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# An *in silico* data mining of the ammonium transporter gene family in *Ananas comosus* L.

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## Abstract

Arguably, nitrogen (N) is an important and essential component for plant growth and development. Ammonium is a major inorganic nitrogen source for plants mobilized by ammonium transporter (AMT) among N available sources. In this study, data mining revealed that in the *Ananas comosus* L. genome was identified eight AMT family genes. The eight pineapple AcoAMT proteins were identified and phylogenetically clustered into two groups with AMT proteins from other plants. Two pairs of *AcoAMT* (*AcoAMTa* and *AcoAMTg*) genes located on chromosome 1 and unchromosome appear to be segmental duplications. Based on this information, we conducted a comprehensive analysis using some bioinformatics tools to characterize the identified genes individually. The comprehensive analysis of AMT will provide an essential foundation for further investigation of the regulatory mechanisms of *AcoAMTs* in *A. comosus* L.

Keywords: bioinformatic analysis; pineapple; ammonium; nitrogen transport.

# Uma mineração de dados in silico da família do gene transportador de amônio em Ananas comosus L.

## Resumo

Indiscutivelmente, o nitrogênio (N) é um componente importante e essencial para o crescimento e desenvolvimento das plantas. O amônio é a principal fonte de nitrogênio inorgânico para as plantas, sendo mobilizado pelo transportador de amônio (AMT), dentre as fontes de N disponíveis. Neste estudo, a mineração de dados revelou que no genoma de *Ananas comosus* L. foram identificados oito genes da família AMT. As oito proteínas AcoAMT de abacaxi foram identificadas e filogeneticamente agrupadas em dois grupos com proteínas AMT de outras plantas. Dois pares de genes *AcoAMT* (*AcoAMTa* e *AcoAMTg*) localizados no cromossomo 1 e não cromossômicos parecem ser duplicações segmentares. Com base nessas informações, realizamos uma análise abrangente usando algumas ferramentas de bionformática com a finalidade de caracterizar individualmente os genes identificados. A análise abrangente do AMT fornecerá uma base importante para uma investigação mais aprofundada dos mecanismos regulatórios de *AcoAMTs* em *A. comosus* L.

Palavras-chave: análise bioinformática; abacaxi; amônio; transporte de nitrogênio.

#### Introduction

Agronomically, nitrogen (N) is one of the most necessary macronutrients for plant growth and development (TEGEDER; MASCLAUX-DAUBRESSE, 2018). Nitrogen is a fundamental element of several biological molecules, such as nucleotides, chlorophyll, amino acids, and proteins. Plants can take up the N present in the soil in two ways: either inorganic N (nitrate; FAN et al., 2017) or organic N (ammonium, amino acid, and urea; reviewed by TEGEDER; MASCLAUX-DAUBRESSE, 2018). The use of N during plant growth and development can be divided into three main stages: (I) absorption; (II) assimilation and (III) remobilization of N. In addition to N

absorption from the soil, the nitrogen use efficiency (NUE) also depends on the assimilation of nitrate and ammonium, and on how the crop recycles organic N (MASCLAUX-DAUBRESSE et al., 2010; XU et al., 2012). Mainly, ammonium is the preferential form of N uptake by plants because the ammonium assimilation requires less energy than nitrate (BLOOM, 2015). In plants, the transport of nutrients, water, and metabolites is often mediated by a series of gene families of transporters proteins inserted into cell membranes (NACRY et al., 2013).

Specifically, the members of this family in plants belong to the AMT superfamily, which permeates ammonium via NH4<sup>+</sup> uniport or  $NH_3/H^+$  co-transport, or belong to the MEP subfamily (Methylammonium Permeases), which includes AmtB from the bacterium Escherichia *coli* and human homologs (AMT/MEP/Rh transporter/methylammonium (ammonium permease/rhesus) (LOQUÉ; VON WIRÉN, 2004; LUDEWIG et al., 2007). The presence of the transmembrane domains (TM) in this transporter (AMT) can be ranged from 10-12 assembled into (TM) and are homo/or heterotrimers (LUDEWIG et al., 2003). Another important description of this transporter is that in plants, they are subdivided into two subfamilies: AMT1 and AMT2 (CASTRO-RODRÍGUEZ et al., 2016).

