Nitrogen Uptake in Plants: The Plasma Membrane Root Transport Systems from a Physiological and Proteomic Perspective

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Abstract: Nitrogen nutrition in plants is a key determinant in crop productivity. The availability of nitrogen nutrients in the soil, both inorganic (nitrate and ammonium) and organic (urea and free amino acids), highly differs and influences plant physiology, growth, metabolism, and root morphology. Deciphering this multifaceted scenario is mandatory to improve the agricultural sustainability. In root cells, specific proteins located at the plasma membrane play key roles in the transport and sensing of nitrogen forms. This review outlines the current knowledge regarding the biochemical and physiological aspects behind the uptake of the individual nitrogen forms, their reciprocal interactions, the influences on root system architecture, and the relations with other proteins sustaining fundamental plasma membrane functionalities, such as aquaporins and H+-ATPase. This topic is explored starting from the information achieved in the model plant Arabidopsis and moving to crops in agricultural soils. Moreover, the main contributions provided by proteomics are described in order to highlight the goals and pitfalls of this approach and to get new hints for future studies.

Keywords: plant mineral nutrition; organic nitrogen; metabolic networks; glutamate; subcellular proteomics

1. Introduction

Nitrogen (N) is the most abundant mineral element present in plant tissues, in which it constitutes about 1 to 5% of total dry matter [1]. Plants acquire N by roots throughout the life cycle, and the availability of this macronutrient, in terms of total amount and forms, deeply affects plant development and interactions with the environment [2]. In cultivated soils, N availability is a key factor often limiting crop productivity. Hence, there is a worldwide massive use of N fertilizers, despite the detrimental effects on ecosystems and high socioeconomic costs [3]. Improvement of the current knowledge about N nutrition in plants is, therefore, required to reduce the impact of these anthropogenic activities on a global scale.

Leaving aside protein-humic complexes not directly bioavailable to plants, in soil N is present as inorganic forms, such as nitrate (NO₃⁻) and ammonium (NH₄⁺), and as organic forms, mainly consisting of urea, free amino acids, and short peptides. The accessibility of these resources by roots varies considerably through space and time, due to soil heterogeneity and to dynamic microbial conversions, two aspects in turn affected by agronomic practices and environmental conditions [4]. In aerobic soils, NO₃⁻ is the most abundant form, with concentrations between 1 to 5 mM, while NH₄⁺ concentration typically ranges between 20 and 200 μM. However, NO₃⁻ is readily leached, while NH₄⁺ is strongly adsorbed by soil particles and slowly released [5]. Differently, free amino acids
and urea are generally present in concentrations ranging from 1 to 150 μM and <70 μM, respectively [5–7], representing a minor proportion of the available N for crops.

The relevance of organic N for crop nutrition is a matter of debate. Although there is evidence that plants can acquire amino acids, small peptides, and (partial) proteins from the soil [8], a high impact in agricultural contexts was traditionally ruled out [5,9]. However, in recent years, many studies opened new questions. First observations proved that the supply of sole amino acids sustains plant growth, and were soon followed by the characterization of root transport systems for amino acid uptake, whose molecular bases were partly elucidated in Arabidopsis (Arabidopsis thaliana L.) and confirmed in crops [10,11]. At the same time, it was proven that the provision of amino acids, even at a very low concentration, can affect root morphology and plant growth [12]. Similarly, considering that urea-based formulations account for over 50% of total N fertilizers applied in agriculture, the discovery of urea transporters in the plasma membrane (PM) of root cells has drawn attention to a direct use of this nutrient by plants [13].

Total N availability, and the forms supplied, affect seed germination, plant growth, root and leaf functionalities, hormonal balance, and seed production. Recent literature stresses the importance of interpreting N nutrition as a composite scenario, taking into account the contribution of individual nutrients, their reciprocal interactions, and the plethora of effects on plant metabolism [14,15]. Needless to say, the “‐omic” approaches, given their intrinsic holistic nature, seem to adequately respond to this need.

In roots, the major adaptations to N availability consist of the changes in uptake activity and in the modulation of the root system architecture (RSA), both of which are related to the ability of N forms to act as nutrients and/or regulatory signals for plant growth and metabolism (Figure 1).

![Figure 1](attachment:image.png)

**Figure 1.** The figure summarizes the regulatory pathways and the metabolic relations among the main transporters involved in N uptake by roots, taking as a model Arabidopsis. For simplicity, transporters, transceptors, aquaporins, and H+‐ATPase are reported in the same root cell, but it is not the common real case. Acronyms and details are explained in the text.
Transport and sensing of N forms involve proteins located in the PM of root cells. In recent years, there was a huge increase in knowledge about the control of uptake at the transcriptional level, as well as about the components involved in signaling. It was also highlighted that post-translational modifications (PTMs) of transporters, such as phosphorylation events and formation of protein complexes, have key roles in rapid adaptations to sudden changes in N availability [11,13,16,17].

From a physiological perspective, N uptake is also related to other main PM activities, such as the formation of the electrochemical proton gradient and water homeostasis. Hence, this review provides an overview of N nutrition in plants, trying to integrate the notions about single N forms, their interactions, as well as the relations with other PM functionalities (Figure 1).

Many studies have been focused on the identification of proteins involved in N uptake, reaching the most complete molecular characterization in Arabidopsis. We used Arabidopsis as a starting reference point, but several parallelisms in crops are reported. However, the description of these aspects in legumes (*Fabaceae* spp.) and actinorhizal crops, as well as in rice (*Oryza sativa* L.), is out of our scope due to their numerous peculiarities.

