracts of APT1::Ex1:GFP transformants showed that Exon1 was cleaved from GFP. This construct was amplified from genomic DNA and only the genomic fragment of Exon 1 upstream of Intron 1 was used; the portion of “Intron 1” that contains the first six amino acids of APT1.2 were not included. This transgene was constructed to determine whether translation initiation occurred at Exon 1 in vivo and the localization of the resulting polypeptide. It remains unclear as to whether the fusion of Exon 1 splice site with GFP resulted in a false cleavage site. The subcellular prediction tool, MitoProtII, suggests that APT1.1 has a transit peptide the end of which lies within the 6 amino acid region left out of this construct. To restore the native amino acid sequence the full-length APT1.1 ORF was used in the
actP::APT1.1:2xGFP plant lines. Immunoblots of leaf extracts from these transgenic lines showed that in the absence of Intron 1 splicing, Exon 1 was not cleaved from APT1.1. The discrepancy as to whether APT1.1 is proteolytically cleaved or not, is important since it impacts two other questions: the method by which an un-cleaved protein is imported into chloroplasts, and second, the degree of post-transcriptional regulation on APT1.1.

With respect to the chloroplast import issue, it was recently estimated that 86% of 1325 nuclear-encoded plastid proteins are translocated across the chloroplast envelope via the TOC-TIC machinery (Zybailov et al., 2008). All these proteins possess an N-terminal cleavable transit peptide that is released from the preprotein after translocation across the chloroplast envelope (Oblong and Lamppa, 1992). However there is growing evidence that some nuclear-encoded chloroplast proteins do not have cleavable transit peptides. One example is Tic32 that contains a small, N-terminal transit peptide sufficient for chloroplast targeting to the inner-membrane yet is not cleaved (Nada and Soll, 2004). The import method for Tic32 has yet to be elucidated, however it has been determined that inhibition of Toc159, Toc34, and Toc75 receptors does not inhibit the translocation of Tic32 into chloroplasts. Inaba and Schnell (2008) postulate in a recent review that although the import of a few proteins have been shown to be independent of the most characterized TOC-TIC proteins (i.e. Toc159), an alternative pathway may use other characterized TOC translocons. The results of a recent Arabidopsis plastid proteome analysis supports the existence of an alternative import method. Using mass spectrometry 3800 proteins were identified as part of the plastid proteome, but only 67% of these proteins were predicted by TargetP to have cleavable transit peptides (Kleffmann et al., 2004). In vitro chloroplast import assays could be used to determine whether APT1.1 is dependent or independent of Toc159, Toc34 and Toc75 machinery for import.
If APT1.1 is proteolytically cleaved when it is imported into plastids then the issue of its post-transcriptional regulation will need to be reviewed. The cleavage of APT1.1 would result in a peptide of 24 kDa, equal to that of APT1.2. Thus the intensity of 24 kDa polypeptide band visualized on immunoblots of crude leaf extracts would reflect the sum of APT1.2 and the processed APT1.1. To date, immunoblot analysis of over-expressed APT1.1 ORF and APT1.1 fusion proteins has always detected the full-length un-cleaved protein. This evidence supports the notion that APT1.1 is post-transcriptionally regulated and not subject to proteolytic cleavage. However these proteins are being expressed from transgenes residing outside the native genomic context of APT1 and it could be argued that this may affect their expression. Moreover, their expression from a cDNA removes numerous potential regulatory steps that may be a component of the processing. Future tests could involve constructing transgenes harbouring mutations at the predicted cleavage sites by TargetP and MitoProtII. If these predicted sites are indeed necessary for proteolytic processing of APT1.1 than an increase of APT1.1 protein abundance should be detected by immunoblotting.

**Punctate localization of APT1.1 may be related to Chloroplasts**

The localization of APT1.1 in punctate-bodies led me to consider its possible localization in mitochondria or peroxisomes because of their similarity in size and shape. However, these organelles are well known for their dynamic movement (Logan and Leaver, 2000; Titorenko and Mullen, 2006) and this movement was not observed for any of the punctate bodies in *actP::APT1.1:2GFP* plants. Additionally, the estimated size of the GFP punctate bodies was 2µm in diameter, whereas mitochondria and peroxisomes are normally 0.5 to 1µm in diameter.
The question arose as to whether these punctate bodies could reside in the chloroplast or perhaps be proplastids. This was considered because the majority of the punctate-bodies were localized adjacent to chloroplasts and detected only in optical sections where mesophyll cells were in focus. A literature search has identified two distinct proteins that are localized to chloroplasts and report similar confocal images of GFP-fusion proteins to those obtained in this study.

Plastid Envelope DNA binding protein (PEND) is a chloroplast-localized protein that is specifically targeted to the plastid nucleus, which is also known as the plastid nucleoid (Terasawa and Sato, 2005). Plastid nucleoids harbour chloroplast DNA that codes for proteins involved in photosynthesis, gene expression, or essential to plastid metabolism. PEND aids in anchoring the plastid nucleoid to the stromal side of the inner envelope. Interestingly, Myouga et al., 2008 showed that Iron Superoxide Dismutase 3 (FSD3) co-localizes with PEND in plastid nucleoids. Both PEND-GFP and FSD3-GFP confocal images show punctate-like structures, similar to those seen in the plants expressing APT1.1-GFP. In all three cases, there was no clear overlap of the punctate bodies with chlorophyll autoflorescence. Thus, it is possible that APT1.1 is being localized to non-chlorophyll containing regions of the chloroplast, similar to PEND and FSD3.