AMTs genes were identified in prokaryotic and eukaryotic organisms (reviewed by McDONALD; WARD, 2016). The first ammonium transporter genes were identified in (MARINI et studies with yeasts al., 1997). AMT family genes were shown to be highly expanded in plants and have already been characterized in several species, such as Zea (GU et 2013), Arabidopsis mays L. al., thaliana (HUANG et al., 2015), Oryza sativa L. (FERREIRA et al., 2015), Triticum aestivum L. (DUAN et al., 2016), Coffea canephora (DOS SANTOS et al., 2017) and Solanum lycopersicum L. (FILIZ; AKBUDAK, 2020). Because of the numerous studies in plants that describe the AMT gene's identification and characterization, there are no reports on this transporter in pineapple (Ananas comosus L.).

Pineapple is a tropical fruit, perennial monocot from the Bromeliaceae family, and has significant economic importance in tropical and sub-tropical areas. In world production, the pineapple is considered the third most important tropical fruit after banana and citrus

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(reviewed by XIE et al., 2018). Ming et al. (2015) made available pineapple completed genome sequencing. These databases provide valuable information in the search for agronomic interest genes, enabling us to identify the AMTs genes in the pineapple genome. Given the above, we performed the analyses of sequence characteristics, gene structures, chromosome distribution. motif compositions. and evolutionary relationships. The present study may understand the transcriptional mechanisms of the AMTs genes in response to pineapple's nutritional efficiency.

# **Material and Methods**

We downloaded the protein sequences, genomic sequences and coding sequence (CDS) of pineapple obtained from Phytozome database (GOODSTEIN et al., 2012). To confirm the selected candidate gene sequences, belong to the AMT gene family, we used the National Center for Biotechnology Information (ALTSCHUL et al., 1997) database, using the BLASTP tool. The isoeletric point (pl) and theoretical molecular weight (kDa) of AMTs genes were calculated using the ExPASy server (http://web.expasy.org/compute pi/). Also was evaluation of the grand average of hydropathicity (GRAVY) of all identified proteins through the GRAVY calculator (GASTEIGER, 2003). The predictions of subcellular localization were verified with the help of Plant-mPloc tool (CHOU; SHEN, 2010). The gene structures (exon/intron) of the AMTs genes were determined using the Gene Structure Display Server (HU et al., 2015). For these analyses, we used the predicted coding sequence (CDS) and their corresponding genomic DNA sequences. Additionaly, we used the TMHMM server v. 2.0 (KROGH et al., 2001) for prediction of transmembrane helices in AMT proteins. To phylogenetic analysis initially, protein sequence alignments and analysis were conducted using ClustalW (LARKIN et al., 2007). Posteriorly, the phylogenetic trees were constructed with MEGA7.0 software (KUMAR et al., 2016) using the Neighbor-Joining (NJ) method. We also verified the reliability of the obtained trees topology by the bootstrap method (1.000 replicates). Individual, the physical locations of AcoAMTs genes were obtained from the database of pineapple genome and posteriorly, the map of the chromosome location of genes was constructed through the Mapchart 2.2 software (VOORRIPS, 2002). We then further investigated the synonymous (*Ks*) and non-synonymous (*Ka*) substitution rates of the paralogs genes by using the *Ka\_Ks* calculator 2.0 (ZHANG *et al.*, 2006). To analyze the synteny relationship of the *AMT* genes, we used the orthologous genes between *A. comosus* L. and other two species: *A. thaliana* and *O. sativa*. Then, the Circos software was used to represent the syntenic relationships (KRZYWINSKI *et al.*, 2009).