This topic is examined describing the contribution of proteomics, in order to highlight goals and pitfalls and to get new hints for future studies. Deciphering how the root membrane proteome changes in response to different N sources could provide new knowledge useful to enhance agriculture sustainability.

2. Transporters and Transceptors Involved in Nitrogen Uptake by Roots

The uptake of N nutrients is finely controlled and influenced by the interplay among three main protein classes, namely transporters, receptors, and transceptors. In this review, the term transporter is used as a synonym of selective carrier proteins, while the term transceptor refers to proteins able to fulfil a dual transport/sensing function [18]. Finally, it is worth remembering that the total concentrations of N forms in root cells depend on external availability as well as on the balance among net uptake (defined as the difference between influx and efflux), root metabolism, vacuolar accumulation, and xylem/phloem (un)loading [1].

2.1. Nitrate Uptake

Nitrate is the primary N source for plant growth in most agricultural soils, and its availability significantly affects crop productivity [9]. At high external supplies, total NO$_3^-$ concentration in root cells can reach up to 100 mM, mostly stored in the vacuole, while cytosolic concentration is kept low, as measured by selective-microelectrodes in cereals [19]. Electrophysiological studies indicated that NO$_3^-$ uptake by roots is always an active process, mediated by a 2H$^+$/1NO$_3^-$ symport mechanism, while NO$_3^-$ efflux is passive, saturable, and inducible [20,21].

In plants, NO$_3^-$ uptake is mediated by transporters of the NPF family (previously named NRT1/PTR family) and of the NRT2 family, which in Arabidopsis consist of 53 and 7 members, respectively [22,23]. Although there are no sequence homologies between the two families, these transporters share the same topology consisting of 12 transmembrane domains and have both the N- and C- termini lying on the cytosolic side of the membrane [24]. The activity of these proteins is strictly regulated by NO$_3^-$ availability and plant N nutritional status. In N starved plants, renewed availability of NO$_3^-$ triggers the typical NO$_3^-$ primary response (NPR), which comprises the rapid induction of NO$_3^-$ transporters and N assimilating enzymes. This adaptation is followed by later down-regulation of uptake, correlated with the accumulation of NO$_3^-$ itself and of its downstream metabolites, such as glutamine (Gln, Figure 1) [25]. A prominent role for the transcriptional control of NO$_3^-$ transporters was demonstrated in Arabidopsis and crops, and the importance of PTMs was also highlighted [26–28].
In Arabidopsis, three members of the NPF family are located in the PM of root cells and participate in NO\textsuperscript{-} uptake (Table 1).

Table 1. Transporters involved in N uptake in roots of Arabidopsis thaliana. The table reports the main information about transporters, family, affinity range, and tissue specificity in roots.

| N Forms | Family | Protein | Affinity | Function | Tissue Specificity in Roots | Ref |
|---------|--------|---------|----------|----------|-----------------------------|-----|
| NPF     | NRT1.1 | Dual    | Uptake   | NO\textsuperscript{-} transporter | Primary root tip, emerging lateral roots, epidermis near root tips, cortex, and endodermis in the mature portion of the root | [29,30,31,32] |
|         | NRT1.2 | Low     | Uptake   | Root     | Root hairs and epidermis    | [33] |
|         | NAXT1  | Low     | Efflux   | Cortex   | Cortex of mature roots       | [34] |
| Nitrate | NRT2.1 | High    | Uptake   | NO\textsuperscript{-} transporter (?) | Epidermis, cortex, and endodermis in the mature portion of root | [32,35] |
|         | NRT2.2 | High    | Uptake   | Root     | Root                         | [36] |
|         | NRT2.4 | High*   | Uptake   | Epidermis of lateral roots     | [37] |
|         | NRT2.5 | High*   | Uptake   | Root hairs, epidermis, and cortex | [38] |
| Ammonium| AMT1;1 | High    | Uptake   | Root tip, root hairs, epidermis, and cortex | [39] |
|         | AMT1;2 | High    | Retrieval from apoplast | Endodermis near the root hair differentiation zone, cortex in the zone of emerging lateral root | [40] |
|         | AMT1;3 | High    | Uptake   | NO\textsuperscript{+} transporter (?) | Root tip, root hairs, epidermis, and cortex | [39] |
|         | AMT1;5 | High*   | Uptake   | Root tip, root hairs, epidermis | [40] |
|         | LHT1   | High    | Uptake   | In young seedlings in the epidermis of later and emerging roots; in older plants, in root tip | [41,42] |
|         | LHT    |         |          | Root hairs, epidermis, cortex, endodermis | [7] |
|         | LHT6   | High    | Uptake of neutral and acidic amino acids, His, Ala, (Phe?)  | | |
| Amino acids | AAP5   | High    | Uptake of Arg and Lys | Cortex | [42,43] |
|         | AAP1   | High    | Gln, Ala, Gln, Pro, Ser | Root tip, root hairs, epidermis, cortex, endodermis, and vascular cylinder | [7,44] |
|         | ProT   | Low**   | Pro and glycine betaine | Epidermis and cortex | [45] |
| Urea    | SSS    | High    | Uptake   | Epidermis, cortex, vasculature tissues near the xylem | [46] |
|         | DUR3   |         |          |                      | |

\* very high affinity; " the activity was tested only in high availability of Pro, to our knowledge, the high affinity could not be excluded. For acronyms see the text.