Immunogold labeling was performed on leaves of 3-week-old aptl-3 plants containing an actp::APT1.1 transgene to further clarify the localization of APT1.1 (Figure 27). The epitope was detected using a secondary antibody conjugated to 10 nm gold balls, via transmission electron microscopy. The results have yet to be analyzed quantitatively but it was quite evident that the gold balls were concentrated in bodies located within chloroplasts (B. Moffatt, personal
Interestingly, these bodies appeared to be located near the edges of the plastids in regions with few thylakoids. To date these bodies have only been observed in the APT1.1 overexpressing line and not in the WT, but further analysis is needed to confirm this conclusion. The localization of these bodies may well correspond to the punctate GFP signals observed using confocal microscopy as they both reside in regions of the chloroplast with little or no chlorophyll.

However, one discrepancy between the two detection methods is the predicted size of the punctate-bodies. Using confocal microscopy the GFP-tagged punctate-bodies were estimated to be 2 µm in diameter, whereas those imaged using the TEM were under 1 µm in diameter. One possible explanation for this could be that the intensity of the GFP fluorescence is contributing to its larger appearance by confocal microscopy.

Figure 27. TEM images of aptl-3 actP::APT1.1 containing plant lines. Immunogold labeling of APT1.1 resulted in 10 nm gold beads localized to chloroplasts within peripheral located bodies. Chl (Chloroplast). Inset is a magnified image of the marked area. Tissue preparation and imaging performed by B. Moffatt. Scale bar represents 0.2µm.

The punctate localization of APT1.1 is perhaps of greater interest than its identified chloroplast localization. As observed in both GFP lines, expression of Exon 1-GFP fusion from
the native $APT1$ promoter and over-expression of the full-length APT1.1 consistently showed this punctate signal. If these punctate bodies are representative of a specific region within chloroplasts then the $\textit{APT1}::\text{Ex1:GFP}$ plant lines give an accurate representation of of APT1.1’s predominant localization in WT plants. Thus, the co-localization with chlorophyll auto fluorescence seen in $\textit{actP}::\text{APT1.1:2GFP}$ may be a result of over-expressed protein and not be representative of localization under native conditions.

**Communication between de novo synthesis and purine salvage activities**

With the localization of APT1.1 determined, future studies will need to investigate the purpose behind compartmentalizing APT1.1 within plastids. In non-plant cells de novo synthesis and purine salvage are both localized in the cytosol (An et al., 2008). This is contrary to plant cells that situate de novo synthesis within chloroplasts, and the purine salvage pathway in the cytosol. As a result, non-plant cells have the majority of their adenylate pool in the cytosol, whereas in plant cells the adenylate pool is equally divided between the chloroplast and cytosol. This was shown in leaf protoplasts of wheat where 47% of adenylates were found in the chloroplasts, 44% in the cytosol, and 9% in the mitochondria (Stitt et al., 1982). It could be argued that localizing APT1.1 to chloroplasts may mediate some kind of metabolic cross talk or coordination between the two purine biosynthesis pathways. Alternatively, the purine salvage pathways may have evolved to be targeted to chloroplasts to assist in sustaining its nucleotide pool in an energy efficient manner, as it does in the cytosol of both plant and non-plant cells.

Coordinating de novo synthesis and purine salvage could result in regulation of de novo synthesis by purine salvage. For example, in yeast, fusions of de novo purine gene promoters to lacZ have a 3-18 fold lower expression in the presence of adenine (Daignan-Fornier and Fink,
A potential role for APT in this system was suggested to involve metabolism of adenine via APT activity, thereby removing repression from the \textit{de novo} synthesis pathway. This mechanism would further contribute to the salvage of adenine to AMP. However, the repression of adenine on \textit{de novo} synthesis has yet to be demonstrated outside of yeast. Rather, in plants it was shown that adenylates arising from either \textit{de novo} synthesis or purine salvage act as negative regulators that feedback to the first enzyme (amidophosphoribosyltransferase; ATase) in \textit{de novo} synthesis (Reynolds \textit{et al.}, 1984). Thus, in both yeast and plants purine metabolites may affect the regulation of \textit{de novo} synthesis. The salvage pathway may supplement the adenylate pool more in these situations, in a less energetically demanding manner. As the synthesis of purine nucleotides via the \textit{de novo} route is energetically expensive, acknowledging the output of a less energy-demanding pathway would assist the cell in synthesizing purine nucleotides in the most energy efficient manner.

Apart from regulation, the \textit{de novo} purine biosynthesis enzymes may be co-localized with purine salvage enzymes to coordinate metabolism during development. This would be beneficial, as nucleotide pools need to be sustained during times of cell growth and differentiation. Two studies have followed the metabolic fate of labeled purine precursors during developmental processes in plants. Both studies highlight the importance of the purine salvage pathway in growing and sustaining nucleotide pools early in development.