#### **Resuts and Discussion**

Ammonium acquisition by plants is transporter AMT-mediated that are ubiquitous plasma membrane proteins and essential for the nitrogen demand of plants (VON WIRÉN et al., 2000; HAO et al., 2020). In this study, we identified and cataloged eight genes coding for putative AMTs. We described the details of all eight pineapple AMT transporter proteins, including gene identifier, chromosome location, protein length, molecular weight, isoelectric point (pl), prediction of the hydrophobicity (GRAVY), subcellular location and their family. Collectively, all the information is listed in Table 1. The number of the genes that we obtained is similar to that in Setaria italica, O. sativa and Sorghum bicolor (Table 2) (see a review of VON WITTGENSTEIN et al., 2014; SANTOS et al., 2017). In the present study, it is possible to observe the conservation and confirmed of number AMT genes in monocot species. Two genes are located on scaffolds (scaffold\_558 named AcoAMTq) and (scaffold 881 named AcoAMTh). We believe that this gene (AcoAMTa - Aco021941.1) is an alternate form of the AcoAMTa gene, as they have the same characteristics, according to data presented in Table 1. All the genes identified were renamed based on their chromosomal location (Table 1).

The results showed that the lengths of AMT transporter proteins range from 90 (AcoAMTe - Aco003227.1) to 495 (AcoAMTh - Aco030863.1) amino acids. The predicted molecular weights of proteins ranged from 9.95 (AcoAMTe - Aco003227.1) to 52.73 kDa (AcoAMTh - Aco030863.1) and the pl values were between 4.79 (AcoAMTe - Aco003227.1) and 8.53 (AcoAMTa - Aco009722.1 and AcoAMTg - Aco021941.1), respectively. In *S. lycopersicum* L. the molecular weight and pl values ranged from 49.65 to 55.37 kDa and from 5.33 to 7.16 (FILIZ; AKBUDAK, 2020). Generally,

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the structure of members AMTs family can range from 45–50 kDa and approximately lenght between 400 – 450 amino acids (NINNEMANN *et al.*, 1994; BLAKEY *et al.*, 2002). The predictions of the hydrophobicity of the deduced amino acid sequences indicated that the GRAVY of all AcoAMT proteins were above zero (Table 1).

**Table 1.** Characterization in silico of AMT genefamily in pineapple.

Gene	Gene identifier	Chromossome location	Protein (aa)	MW (kDa)	рI	GRAVY	Subcellular prediction	Category (family)
AcoAMTa	Aco009722.1	LG01:108464110894	460	49.73	8.53	0.616	Cell membrane	AMT2
AcoAMTb	Aco011210.1	LG01:1341228913419316	473	51.51	8.10	0.411	Cell membrane	AMT2
AcoAMTc	Aco014095.1	LG13:5757858987	469	49.81	6.14	0.570	Cell membrane	AMT1
AcoAMTd	Aco012484.1	LG13:14907081492001	342	36.11	7.11	0.594	Cell membrane	AMT2
AcoAMTe	Aco003227.1	LG17:12132481213539	06	9.95	4.79	0.361	Chloroplast/ Mitochondrion/Nucleus	AMT1
AcoAMTf	Aco007888.1	LG21:98038459808433	482	51.51	5.89	0.543	Cell membrane	AMT2
AcoAMTg*	Aco021941.1	scaffold_558:7233174767	460	49.73	8.53	0.616	Cell membrane	AMT2
AcoAMTh	Aco030863.1	scaffold_881:3627337760	495	52.73	6.69	0.440	Cell membrane	AMT2

<sup>\*</sup> Possible duplication of the *AcoAMTa* gene (see Table 3).

Almost all AMT transporters were predicted to be located in the cell membrane. Only one AMT (AcoAMTe - Aco003227.1) has been indicated with subcellular localization in the chloroplast, mitochondrion, and nucleus (Table 1). Detailed information about copy number of *AMT1* and *AMT2* transporters of different plant species are presented in Table 2.