NRT1.1 (AtNPF6.3) and NRT1.2 (AtNPF4.6) participate in NO\textsuperscript{-} influx, and the second is the major one responsible for the constitutive influx in the low-affinity range (>0.25 mM) [33], while NAXT1 (AtNPF2.7) mediates NO3\textsuperscript{-} efflux to the external medium [34].

The characterization of NRT1.1, as the first transceptor identified in higher plants, represents a milestone in plant nutrition research [29]. NRT1.1 is an inducible dual-affinity transporter able to participate in the uptake of NO3\textsuperscript{-} both in the low-affinity (Km~4 mM) and high-affinity (Km~50 \textmu M) ranges, depending on the dephosphorylation/phosphorylation of threonine (Thr)-101 [30]. Interestingly, the determination of the crystal structure of NRT1.1 suggested that phosphorylation also influences the oligomerization state of the protein in the PM. According to the proposed model [47], at high external concentrations of NO3\textsuperscript{-}, the absence of phosphorylation of Thr-101 permits that NRT1.1 forms dimeric complexes operating in low-affinity mode. Instead, when NO3\textsuperscript{-} concentration is low (<0.2 mM) [30], the phosphorylation of NRT1.1 determines dimer decoupling, by which individual protomers adopt a high-affinity transport mode. The phosphorylation of Thr-101
is mediated by CIPK23 kinase (CBL-interacting serine/threonine-protein kinase 23). In detail, CIPK23 is recruited at the PM by binding with one of the two proteins CBL9 or CBL1 (calcineurin B-like 1 or 9), which also enables its activation by auto-phosphorylation (Figure 2A) [48].

According to its definition as a transceptor, NRT1.1 plays a key role as an NO\textsuperscript{−} sensor, showing a signaling function that is independent of the transport activity. NRT1.1 regulates a very ample set of plant responses to NO\textsuperscript{−}, by a signaling cascade which is partly elucidated [16]. At the root level, these responses include both changes in RSA (Section 3) and the regulation of NO\textsuperscript{−} uptake (Figure 1). Once again, the functionality of NRT1.1 as an NO\textsuperscript{−} sensor (and of its concentration) depends on its phosphorylation state. For instance, the phosphorylated and not-phosphorylated forms participated in the short-term biphasic up-regulation of the NRT2.1 gene (see below) in response to low and high external NO\textsuperscript{−} inputs, respectively, while the phosphorylated one mediates the long-term down-regulation of NRT2.1 at high NO\textsuperscript{−} concentrations [29,49].

![Figure 2. The model proposed for the NRT1.1 dimerization mediated by phosphorylation (adapted from [47]) (A). Model proposed for the tetramer NRT2.1/NAR2.1 according to [22,50] (B). Details are explained in the text.](image)

Among the NRT2 family in Arabidopsis, four members were characterized as responsible for the high-affinity influx of NO\textsuperscript{−} in roots (<0.25 mM, Table 1). NRT2.1 and NRT2.2 are the major components of the inducible HATS (High Affinity Transport System), being highly induced at the transcriptional level in roots by exposure to NO\textsuperscript{−} [36]. NRT2.4 and NRT2.5 are characterized by a very high-affinity and are strongly induced by N deprivation but rapidly repressed by NO\textsuperscript{−} and NH\textsubscript{4}\textsuperscript{+}, suggesting their primary role in the earliest influx of NO\textsuperscript{−} after starvation [37,38,51].

The molecular mechanisms governing NRT2.1 were largely described. The transcription of the NRT2.1 gene is rapidly induced by NO\textsuperscript{−} in an NRT1.1-dependent manner (see above), at a low or high level according to the concentration of the anion [29]. Moreover, it is repressed by Gln and NH\textsubscript{4}\textsuperscript{+}, and diurnally regulated, probably by a carbohydrate pool translocated from the shoot (Figure 1) [24]. In addition, NRT2.1 was proposed as a potential NO\textsuperscript{−} transceptor, involved in the modulation of RSA (Section 3). The recruitment of NRT2.1 to the PM requires binding with the protein NAR2.1 (Nitrate Assimilation Related protein), forming a stable tetramer composed of two subunits for each protein (Figure 2B) [50]. However, in Arabidopsis, the dependency of functionality on NAR2.1 was demonstrated for NRT2.5 and excluded for NRT2.4 [37,51], but the physiological meaning is not yet elucidated. Interestingly, in recent years, some proteomic investigations contributed to discovering additional molecular events that regulate NRT2.1. First indications were provided by a phosphoproteomic study aimed at the characterization of the early changes.
induced by NO₃⁻ or NH₄⁺ resupply to Arabidopsis seedlings, based on the phosphopeptide enrichment by the titanium dioxide methodology. This approach revealed that NRT2.1 is phosphorylated in serine (Ser)-28 in N starved seedlings and is rapidly dephosphorylated after resupply of high NO₃⁻ concentrations, suggesting that this mechanism contributes to inactivate NRT2.1 when the activity of low-affinity transporters becomes predominant [52]. Later on, it was clarified that the phosphorylation at Ser-28 contributes to increase the stability of NRT2.1 and is required for the accumulation of this protein in response to low NO₃⁻ availability [53]. Furthermore, by a phosphoproteomic approach specifically devoted to NRT2.1 characterization, it was recently proven that NRT2.1 can be also phosphorylated in the C-terminus (at Ser-510). This PTM leads to NRT2.1 inactivation in response to high N supplies, without affecting its interaction with NAR2.1, proving a novel and fundamental mechanism for the regulation of NO₃⁻ uptake in Arabidopsis roots [54].