Brassica embryos early in development incorporate up to 75\% of [8-\textsuperscript{14}C] adenine into adenine nucleotides with a gradual decrease to 46\% incorporation during later developmental stages (Ashihara \textit{et al.}, 2008). The next largest portion of adenine was recovered in RNA, with 15-20\% being incorporated throughout the four developmental stages observed. In white spruce,
a similar pattern of high purine salvage activity occurs during shoot organogenesis (Stasolla et al., 2006). These authors found that under shoot inducing conditions (i.e. in the presence of an exogenous supply of cytokinins) the uptake of [8-14C] adenine increased up to 2.5-fold over non-shoot inducing conditions, at the peak period (day 20) when shoots began to form. Furthermore, APT activity increased 1.5-fold under shoot-inducing conditions over non-shoot-inducing conditions, throughout the 30-day observation period. Additionally, APT activity was consistently higher than that of the secondary salvage pathway. Thus, these studies show that purine salvage activity contributes largely to nucleotide pools and are most active during early stages of growth and development. However, none of these studies followed the metabolic fate of de novo synthesis precursors in comparison to the salvage activities.

One study followed both de novo and salvage purine metabolic pathways in tobacco cells undergoing programmed cell death (PCD) (Stasolla et al., 2005). NO and H2O2 production were induced in tobacco cell culture to initiate PCD. After PCD induction the metabolic fates of [8-14C] adenine (purine salvage precursor) and [8-14C] AICAR (de novo synthesis precursor) were monitored for 8 hours (Stasolla et al., 2005). After 2 h, APT enzymatic activity reached its maximum and the incorporation of adenine into the nucleotide fraction was higher (40% incorporation) than AICAR (12% incorporation). After 4 h, the incorporation of adenine declined to 13%, while the incorporation of AICAR increased to 18%. In addition, at 4 h a large fraction of [8-14C] AICAR was shifted from degradation products to the nucleotide fraction, representing the time at which de novo synthesis initiates. At no point throughout the time course was there a significant incorporation of either adenine or AICAR into the nucleic acid fraction. This study was one of the first to demonstrate the early activation of purine salvage and later initiation of de novo synthesis. This showed the importance of coordinating purine nucleotide biosynthesis
pathways to provide quick and constant growth of nucleotide pools helping to sustain them through periods of high demand.

Based on the literature, a metabolic role for APT1.1 within chloroplasts could be proposed. The previous studies all highlight the importance of adenine salvage during intense growth periods as the majority of adenine was recovered in nucleotide fractions. It is possible that APT1.2 is insufficient to maintain nucleotide pools in the cytosol and the chloroplasts. If \textit{de novo} synthesis initiates later than the purine salvage pathway, as reported in tobacco cells, APT1.1 may function in sustaining nucleotide pools in the chloroplast prior to start of \textit{de novo} synthesis. Thus, one possibility is that APT1.1 and APT1.2 might be functionally similar enzymes involved in sustaining nucleotide pools.

**Purine Salvage and CK metabolism in chloroplasts**

A second possible role for APT1.1 in chloroplasts could relate to cytokinin metabolism. In plants, plastids are the predominant site of cytokinin biosynthesis (Sakakibara \textit{et al}., 2006). Localization of the adenosine phosphate-isopentenyltransferase (IPT) genes that catalyze the synthesis of cytokinins has been studied in Arabidopsis using GFP protein fusions. It was shown that half of the IPT gene family (IPT1, 3, 5, 8) are localized to plastids (Kasahara \textit{et al}., 2004). The IPT genes catalyze the synthesis of cytokinins by transferring the isoprenoid side chain to either ADP or ATP. The isoprenoid side chain can arise from two pathways: the plastid-localized methylerythriol phosphate (MEP) pathway or the cytosolic mevalonate (MVP) pathway (Rohmer, 1999). Kasahara \textit{et al} (2004) showed that only 20\% of the labeled $^{13}$C precursors for the MVP pathway were incorporated into tZ and iP cytokinins; in the MEP pathway 90\% of the $^{13}$C precursor was recovered for the same cytokinins. These results show that MEP is the
predominant pathway for the synthesis of isoprenoid side chains as well as demonstrate that plastids are a major site of CK biosynthesis in plants.

As ADP/ATP are essential precursors for the biosynthesis of cytokinins, APT1.1 may be localized to the plastids to support the synthesis of these molecules. Cytokinins are important hormones in plant growth and development, thus the synthesis of this precursor by a rapid and efficient pathway is essential. Alternatively, APT1.1 may function in CK homeostasis by interconverting cytokinin bases to their corresponding less active nucleotide forms. These molecules can be converted back to their active base form, reducing the need for their constant de novo biosynthesis (Kurakawa et al., 2007).