**Table 2.** List of members from the *AMT1* and *AMT2 genes* identified in different plants species, based in von Wittgenstein *et al.* (2014) and dos Santos *et al.* (2017).

	Number of members		
Species - Monocot/Dicot	AMT1	AMT2	
Arabidopsis thaliana	5	1	
Glycine max	5	5	
Vitis vinifera	1	1	
Populus trichocarpa	6	5	
Coffea canephora	4	4	
Setaria italica	2	6	
Oryza sativa	2	6	
Sorghum bicolor	2	6	
Manihot esculenta	5	4	
Cucumis sativus	4	2	
Ricinus communis	4	3	
Ananas comosus L.	2	6	

In addition, we note that only two AMT1 transporter members have been identified, which leads us to believe that the duplication mechanisms for this type of transporter in the evolution of the A. comosus L. genome were not relevant (Table 1 and Table 2).

Evolutionarily, it is known that the structural diversity of the gene is one of the main evidence, besides that it also provides valuable

information for the evolution of the innumerable multigenic families (CAO; SHI, 2012). In this sense, we analyzed the putative gene structure based on the CDS and genomic DNA sequences of each gene (Figure 1). The number of introns in *AcoAMT* genes varied from one to three. These results indicated that *AcoAMT* in pineapple shows lower gene structure diversity.

**Figure 1.** Representation of the *AcoAMT* genes structure, the yellow box, blue and black lines represent exons, upstream/downstream regions of the gene and introns, respectively.



The number of transmembrane helices (TMHs) in pineapple showed variations between 1 and 11 TMs (Figure 2).

Figure 2. Prediction of transmembrane domains of AcoAMT sequences were performed using TMHMM server v. 2.0.



It is important to clarify to readers that the little transmembrane domain presented by AcoAMTe is due to the size of the protein sequence (90 aa). Although only one AcoAMT sequence showed little TM, ours analyzes corroborate with previous studies, about on the indicative of the amount of the transmembrane domain (McDONALD *et al.*, 2012; FILIZ; AKBUDAK, 2019). This result may be indicative of the complexity of ammonium homeostasis in pineapple. The structural formation of an AMT is



formed by 11–12 transmembrane regions, standing out with feature sequences "D (F Y W S) A G (G S C) X2 (L I V) (E H) X2 (G A S) (G A) X2 (G A S) (L F)" located at its transmembrane region 5 and "D D X (L I V M F C) (E D G A) (L I V AC) X3 H (G A L I V) X2 (G S) X (L I V A W) G" at transmembrane region 10 (VON WIRÉN; MERRICK, 2004; reviewed by HAO *et al.*, 2020).

Firstly, to understand the evolutionary relationships of AMTs between pineapple and *A. thaliana*, we aligned multiple AMTs protein

sequences and constructed an unrooted phylogenetic tree for the identified multiple

AMTs genes using ClustalW and MEGA 7.0 (Figure 3).

**Figure 3.** Phylogenetic tree of AMT proteins sequence from *A. comosus* L. and *A. thaliana*. Phylogenetic tree was constructed with the Neighbour-Joining (NJ) method using MEGA7.0 program with 1.000 bootstrap replicates. Branches with less than 50% bootstrap support were collapsed. The different colors highlight different groups of AMT1 and AMT2 family.



These results indicated that the AcoAMTs proteins were clustered into two subgroups, AMT1 and AMT2, according to their affinity type. Group AMT1 is composed by transporters of high-affinity  $NH_4^+$  (represented in red, Figure 3) (NINNEMANN *et al.*, 1994; VON WIRÉN; MERRICK, 2004; YUAN *et al.*, 2007). According to the literature, the group AMT2 members (represented in black, Figure 3) are consists of representatives can participate in the to the transfer of net  $NH_3$ , however without the presence of current across the membrane (GUETHER *et al.*, 2009) and also are not permeable to the  $NH_4^+$  analog methylammonium

(SIMON-ROSIN *et al.*, 2003; review by SUN *et al.*, 2018).