Although the huge amount of information obtained in Arabidopsis is exciting, it is important to remember that direct parallelisms in crops must be done with caution. For instance, a recent characterization of ZmNPF6.6, an NRT1.1 homolog in maize (Zea mays L.), confirmed the induction by NO₃⁻ and the involvement in its low and high-affinity uptake. However, the biphasic kinetic, the regulation by phosphorylation, and the auxin transport were not confirmed [55]. Similarly, the literature indicates that in crops the binding of NRT2 could involve different members of the NAR family, since partner pairs are species-dependent [56].

2.2. Ammonium Uptake

Ammonium is an important N nutrient rapidly absorbed and assimilated by plants, as well as a signaling molecule influencing plant growth and RSA (Section 3). However, many crops are sensitive to NH₄⁺ toxicity, especially at high dosages, and generally results in stunted growth, leaf chlorosis, and poor root development. Indeed, for optimal growth most crops require the contemporaneous availability of NO₃⁻ and NH₄⁺, even if the best ratio depends on plant species, developmental phase, and environmental conditions [9,57].

In plants, the uptake of NH₄⁺ and its allocation among organs are highly influenced by nutritional conditions. At the cellular level, it was estimated that NH₄⁺ concentrations in cytosol and vacuole range from 1 to 10 mM and 1 to 45 mM, respectively [58], while, interestingly, in roots, the apoplastic NH₄⁺ concentration is buffered around 1 to 2 mM, both under low and high NH₄⁺ supplies [40].

At higher external NH₄⁺ concentrations, there is evidence that the nutrient probably permeates into root cells both as NH₃ and NH₄⁺ through two distinct mechanisms. The first one is the passive and electroneutral influx/efflux cycle of NH₃, putatively facilitated by aquaporins (Section 4.1), that results in the hyper-accumulation of the charged form into the vacuole. The second one includes the NH₄⁺ influx likely mediated by other PM channels, such as non-selective cation (NSCC) and potassium (K⁺) specific channels, among which AKT1 (Figure 1) [57,59].

At low external NH₄⁺ concentrations (<1 mM), NH₄⁺ influx is a saturable and highly controlled process, mediated by members of the AMT1 subfamily (of the Ammonium Transporter/Methylammonium Permease family) through NH₄⁺-uniport or NH₃/H⁺ co-transport. These are 45 to 65 kDa proteins with 11 hydrophobic transmembrane domains, a cytosolic C-terminus and an N-terminus in the apoplast [58,60,61]. In Arabidopsis, four members of the AMT1 family are responsible for high-affinity NH₄⁺ (<1 mM) uptake in the roots (Table 1).

In detail, AMT1;1, AMT1;3 and AMT1;5 are accumulated in the epidermis and root hairs, which contribute in an additive manner to NH₄⁺ uptake via the symplastic route [39,40]. However, the different substrate affinities suggest that AMT1;1 and AMT1;3 (Kₘ ~50 and 60 μM, respectively) mainly operate at NH₄⁺ concentrations common in soil, while AMT1;5 (Kₘ ~5 μM), which is accumulated only under prolonged N starvation, might
significantly contribute when the availability of NH₄⁺ is very low [40]. Finally, since AMT1;2 has a lower affinity (Km ~230 μM) and is accumulated in the endoderm, it is thought to be mainly involved in the retrieval of NH₄⁺ that enters root through the apoplastic route [40].

At the transcriptional level, the AMT1 genes are generally subjected to diurnal changes, probably regulated according to the rate of carbohydrate translocation from the shoot. Moreover, AMT1 gene expression is induced during N starvation and reduced under high NH₄⁺ availability, probably through mechanisms exerted by its downstream metabolites, such as Gln (Figure 1) [40,58,62].

At the protein level, these transporters form trimeric complexes localized in the PM [63], for which post-translational regulations were suggested already from the earliest molecular studies [62]. In recent years, several proteomic investigations have significantly contributed to elucidate this aspect, pointing out the key role of phosphorylation at the C-terminus in AMT1 subunits. Firstly, a phosphoproteomic study, conducted by IMAC methodology (Ion Metal Affinity Chromatography) on PMs enriched fraction from cell suspensions of Arabidopsis, identified a phosphorylation site in the C-terminus of AMT1 conserved in three members of the family [64]. Then, studies on AMT1;1 and AMT1;2 proved that the C-terminus of the AMT1 subunit acts as an allosteric regulator of the complex. According to the proposed molecular model, in the non-phosphorylated form, the C-terminus interacts within its own monomer and with the adjacent one, assuring the transport. Its phosphorylation (at Thr-460 in AMT1;1) leads to trans-inactivation of the whole complex (Figure 3) [63,65].

Later on, a dedicated study on Arabidopsis roots was conducted combining phosphoproteomics with complementary analyses, such NH₄⁺ pulse treatments, determination of ¹⁵N-NH₄⁺ uptake, and protein blots [66]. This combined approach proved that phosphorylation of Thr-460 in AMT1;1 is induced by NH₄⁺ in a time- and concentration-dependent manner, while neither NO₃⁻ nor Gln (nor endogenous nor external) trigger this response, overall proving a feedback mechanism able to tune NH₄⁺ uptake capacity to prevent toxicity [66]. The same study proposed that AMT1;1 could act as a transceptor, but this hypothesis was successively questioned. For instance, a comparative study of proteomic and transcriptomic profiles of Arabidopsis roots within three hours of NO₃⁻ or NH₄⁺ deprivation indicated that the dephosphorylation of AMT1;1 was not concomitant with large transcriptomic changes, suggesting that the predominant regulative signal is the endogenous NH₄⁺ concentration [67]. However, the same study paved the way to discover the putative roles of AMT1;3 as the transceptor involved in the modulation of RSA (Section 3).