To gain insight into the function of APT1.1 and APT1.2, I sought to identify their protein interactors. The objective was to find potential protein-protein interactors that would provide insight into the function or metabolic pathway in which each isoform participates. Furthermore, it was anticipated that this approach might confirm the subcellular localization of each isoform depending on the identity of the protein(s) it interacted with. The method chosen involved over-expressing each APT1 isoform fused to the eight-amino acid StrepII epitope tag, in WT plants. The hope was that this would allow for each APT1 to form its natural protein complexes in planta and these complexes could be recovered by affinity chromatography. Unfortunately, this method did not reveal any protein interacting partners for either isoform. However this is not a definitive result; other methods should be followed up to capture protein interacting partners since some interactions are quite dynamic and weak, although yet still functionally important. In addition, there are a number of other factors that may have contributed to a negative result. One factor could be the buffers and conditions used to purify protein complexes using the StrepII tag that may be incompatible with the APT1 proteins. This even includes the addition of other
metabolites necessary for mediating interacting partners as demonstrated by Katsir et al (2008). These authors showed that the presence of the hormone-conjugated amino acid (JA-Ile) was necessary to pull down tomato COI1 protein from leaf extracts using the his-tagged JAZ1 protein. For APT1.1, this may implicate the need to add adenine or cytokinins into the purification mix. A second cause of this negative result could apply to the timing at which APT1.1 is expressed. If APT1.1 and its potential protein interacting partners are co-expressed then over-expression of APT1.1 will not yield any protein interacting partners until the others are expressed.

**Functional complementation of apt1-3**

Previous work suggested that changes in both adenylate and/or cytokinin metabolism via APT1 contribute to the male sterility phenotype in apt1-3 mutants (Zhang et al, 2002). To test whether APT1.1 alone could rescue male fertility the APT1.1 ORF was introduced into apt1-3 by genetic crossing. Fertility was completely restored in all three lines of apt1-3 mutants that over-expressed APT1.1. No abnormal phenotypes were observed in these transgenic lines with regards to their vegetative or reproductive growth.

There are several interpretations of this result. One, over-expression of APT1.1 may provide sufficient cytosolic expression for complementation of fertility. Support for this argument can be clearly seen in the oxt1 mutant where a small fraction of cytosolic APT activity (4-6%) is able to sustain fertility. The second interpretation is that restoring one adenylate pool, that within the plastids, was sufficient to provide needed adenylates required for pollen development. Alternatively, if altered cytokinin metabolism is contributing to the male sterile phenotype then maybe over-expressed APT1.1 can restore this imbalance either in chloroplasts
or in the cytosol. Use of the native \textit{APT1.1} promoter might reveal if the natural level of APT1.1 expression is sufficient to restore male sterility. Nonetheless, previous complementation using the APT1.2 isoform by Gaillard \textit{et al} (1998) and in this thesis using APT1.1 provides strong evidence that both isoforms are functionally similar. However, these data do not imply that these enzymes have similar enzymatic activities. In both cases male sterility was restored in \textit{apt1-3} plants. However, these results do not distinguish whether adenine or cytokinin metabolism is responsible for this effect.

\textbf{Using amiRNAs in isoform specific gene silencing}

AmiRNA lines were constructed to learn the effect of having only one APT1 isoform in WT and gain insight into the functional contribution of each enzyme in its native context. These constructs were designed to specifically target the 5’ UTR of either \textit{APT1.1} (ami1.1) or \textit{APT1.2} (ami1.2). The ami1.1 lines had approximately 50% reduced APT1.1 in leaves relative to WT. Surprisingly, APT1.2 protein levels were also affected in these lines by up to 50%. In the ami1.2 lines, a 50-75% reduction in APT1.2 protein in leaves was observed in some lines; APT1.1 protein levels were unaffected. For both ami1.1 and ami1.2 plants, APT1 protein abundance in inflorescences was the same as WT.

No phenotypes were observed with respect to vegetative or reproductive growth in any of the five independent plant lines of each amiRNA construct analyzed. As a result, these lines were not assayed for APT enzymatic activity, as the reduced level of expression was not enough to induce any phenotypes. However this approach should not be neglected as improvements have been made to amiRNA design software. Clearly demonstrated in these lines is the need for amiRNAs with higher efficiencies to reduce protein levels much lower than observed in the
present lines. As of now, these lines could be analyzed by mass spectrometry for insights into small perturbations in purine, nucleotide, and cytokinin pools.

**Annotating APT1.1 isoforms in other plant species**

Currently, the only plant genome annotated to contain a second *APT1* transcript variant (APT1.1) is Arabidopsis. APTs from other plant and non-plant species are annotated as having one transcript variant with high sequence similarity to that of APT1.2. This allows one to question whether any other species, aside from Arabidopsis, has an APT transcript containing an additional N-terminal Exon that functions as a transit peptide. As Exon 1 was shown to function as a transit peptide and there is low sequence conservation among transit peptides it is inefficient to find homologous sequences using BlastP, tBLASTn, or PSI-BLAST. However, transit peptides from closely related species are less divergent making these algorithms useful in identifying putative Exon 1 sequences within a family.

The approach I took involved searching through APT EST collections and identifying the highly conserved Exon 2 start site found in all APTs. Once identified, the EST was virtually translated and then I manually searched for upstream methionines. Putative Exon 1 sequences were defined as sequences fulfilling two criteria. First, it must have only one upstream methionine in-frame with the methionine of Exon 2. The presence of additional methionines could confuse which one provides the start site of translation. In addition, setting the criterion for only one methionine adds greater confidence that this event is real. Second, the putative Exon 1 must contain at least 30 amino acid residues. Arabidopsis Exon 1 contains 60 amino acids residues, but as transit peptides are quite variable in length a more open-size limit was selected for this criterion.
This screen has lead to two interesting observations. The first observation was that putative Exon 1 sequences were only identified in plants; none were found in yeast, mammalian, or prokaryotic transcriptomes (Figure 28). Secondly, all of these putative Exon 1 sequences contained only one upstream methionine, and as anticipated, the sequence lengths and compositions are variable, consistent with their roles as transit peptides.