To further confirm that there were two major AMT transporters groups and to study and understand the evolutionary relationships of the AMTs of other plants, we selected proteins sequences of 12 species (including 10 from *Glycine max*, 2 from *Vitis vinifera*, 11 from *Populus trichocarpa*, 8 from *C. canephora*, 8 from *Setaria italica*, 8 from *O. sativa*, 8 from *S. bicolor*, 9 from *Manihot esculenta*, 6 from *Cucumis sativus*, 7 from *Ricinus communis*, 8 from *A. comosus* L.), and we constructed a phylogenetic tree (Figure 4).

Figure 4. Phylogenetic tree analysis for all AMT proteins from 12 plants (A. thaliana, G. max, V. vinifera, P. trichocarpa, C. canephora, S. italica, O. sativa, S. bicolor, M. esculenta, C. sativus, R. communis, A. comosus L.), totally 99 protein sequences. Phylogenetic tree was constructed with the Neighbour-Joining (NJ) method using MEGA7.0 program with 1.000 bootstrap replicates. Branches with less than 50% bootstrap support were collapsed. The different colors highlight different groups of AMT1 and AMT2 family. The sequences of Α. comosus was obtained in Phytozome database L. (http://www.phytozome.net/pineapple.php) and other sequences were obtained based in von Wittgenstein et al. (2014) publication.



Genome chromosomal location showed that AcoAMT genes were randomly distributed on chromosomes (Figure 5). The largest number of AcoAMT genes occurred on chromosomes 1 and 13 (two AcoAMT genes), followed by one gene located on chromosomes 17 and 21. In addition, two genes are located on scaffold558 (one gene) and scaffold881 (one gene), in which they are represented by the unchromosome in Figure 5. Expansion analysis of the AcoAMT genes in the A. comosus L. genome was examined. Based on their chromosomal

distribution and the high rate of sequence similarity, we determined that two duplication pairs arose from segmental events, one between *AcoAMTa* and *AcoAMTg* genes and another between *AcoAMTb* and *AcoAMTf* genes in *A. comosus* L. (Table 3); the lines red in Figure 5 shows the connections among these paralogs. Our results indicated that these *AcoAMT* genes were possibly generated by gene duplication and the segmental duplication events played and a major driving force for *AcoAMT* expansion. The duplication of genes increases the functional divergence, which is an essential factor in adaptability under changing environmental conditions (CONANT; WOLFE, 2008).

We have calculated the divergence and evolutionary relationship by *Ka*, *Ks*, and *Ka/Ks* ratios between two paralogs *AcoAMT* genes. *Ka/Ks* ratio was 0 for *AcoAMTa* and *AcoAMTg* and 0.22 for *AcoAMTb* and *AcoAMTf* (Table 3). In general, *Ka/Ks* ratio less than 1, equal to 1, and greater than 1 means negative or stabilizing selection, neutral selection, and positive selection, respectively (ALI *et al.*, 2017). The *Ka/Ks* ratio trend in the coding sequences of two duplicate pairs indicated that the *AcoAMT* genes have undergone a positive Darwinian or purifying selection (*Ka/Ks* < 1). Positive selection of a gene during evolution means that it increases its potential and has more transcription levels under stress conditions (BOWERS *et al.*, 2003). Figure 5. Chromosomal distribution of AMT genes in A. comosus L. Chromosome number is labeled above the chromosomes. Red lines reflect segmental duplications. (Data extracted from Table 3).