Other studies revealed an additional degree of complexity in the NH₄⁺ uptake regulation. In Arabidopsis roots, it was shown that AMT1;1 and/or AMT1;3 interact in functional homo- and heterotrimers, both subjected to (trans)-inactivation exerted by the phosphorylation of the AMT1;3 subunit [68]. The same study has also put in evidence that the phosphorylation in AMT1;3 does not affect the functionality of AMT1;2 or AMT1;5, suggesting different regulative pathways for individual AMT1 transporters. In the same years, the above-cited proteomic investigations on NO₃⁻/NH₄⁺ resupply or deprivation in Arabidopsis identified novel phosphorylated sites in the C-terminus of AMT1;1 and AMT1;3 differently modulated in response to NO₃⁻ [52,67]. The molecular network behind was partly elucidated a few years ago. Firstly, a large-scale genetic screening in Arabidopsis led to the discovery that the phosphorylation at the C-terminus of AMT1;1 and AMT1;2 is catalyzed by the CIPK23/CBL1 complex in response to high NH₄⁺ availability, although the involvement of other kinases was not excluded (Figure 3) [69].
In Arabidopsis the CIPK23/CBL1/9 complex is also involved in the positive regulation of the K+ transporter AKT1 [48], which in barley (*Hordeum vulgare* L.) was suggested as one of the unspecific low-affinity importers of excess NH₄⁺ [70]. From a physiological point of view, this mechanism is very interesting because it could contribute to enhancing the NH₄⁺ uptake, and hence N resources for plant growth, when the risk of toxicity is avoided by the co-presence with NO₃⁻. Moreover, this novel scenario highlights the interplay of the molecular pathways controlling the uptake of NO₃⁻, NH₄⁺, and K⁺ (Figure 1). Proteomic studies highlighted a major accumulation of the K⁺-transporter HAK5 in roots of Arabidopsis plants when grown in NH₄⁺ instead of NO₃⁻, suggesting a compensatory response [67]. Similarly, in maize and potato roots, an increase in the accumulation of the voltage-gated potassium channel beta subunit was observed in response to NO₃⁻ availability [71,72].

The study of Straub and co-workers also revealed that the phosphorylation of AMT1;3 (Thr-464) is mediated by a kinase not yet identified, evidencing the involvement of different kinases in the regulative pathway of AMT1 [69]. Moreover, it was recently demonstrated that AMT1;3 can be phosphorylated in additional sites (Ser-480, Ser-487, Thr-494) in response to NO₃⁻ availability (Figure 1). In particular, the activity of AMT1;3 (dephosphorylated in Thr-464) can be further increased by dephosphorylation of Thr-494 [73]. Overall, these observations point out how much the NH₄⁺ uptake is finely regulated, proving its relevance in plant physiology.

The molecular determinants of NH₄⁺ uptake are generally conserved across an ample set of plant species, but significant differences were reported in the transcriptional regulation of AMT1 among crops [17]. For instance, in several cereals, the supply of NH₄⁺ triggers an increase in the NH₄⁺ influx, which is kept stable for a few hours (even for 24 h in maize) before the feedback negative regulation occurs. In maize, this response was associated with a peculiar induction of ZmAMT1;1 and ZmAMT1;3 triggered by a local NH₄⁺ signal, independent from plant nutritional status, but the complete elucidation of the regulative events is not yet elucidated [74]. Overall, considering the ample range of degrees in adaptability/sensitivity to NH₄⁺ of crops and cultivars [9], the study of NH₄⁺ nutrition in plants is still an open, and partially unexplored, research field.

### 2.3. Uptake of Amino Acids

In plant tissues, the content of total amino acids usually ranges between 1 and 10 mM, but this value is very variable and can highly increase, especially under high NH₄⁺

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Figure 3. Schematic representation of the AMT1 trimeric complex and its control mode through CIPK23/CBL1-mediated phosphorylation induced by high NH₄⁺ availability, adapted from [69] (see text for further details).
nutrition [9]. Similarly, the distribution of total amino acids in plant cell organelles is variable, with the highest concentration typically found in plastids and cytosol [75,76]. In plant tissues, it is possible to distinguish the group of the “major” amino acids (i.e., more abundant, including glutamate (Glu), Gln, aspartate, asparagine, alanine (Ala)) and the group of the “minor” ones, present at lower levels and whose biosynthesis is subjected to a strong end-product feedback control [77].

As previously stated, amino acid concentration in soils ranges between 1 and 150 μM and, once again, the “major” amino acids are the most abundant [7]. The soil amino acid content is highly influenced by environmental conditions, microorganism activities, and soil buffer capacity that renders basic amino acids (arginine (Arg), lysine (Lys), and histidine (His)) less mobile and less available for roots [5,9]. However, considering the intense exchanges of organic exudates between microorganisms and roots, it is possible that the amino acid compositions in the rhizosphere and in bulk soil significantly differ [10].