Figure 28. Multiple sequence alignment of Arabidopsis APT1.1 with APTs from other plants species. Putative Exon 1 sequences were identified by aligning the highly conserved amino acid sequence of APT1.2 with APT ESTs from other species. A manual search for an upstream open reading was carried out and sequences aligned using ClustalW.

To identify if these putative Exon 1-containing species follow speciation events, other APT sequences were analyzed. A radial tree of all APTs found in the NCBI database shows that APT evolution follows speciation (Figure 29). This is evident as plant, fungi, animal, and bacterial APT sequences all cluster together. Additionally, the phylogeny correctly shows that fungi and animal APTs are more similar than plant APTs consistent with speciation. Thus, this radial tree predicts that plant APTs evolved from a single plant APT ancestral gene.
Figure 29. Radial phylogenetic tree of APT amino acid sequences from bacteria, animals, fungi and plants. All APT amino acid sequences in the NCBI database (1231 total sequences) were used to construct this phylogenetic tree. This tree shows that APT evolution is consistent with speciation as the bacterial sequences are the most ancestral, and due to the fact that and fungi and animal APTs cluster more closely together than plant sequences. Additionally, this tree predicts that all plant APTs arose from one ancestral APT gene. Bioinformatic analysis performed by Andrew Doxey.

Using this information the plant APTs containing a putative Exon 1 sequence were located on a phylogenetic tree (Figure 30). The most obvious observation from this tree is that APT isoforms from the same species exist within different clusters. This is likely due to genome duplication events. Assuming all species start off with one copy of an APT gene, genome
duplication events lead to the formation of gene families and sequence divergence throughout evolution. This explains why APT gene families are dispersed in different clusters.

Interestingly, the identified putative Exon 1-containing APT genes all cluster together. This preliminary analysis predicts that this Exon 1 arose from one common ancestor and is thus a monophyletic trait. Further investigations, will have to explore APT genes within this cluster to confirm they contain a putative Exon 1. The same analysis needs to be performed on those species outside the cluster to correctly identify the absence of Exon 1, and the time in evolution in which this monophyletic trait arose.
Figure 30. Bayesian phylogeny of plant APTs. All plant APT amino acid sequences from the NCBI database were used to construct this phylogeny rooted using a bacterial sequence. Consistent with speciation, this tree shows the most ancestral branches as moss (green) and pine (purple). Based on this phylogeny it is predicted that after moss the APTs diverged into three “types”: the Type 1 and Type 2 sequences and a questionable third type found in-between. Interestingly, the monocot and dicot sequences form subclusters in both the Type 1 and Type 2 clusters. Putative Exon 1 sequences identified in an
earlier analysis (Figure 28) were identified on this phylogeny and remarkably these all group within one cluster. The boxed X in this image represents the ancestral APT protein in which the putative Exon 1 sequence arose. Support values for each cluster are shown on the branches; the scale bar signifies 2 amino acid mutations per 100 amino acids along this evolutionary distance. Analysis done by Andrew Doxey.

Although much remains to be learned, the results of this research indicate that the expression of this housekeeping enzyme is highly regulated and compartmentalized within the cytosol and chloroplasts of plant cells. Bioinformatic analyses support that these dual APT isoforms exist in many other plant species. This in turn will be useful in elucidating what event during plant evolution required this function to arise. The continued studies of the APT1 locus will be of greater value in understanding the control of purine metabolism in plants.

The second part of this research focused on two APT1-deficient mutants, apt1-3 and oxt1. The apt1-3 mutant has a single nucleotide polymorphism at the splice junction of Exon 4. This mutant has about 1-2% of the APT activity of WT plants and is completely male sterile. The oxt1 mutant contains a T-DNA insertion within Intron 1, which reduces its APT activity to about 3.1% of WT levels. However, unlike apt1-3, oxt1 is as fertile as WT. The objective of this study was to investigate the basis of fertility in this APT-deficient mutant.

**APT1 activity does not correlate with decreased fertility in oxt1 and oxt1apt1-3 mutants**

Initial characterization of residual APT enzymatic activity showed that apt1-3 leaves contained 2.1% residual WT APT activity while oxt1 had 3.1% residual APT activity. These activity determinations were made using a radiochemical assay that measures the amount of $^3$H-adenine converted to $^3$H-AMP over time. The activities of different extracts are compared on a protein basis. Since these assays are done on crude extracts it is possible that the enzymes of the two-step adenine salvage pathway (adenine phosphorylase and ADK), which are presumably
also in the mixture, may contribute to the apparent APT activity. Since the crude extracts used for the APT assay are not desalted to remove small molecules, the ATP needed for the ADK reaction is available in small amounts (B. Moffatt, personal communication). I once tested the use of desalted extracts in the APT assay and noted a small decrease in $^3$H-AMP production (data not shown). This decrease may be due to removal of ATP by the desalting step, thereby inhibiting ADK activity, and any contribution of the 2-step pathway. However this observation needs to be further investigated since adenine phosphorylase activity has been shown to be very low in plants and thus the contribution of the secondary pathway is likely very low as well (Ross, 1981). Thus it is doubtful that the use of desalted extracts would have changed my results or conclusions to any significant extent.