**Table 3.** Duplication date of paralogous genes pairs in among *A. comosus* L. *AcoAMT* genes. *Ka* represents the non-synonymous substitution number per non-synonymous site, *Ks* is the number of the synonymous substitution site; *Ka/Ks* represents the ratio of non-synonymous (*Ka*) to synonymous (*Ks*) substitutions.

Paralogous Pairs	<b>Chromosomal location</b>	Duplication event	Ка	Ks	Ka/Ks	Selection
AcoAMTa/ AcoAMTg	Chr1/ scaffold_881	segmental	0.00	0.003	0	Purifying
AcoAMTb/ AcoAMTf	Chr1/Chr21	segmental	0.16	0.74	0.22	Purifying

To examine the origin and evolutionary history among A. comosus L. with other species (A. thaliana and O. sativa), a comparative analysis was performed to identify the orthologous pairs of AcoAMT genes (Figure 6; Table 4). A total of 5 AcoAMT genes showed a syntenic relationship with A. thaliana, followed by 8 in O. sativa (Table 4). AcoAMTh was found to be associated with five genes pairs in A. thaliana (AtAMT11. AtAMT12. AtAMT15, AtAMT13 and AtAMT14) and two genes pairs in O. sativa (OsAMT11 and OsAMT13), indicating that AcoAMTh gene might be derived from the same ancestor and arose before the divergence of eudicots and monocots. Indeed, we suggest that this gene may have played an essential adaptative role during evolution. Besides, we also found that all other AcoAMT genes were present in A. comosus L. and

were not found in Arabidopsis (AcoAMTa, AcoAMTb, AcoAMTd, АсоАМТс, AcoAMTe, AcoAMTf and AcoAMTg - Figure 6; Table 4), which may indicate that these genes were formed after the divergence of eudicot and monocot plants. Interestingly, we found an extensive genetic synteny between A. comosus L. and O. sativa, two monocot species. This was expected because the genome-sequenced CAM (pineapple) share conserved syntenic relationships with several important cereal species (like rice and sorghum) and has been regarded as an important crop for studying CAM photosynthesis and abiotic stress tolerance (MING et al., 2015).



Figure 6. Synteny between A. comosus L. and A. thaliana (A) and O. sativa (B). (Data extracted from Table 4).

Table 4. AcoAMT genes synteny among A. comosus L. and A. thaliana and A. comosus L. and O. sativa.

Ν	A. comosus	Chromosome Position	A. thaliana	Chromosome Position
1	AcoAMTh	scaffold_881:3627337760	AtAMT11	Chr4:78581837859918
2	AcoAMTh	scaffold_881:3627337760	AtAMT12	Chr1:2406077024062630
3	AcoAMTh	scaffold_881:3627337760	AtAMT15	Chr3:88014008802890
4	AcoAMTh	scaffold_881:3627337760	AtAMT13	Chr3:88056318807388
5	AcoAMTh	scaffold_881:3627337760	AtAMT14	Chr4:1416168114163195
Ν	A. comosus	Chromosome Position	O. sativa	Chromosome Position
1	AcoAMTb	LG01:1341228913419316	OsAMT31	Chr1:3773528037737641
2	AcoAMTb	LG01:1341228913419316	OsAMT33	Chr3:3523825535241976
3	AcoAMTc	LG13:5757858987	OsAMT13	Chr2:2469088424692884
4	AcoAMTf	LG21:980384598084333	OsAMT32	Chr2:2073407620737331
5	AcoAMTf	LG21:980384598084333	OsAMT31	Chr1:3773528037737641
6	AcoAMTf	LG21:980384598084333	OsAMT33	Chr3:3523825535241976
7	AcoAMTh	scaffold_881:3627337760	OsAMT11	Chr4:2550015825502633
8	AcoAMTh	scaffold_881:3627337760	OsAMT13	Chr2:2469088424692884

#### Conclusion

The current study identified eight ammonium transporters genes in the pineapple genome and provides an overview of the structure, phylogeny, chromosomal distribution, and synteny relationship of the *AMT* genes. This

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