The effectiveness of amino acids as N nutrients for plants is a controversial issue. The major objections reside in the low diffusion coefficient of amino acids in the soil as well as in their short half-life (about 4 h), two factors that could limit the competitiveness of plants versus microorganisms in the uptake of these compounds [9]. On the other hand, there is increasing evidence that plants acquire amino acids from the growing media, also when both NO\textsuperscript{3} and NH\textsubscript{4} are available [7]. Physiological studies on crops provided contradictory results. For instance, barley plants, grown in hydroponics and supplied with amino acid concentrations similar to field conditions, showed an uptake rate with Michaelis-Menten kinetics [78]. Conversely, a \textsuperscript{14}C/\textsuperscript{15}N tracer study in maize seedlings, grown in rhizosphere tubes filled with soil and treated with Ala or NO\textsubscript{3}, showed a scarce relevance of organic N uptake [79]. In our opinion, some aspects should deserve specific attention. Firstly, the studies conducted in hydroponics or in experimental media generally provide results supporting the nutritional value of amino acids, while those conducted in soil conditions give opposite results. It is, therefore, plausible that some results are biased due to the interferences of soil particles and to the experimental set-up that somehow could modify the expected amino acid bioavailability. Secondly, the provision of a single amino acid could be misleading because one amino acid is not the other, nor mimics the soil chemical composition.

In Arabidopsis, amino acids prompt different effects when provided at relatively high concentrations (3 mM). Some of the “major” amino acids promote plant growth, Glu has no effects, while the supply of Arg or Ala is positive singly but negative when in combination with NO\textsubscript{3}. Finally, the provision of valine (Val), Ser, and isoleucine (Ile) seem to be detrimental to plant growth [80]. These observations support the use of amino acid by plants, hint interlinks with the inorganic N nutrients, and suggest that “minor” amino acids, when provided in high concentrations (not common in soil), could be toxic due to metabolic interferences [10].

Some high-affinity amino acid transporters, with kinetics comparable to those of microorganisms, have been characterized in the last years [10,81]. In the Arabidopsis genome, more than 100 genes encoding for amino acid carriers were predicted, which belong to the amino acid/polyamine/organocation (APC) superfamily [82] and to the UMAMIT (Usually Multiple Amino acids Move In And Out Transporters) group of the drug/metabolite transporter superfamily [83]. Considering the complexity of amino acid metabolism in plants, the tissue specificity and cell localization of these transporters is fundamental to define their actual physiological functions [11,77].

To date, five carriers that were characterized as amino acid importers in roots of Arabidopsis, all comprised in the AAAP (amino acid/auxin permease) family of the APC superfamily (Table 1). Gene expression in heterologous systems together with complementation studies in yeast characterized these carriers as electrogentic H\textsuperscript{+}/amino acid symporters. Moreover, the structural analyses of AAAP1 (Amino Acid Permease 1), taken as a model, showed that these proteins contain 11 trans-membrane domains with the N-termi-
nus inside the cytoplasm and the C-terminus facing the outer PM surface [84]. The characterization of Arabidopsis mutants fed with amino acids at low concentrations, together with β-glucuronidase and GFP-tagging studies, provided information about the functionalities of these carriers in plants, their expression patterns, and their location in the PM. In detail, LHT1 (Lysine Histidine Transporter 1) is a high-affinity carrier involved in the uptake of neutral amino acids, His, and acidic amino acids, located in the root and leaf cells [41,42]. AAP5 mediates 68 to 88% of the uptake of Lys and Arg in the high-affinity range and, despite being present in all plant tissues, is preferentially accumulated in the root cortex [42,43]. In addition, LHT6, a carrier localized in root hairs, epidermis, cortex, and endodermis, participates in the high-affinity uptake of acidic amino acids, Ala, Gln, and probably phenylalanine (Phe). The same authors also proposed that AAP1, localized in cotyledons and roots, could mediate the high-affinity uptake of Ala, Gln, proline (Pro), Ser, and Glu, although the need for verification was stated [7,44]. Finally, ProT2 is an importer of Pro and glycine betaine located in the root epidermis and cortex, but in Arabidopsis the activity in the high-affinity range (500 μM) was proven only for the second compound [45].

Amino acid efflux is fundamental in the rhizosphere relations, as demonstrated by its change in response to microbial compounds [85] and nutrition limitations [86]. However, its quantitative significance in normal conditions was doubted by several authors, which ascribed it to a small leakage down a concentration gradient [10]. The analyses of root exudate in several crops in non-axenic condition indicated a predominant extrusion of Gly, Ser, and Ala, supporting the involvement of selective transport systems [87]. In recent years, the UMAMIT family was characterized as a class of facilitators with bidirectional properties, and several members are expressed in roots [11], but to date members specifically involved in amino acid exports into the soil are not yet identified [88].

Considering this framework, it is highly conceivable that these transporters have complementary roles in amino acid uptake from the soil, with individual contributions probably varying during plant development and growth conditions. Additionally, the presence of LHT1 and AAP5 in the aerial organs [41,43], as well as the localization of AAP1 in the cotyledon vasculature [44], supports their involvements in other processes of amino acid transport and allocation throughout the plant. For instance, in seedlings LHT1 is expressed in the root epidermis, but it is confined to the root tips in older plants (Table 1). Similarly, AAP5 is highly expressed in roots of young plants [89], but in older plants, its expression seems to become predominant in aerial organs [90]. From these considerations, it was proposed that the contribution of LHT1 and AAP5 in amino acid uptake by roots is substantial during the early developmental stages but could become indirect later [91]. Currently, there is no evidence for the existence of amino acid transceptors in plants [92].