Previous phenotypic analysis of oxt1 showed that it has 75% increased tolerance to chemically induced oxidative stress, when plated on $\frac{1}{2}$ MS media supplemented with the catalase inhibitor, AT, and the glutathione biosynthesis repressor, BSO. In comparison, WT showed signs of chlorosis and root growth inhibition (D. Falcone, personal communication). Interestingly, apt1-3 did not exhibit this stress tolerance as signs of chlorosis were observed. Root length measurements of apt1-3 were not done due to the difficulty of growing these seedlings to be developmentally matched with WT and oxt1. These seedlings had to be initially germinated on $\frac{1}{2}$ MS media supplemented with DAP to select for homozygous seedlings among the segregating F2 population. Although the apt1-3 seedlings are more resistant to this compound than WT plants, their residual APT activity does cause some DAP-associated toxicity and has marked effects by decreasing their growth rate relative to WT. Additionally, apt1-3 plants are developmentally delayed relative to WT and oxt1 throughout their life cycle in the absence of DAP selection.
To investigate whether the small difference in APT activity between *apt1-3* and *oxt1* was the basis for the difference in their fertility and stress tolerance, I created an F1 hybrid which would theoretically have an intermediate residual APT activity. The resulting F1 plant had a developmental chimeric phenotype in that the early flowers were fertile but later ones were sterile. To explain this effect, I hypothesized that the APT activity in the fertile buds was likely higher than that in the sterile buds. However, quite the contrary was observed: APT activity in the sterile buds was 4.6% higher than the fertile buds. Even more remarkable was that the APT activity in sterile buds of *oxt1apt1-3* plants was 2-4% higher than in *oxt1* buds. The mechanism for how or why this occurs is unknown.

One possibility was that the other less predominant APTs, APT2 and APT3, might be contributing activity in the flowers. These APTs were previously shown to have a much lower affinity and $V_{\text{max}}$ for adenine relative to APT1, but a higher affinity for cytokinins (Schnorr *et al.*, 1996; Allen *et al.*, 2002). Semi-quantitative RT-PCR was used to detect any differences in the transcript abundances of APT2 and APT3 relative to WT. These data showed that there was no change in APT2 or APT3 transcript abundance in inflorescences of *oxt1*, *apt1-3*, and *oxt1apt1-3* relative to WT. While this may be true, these results are not consistent with those previously reported by Zhang (2000). Using *in situ* hybridization it was shown that APT2 and APT3 transcripts increase in *apt1-3* inflorescences relative to their abundance in WT inflorescences. The basis of this discrepancy may be attributed to the method used to detect transcript abundance in each case. The advantage of *in situ* hybridization is that changes in transcription can be detected in specific cells or tissues of an organ whereas for an RT-PCR an entire leaf or inflorescence is used as the source of RNA. In the case of these assays, I used entire inflorescences for all my assays (enzyme activity, RT-PCR and immunoblots). Thus my RT-PCR
analysis measures the total transcript abundance of all cells in the inflorescence and likely dilutes minor increases that occur in only a few cells, as was observed in the in situ hybridization data. A more accurate approach may involve measuring protein levels of APT2 and APT3 in oxt1, apt1-3, and oxt1apt1-3 to confirm that changes in transcript abundance are resulting in changes in protein synthesis.

The contribution of the secondary salvage pathway was also assessed in oxt1 and apt1-3 for any differences that may account for their differing fertility. Radiochemical assays showed that while both had slightly higher ADK activity relative to WT (2-fold and 1.7-fold, respectively), the increases in the two mutants was not significantly different.

Although this result indicates that increased ADK activity does not account for the fertility of oxt1, the fact that ADK activity increases in both APT-deficient mutants is interesting. The increase in ADK activity may reflect a response to accumulating adenine. This would be accomplished by adenine phosphorlyase metabolizing adenine to adenosine, and ADK responding to this increased adenosine pool. Preliminary analysis of adenine levels in oxt1 and apt1-3 by HPLC and mass spectrometry analysis has shown that adenine levels are 1.5 to 3-fold greater in oxt1 and apt1-3 plants relative to WT (D. Falcone, personal communication). Thus it is possible that even modest increases in adenine are sensed by the cell and triggering a compensatory response in activity by the two-step adenine salvage pathway. It should be noted that these values are very preliminary and further work is underway to establish more reliable and consistent readings.
Altered cytokinin levels in oxt1 and apt1-3 plants

Kinetic analysis of recombinant APTs indicates that they are capable of accepting cytokinin bases as substrates. It was of interest to investigate whether cytokinin profiles of the APT-deficient mutants were altered. Leaf and bud tissue of oxt1 and apt1-3 plants was sent to the lab of Dr. Neil Emery (Trent University) to be analyzed by LC-tandem mass spectrometry for various cytokinin species. The data obtained to date from these measurements has not been consistent between different samplings so definitive conclusions cannot be made. However it is quite evident from these measurements that cytokinin bases are increased in both oxt1 and apt1-3 leaves and buds. Interestingly, the predominant cytokinin bases in Arabidopsis, tZ and iP, show the greatest fold change relative to WT (data not shown).

Interestingly, the cytokinin ribosides are not significantly altered for any of the cytokinin species measured. Thus, the cytokinin bases are accumulating but not being metabolized by adenine phosphorylase into cytokinin ribosides. This absence of adenine phosphorylase activity is consistent with previous reports that its activity is low in plants (Ross, 1981). Additionally, if cytokinin riboside levels are not affected it suggests that the higher ADK activity in these plants is not a result of increased cytokinin metabolism through the secondary salvage pathway.

As these measurements are preliminary and need further work to establish reliable values, it can be concluded at this time that the cytokinin pools are definitely altered by APT1-deficiency. Precautions have been taken to ensure consistent sampling methods each time. Such precautions include collecting tissue at a specific time in the photoperiod and collecting developmentally matched leaves and inflorescence buds.
Increased adenine is associated with stress tolerance in WT

The observations thus far have demonstrated that APT activity is not tightly associated with fertility. This is most evident in oxt1apt1-3 plants where the APT activity is 4.6% higher in sterile buds as compared to the fertile buds. Equally as important is the approximate 1% difference in APT activity between oxt1 and apt1-3. Furthermore, the secondary salvage pathway cannot account for differences in oxt1 and apt1-3 as the ADK activity measured is similarly increased in both mutants. These observations lead me to conclude that a third component outside of these purine salvage enzymes may be involved.

The identity of this third component might be a protein interactor, miRNA, or a metabolite. To test for a protein interacting partner StrepII epitope tagged APT1.1 and APT1.2 isoforms were introduced into Arabidopsis. As discussed in the previous section, no unique protein interacting partners were found using this particular approach; further research in this direction is necessary before the involvement of interacting partners can be ruled out. Alternatively, metabolite levels have also been investigated for differences that may attribute to the unique phenotypes observed in oxt1 and apt1-3 mutants.

Preliminary data suggests that adenine may contribute to the stress tolerance of oxt1. WT seedlings were grown on ½ MS + AT/BSO media supplemented with 2, 20, and 200 µM adenine (Figure 31). At 2 µM and 20 µM of adenine WT seedlings showed increased stress tolerance identical to oxt1 on ½ MS + AT/BSO alone. However, oxt1 seedlings on 20 µM adenine became stress sensitive as these seedlings grew to half the size of oxt1 grown on lower concentrations of adenine. At 200 µM, oxt1 showed marked defects in growth and chlorosis. Immediate signs of growth inhibition and chlorosis were observed in apt1-3 mutants with any addition of adenine.
As mentioned earlier, HPLC analysis shows that adenine is modestly increased in apt1-3 mutants (D. Falcone, personal communication) coinciding with the residual APT1 activity measured in each plant. Based on these observations, the earlier hypothesis of a threshold of APT activity can be revised. Instead of proposing protein activity perhaps metabolite levels are what distinguish the phenotypes observed in oxt1 and apt1-3. Since metabolite measurements showed adenines levels increasing from oxt1 to apt1-3 then it is expected that further increases of adenine in oxt1 would yield an apt1-3 phenotype. This is consistent with the observation that
oxl mutants grown on 20 μM adenine are no longer tolerant to AT/BSO. Interestingly, adding adenine to WT seedlings resulted in an oxl-like stress tolerant phenotype in these plants. Thus, oxl’s APT1-deficiency may be enough to allow an accumulation of adenine that is advantageous to stress and additionally its residual APT activity is just sufficient to sustain adenylate synthesis, resulting in a fertile plant. In the case of aptl-3, its APT1-deficiency may be below a threshold that results in a cytotoxic accumulation of adenine and insufficient nucleotide biosynthesis to be fertile.

Although this metabolite difference is convincing of a threshold hypothesis for the oxl stress tolerant phenotype, it is unknown whether adenine levels affect male fertility in the same dose-dependent manner. The current hypothesis is that a third component accounts for the phenotypic difference between oxl and aptl-3. This third component may be a metabolite, protein, or non-coding RNA that oxl uses to its advantage to compensate for APT-deficiency, in addition to conferring increased tolerance. This increase in stress tolerance may provide insight into the third component, as it could be the mechanism in which APT-deficiency is compensated for in oxl. RT-PCR analysis of select antioxidant defense genes in oxl indicates that three of these genes (stromal ascorbate peroxidase, thylakoid ascorbate peroxidase, and catalase 3) are up-regulated under non-stress conditions (Sukrong et al., manuscript in preparation). Thus, the next steps could involve investigating the transcriptional response of these genes in aptl-3 in addition to expressing these genes in aptl-3 as a means to test for the restoration of male fertility. Alternatively, one must consider that these genes may be indirectly involved in the observed fertility in oxl. As such, the proteins with which these antioxidant defense proteins interact with and the collection of metabolites involved may also need to be elucidated. I hypothesize that these investigations may yield one of two conclusions. First, APT-deficiency is a source of stress
during pollen development and the increased response of antioxidant defense genes in \( oxt1 \) is sufficient to rescue male sterility. Second, the increased stress response may be an indirect response to the true underlying pathway responsible for restoring male fertility in \( oxt1 \). I predict that the second conclusion is more plausible, as it is difficult to explain differences in adenine and cytokinin metabolism between \( oxt1 \) and \( apt1-3 \) based solely on a stress responsive pathway.