In Arabidopsis seedlings, the expression of LHT1 was found to be inducible by amino acids (5 mM) and by NO₃⁻ (5 mM) if compared to plants supplied with NH₄NO₃ (40 mM) [41]. Although these results are indicative, the high concentrations used do not allow to predict the responses in field conditions. Differently, AAP1 resulted to be induced by light, sugars, amino acids, NO₃⁻, and also by NH₄⁺. The studies on ProT2 were mainly focused on its induction by water and salt stress, which was confirmed in different crops, but little is known about the effects of N nutrients [93]. Gene expression analysis of LHT1, AAP1, and ProT2 in Arabidopsis root and shoot in response to NO₃⁻ induction revealed different regulation depending on timings, doses, and organs. In the long-term (24 h), AAP1 resulted slowly induced equally in both organs, ProT2 seemed slightly de-induced and LHT1 was highly induced, especially in shoots [93]. This scenario depicts the complexity of studying these genes in the context of plant N nutrition (Figure 1). As highlighted by the literature [7], a major limitation consists in the different roles in roots (i.e., amino acid uptake from the soil) and shoots (i.e., supply of amino acids to mesophyll cells from the xylem sap). These functionalities could differently respond to N inputs as well as could be differently affected by factors, including photosynthetic activity and energy metabolism,
with a final outcome that is very challenging to discern the regulatory network. However, an increasing interest is directed to the AAAP family, and, in our opinion, recent genome-wide analyses, such as those conducted in maize [94], potato (Solanum tuberosum L.) [95], tobacco (Nicotiana spp) [96], and wheat (Triticum aestivum L.) [97] are paving the way to unravel this issue in the coming years. On the contrary, proteomics was very rarely applied in the field of amino acid nutrition in plants. To our knowledge, two research projects were conducted about the responses to Gly-based nutrition in Lolium perenne and Brassica campestris [98,99]. Although these pioneering studies revealed new information about Gly metabolism in plants, no amino acid transporters were detected among the responsive proteins. This lack is in part attributable to the 2-DE based analytical approach adopted, known to be unsuitable for resolving membrane proteins (Section 5, Table 2). Indeed, very recently an iTRAQ-based study (Isobaric Tags for Relative and Absolute Quantitation) allowed to characterize LHT1, confirming its key role in N remobilization in Arabidopsis seedlings [100]. That hints that the right time has come for proteomics to help unravel the complex metabolic network behind amino acid nutrition in plants. Future studies could obtain new information useful to ameliorate soil management and agricultural sustainability.

2.4. Urea Uptake

In the last decades, the use of urea fertilizers is increased to account for more than 50% of the world N fertilizer applications, mainly thanks to low costs and high N content. In soil, urea is rapidly hydrolyzed by ubiquitous microbial urease with the release of NH$_3$, subsequently converted into NO$_3^-$ by nitrifying bacteria [101]. Therefore, in agricultural soil, urea concentration is generally low (up to 70 μM) and it was thought for a long time that this kind of fertilizer provided N to crops mainly in the form of NH$_3$. However, the identification of a dedicated high-affinity transporter led to the re-evaluation of urea as a direct N source [102]. In addition, urea is a key molecule in N translocation and recycling in plants. In source tissues, urea is produced by Arg catabolism and is hydrolyzed by cytosolic urease to release NH$_3^-$, which is then re-assimilated in sinks. In this view, the presence of endogenous urease activity in almost all plant tissues reinforces the hypothesis that plants can use the molecule as a nutrient [103].

The first urea transporter identified in higher plants was DUR3 in Arabidopsis, a high-affinity urea/H$^+$ symporter that belongs to the SSS (Sodium-Solute Symporters) family (Table 1). AtDUR3 is an integral protein with 14 transmembrane domains and the N- and C- termini protruding into the outer side of the membrane [104]. In Arabidopsis roots, DUR3 localizes to the PM and sustains up to 90% of urea influx in the high-affinity range (K$_m$ ~4 μM), while the urea influx in the low-affinity range was ascribed to diffusion throughout aquaporins (Section 4.1) [46].

When provided as the sole N source in axenic conditions, urea is taken up as an intact molecule, partly translocated to the shoots, and partly assimilated into amino acids, despite being less efficient than inorganic N forms. These metabolic responses are associated with an increase in expression of AtDUR3 and of genes involved in amino acid metabolism and transport [105]. Interestingly, the urea contents in roots scarcely differ in plants supplied with urea or with inorganic N, suggesting that the endogenous urease activity is sufficient to metabolize the urea taken up from growing media [46,103,105].

The expression of the AtDUR3 gene is high during N starvation, dramatically induced further after resupply of urea, but repressed after resupply of NH$_3^-$ or NO$_3^-$. GUS-promoter analyses showed that the expression of AtDUR3 significantly increases during N starvation in the epidermis and cortex, and to a lesser extent in the vasculature tissues near the xylem (Table 1; Figure 1) [46].