Thus, the basis of male fertility of \( oxt1 \) and the sterility of \( apt1-3 \) has yet to be learned. The data are clear that APT and ADK activities are not the only factors that determine fertility or sterility in \( oxt1 \) and \( oxt1apt1-3 \). It is recognized that metabolites may be an important contributor to the fertility of these mutants. In \( oxt1 \), adenine may be key as this metabolite was able to confer the same degree of stress tolerance to WT seedlings as initially observed in \( oxt1 \) plants. In both APT1-deficient mutants, the importance of cytokinin bases was clearly demonstrated by altered levels detected in both leaf and floral organs. Further resolution of the differences in cytokinin levels will no doubt provide valuable insight of these hormones in both stress tolerance and male fertility. The discovery and characterization of the unusual phenotype of \( oxt1apt1-3 \) hybrids will lead the way in providing a model system to study changes in metabolite levels that contribute to the shift from fertility to sterility.
Appendices

Appendix 1

Figure 32. Testing the sensitivity and specificity of the anti-APT1 polyclonal antibody. WT (lanes 2 and 3) and oxt1 (lane 1) leaf protein extracts were probed with the anti-APT1 polyclonal antibody at a dilution of 1:10,000 (Left panel). The anti-APT1 antibody proved to be specific as an abundant polypeptide at 24 kDa was detected in WT extracts, whereas very little polypeptide was detected for oxt1, consistent with this mutant being APT1-deficient. Ponceau S staining was used to test for equal loading. To test the sensitivity of the anti-APT1 antibody, 10 ng, 1 ng, and 500 pg of recombinant APT1.2-6xHIS protein was probed with the anti-APT1 antibody using two dilutions (Right panel).
Appendix 2

Figure 33. Computational modeling of the 5’ UTR of the \textit{APT1.1} mRNA. The region depicted is the 5’ terminal end of the \textit{APT1.1} mRNA consistent with that recovered in the RIKEN Arabidopsis full-length cDNA clone collection. RNA folding was performed by Yingfu Li, McMaster University.
Appendix 3

Figure 34. Partial genomic sequence of the APT1 gene. Two mRNAs arise from the APT1 gene in Arabidopsis. The APT1.1 primary transcript initiates at the first red arrow and is spliced at the splice acceptor and splice donor sites underlined in blue, to remove the first Exon. The translation of this mRNA begins at Exon 1 (highlighted in yellow) and the resulting polypeptide includes the 6 amino acids adjacent
to Exon 2 (highlighted in red). The APT1.2 primary transcript initiates at the second red arrow with part of Intron 1 forming the 5′UTR. The translation of this mRNA initiates at Exon 2. Note that the APT1.2 polypeptide does not contain the 6 amino acids adjacent to Exon 2; these are only present in an APT1.1 polypeptide. Blue arrows represent forward primers AP183, AP184, and AP187 used to PCR fragments of the 5′ UTR in Figure 5. Primer AP183 (first blue arrow) is located in the same region as the predicted stem-loop structure shown in Figure 33. Orange arrows are forward primers AP186 and 187 using in semi-quantitative RT-PCR. Yellow triangle represents location of T-DNA insertion in oxt1 allele; black line represents point mutation of apt1-3 (GT → AT).
Appendix 4

oxt1 transgenic lines expressing either the APT1.1 or APT1.2 ORF were constructed (oxt1/actinP::APT1.1 and oxt1/actinP::APT1.2, respectively). The APT1.1 and APT1.2 ORFs were cloned under the expression of the actin promoter in a vector expressing glufosinate ammonium resistance (which confers BASTA resistance). These constructs were introduced into oxt1 mutants by the floral dip method. Three stable transformants were recovered for each line. Immunoblot analysis of oxt1/actinP::APT1.1 plant containing lines detected a 26 kDa APT1.1 polypeptide; a 24 kDa APT1.2 polypeptide was detected in oxt1/actinP::APT1.2 plants. For both plant lines no effects on vegetative and reproductive growth were observed relative to WT.

The detection of APT1.1 and APT1.2 polypeptides in the oxt1 transgenic mutants is an interesting observation considering oxt1 heterozygous plants (oxt1/WT APT1). oxt1 heterozygous plants grown on ½ MS media supplemented with AT + BSO exhibit increased stress tolerance and phenotypically resemble oxt1 homozygous plants (Sukrong et al., manuscript in preparation). These data indicate that the oxt1 allele is dominant to the WT APT1 allele. However, this dominance is not exhibited over the actinP::APT1.1 or actinP::APT1.2 transgenes in oxt1 as the respective APT1 polypeptides were detected by immunoblotting. In these transgenes only the APT1 ORFs were present, the APT1 regulatory elements (promoter, UTR, intronic regions) were absent. Taken together, these results point towards the importance of the WT APT1 regulatory elements as targets for oxt1 to exhibit its dominance over this allele.
References


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