Physiological and transcriptomic studies in Arabidopsis plants confirmed that urea uptake is stimulated by substrate and reduced when inorganic N is available [105]. Further studies in maize and wheat agree in the observation that co-provision of urea and NO$_3^-$ has overall positive effects on plant growth and N use. The explanation resides in a
reprograming of the assimilation pathways that seems to assure a better metabolic balance [106]. This behavior is very similar to what is described under the co-provision of NO\textsuperscript{−} and NH\textsubscript{3}\textsuperscript{+} by physiological [9] and proteomic studies. For instance, a recent characterization of proteomic profiles in roots and leaves of maize showed that the co-provision of NO\textsuperscript{−} and NH\textsubscript{3}\textsuperscript{+} is related to specific changes in the abundance of enzymes involved in C and N metabolism, water balance, and stress responses [71]. In the future years, it will be of interest to investigate distinct signatures between urea and NH\textsubscript{3}\textsuperscript{+} in co-presence with NO\textsuperscript{−}.

To date, DUR3 orthologues were identified in several crops, indicating that higher plants have only a single DUR3 gene [13]. The functional characterization of DUR3 in maize confirmed the main features of its involvement in urea uptake in roots [107] and suggested a role in urea vascular loading in leaves [108]. Interestingly, a recent study in tomato revealed different DUR3 transcript abundances in cultivars showing different nitrogen use efficiency (NUE) [109].

To the best of our knowledge, no proteomic study investigated urea nutrition in plants. However, Menz and co-workers detected that in Arabidopsis, DUR3 was more abundant under NH\textsubscript{3}\textsuperscript{+}-adapted conditions [67]. Although this result is apparently in contradiction with DUR3 expression studies, it could suggest possible changes in Arg catabolism after long exposure to NH\textsubscript{3}\textsuperscript{+}. Even more interestingly, these authors detected phosphorylation of Ser-568 in DUR3 which showed a threefold increase after 15 min of NH\textsubscript{3}\textsuperscript{+} depletion and a decrease after 3 h. So far, the influence of this PTM on DUR3 functionality was not yet further investigated.

Overall, new perspectives about the roles of urea in plant nutrition have been proposed in the last years. However, a better understanding of the physiological and biochemical interplay among urea, NO\textsuperscript{−} and NH\textsubscript{3}\textsuperscript{+} is still required. In addition, some environmental concerns exist about possible urea runoff from the soil following the massive use of urea fertilizers, especially if in combination with urease inhibitors [101]. This aspect could be further exacerbated considering that urease inhibitors can negatively affect urea metabolism in plants [110]. Therefore, future studies are needed to reach a fully aware use of urea in agriculture.

3. Influences of Nitrogen Forms on Root System Architecture

The root system architecture (RSA) is defined as the three-dimensional spatial configuration of root components within the soil. The primary root (PR) length defines the depth of the vertical soil layers reached by the plant, while lateral roots (LR) enhance horizontal soil exploration. In addition, root hairs greatly expand the absorptive surface area [111,112]. The arrangement of these components of the root system is determined by the interplay between developmental genetic programs and (a)biotic environmental cues [112,113]. RSA shows high plasticity according to both plant endogenous nutritional status and to the kind of N form available in growing media (Figure 4).

Root architectural responses to NO\textsuperscript{−} supply are strictly dependent on the availability and spatial/temporal distribution of the anion within the soil. Regarding homogenous NO\textsuperscript{−} provision, total root length increases under moderate NO\textsuperscript{−} supply, but decreases under extreme conditions, such as an excess or a severe deficiency. This reflects a systemic response that probably depends on plant nutritional status and hormonal balance [14,114]. Conversely, localized NO\textsuperscript{−} availability directly stimulates the LR elongation and emergence into the NO\textsuperscript{−}-enriched-patch, while LR growth is inhibited outside that zone. These combined responses represent an adaptive strategy to cope with the spatial heterogeneity of NO\textsuperscript{−} availability in soil (Figure 4) [32].

Needless to say, RSA changes require a wide reprogramming of gene expression in roots, which in Arabidopsis NRT1.1 plays a double role (Figure 1). At a null or low NO\textsuperscript{−} concentration, NRT1.1 is phosphorylated and recruited into functional microdomains at the PM, where it facilitates auxin transport out of the LR primordia, hence preventing their outgrowth. At high NO\textsuperscript{−} concentrations, non-phosphorylated NRT1.1 goes toward
oligomerization that, inducing its endocytosis and secretion, triggers the suppression of auxin transport as well as the stimulation of signaling pathways for LR development, partly overlapped by the NPR [16,49,115,116]. In Arabidopsis, it was also proposed that NRT2.1 could participate in the suppression of LR initiation induced by high carbon (C)/N ratios (Figure 1) [114]. However, the network of systemic and local signals behind this response is not yet characterized and several aspects await elucidation [18].

Figure 4. Scheme of the main effects of the different N forms on the responses affecting root system architecture (RSA) that are also related to auxin (aux) signal. Full lines indicate positive effects, dashed lines indicate negative effects. Details are explained in the text.

Similarly, RSA is differently affected by a homogeneous supply or localized provisions of NH₄⁺. When NH₄⁺ is provided as a sole N form, especially under elevated supplies, typical responses consist in the alteration of root gravitropism and in the inhibition of elongation of PRs and LRs (Figure 4). The first is mainly associated with the disturbance of auxin distribution in the root apex, while the second one was related to apoplastic and intracellular pH changes, increase in reactive oxygen species (ROS) formation, and alterations in protein glycosylation [117]. The involvement of the auxin signal is also suggested by the observation that NRT1.1 takes part in NH₄⁺ toxicity in the absence of NO₃⁻ [118]. Conversely, localized NH₄⁺ provision to N starved plants significantly promotes LR branching (Figure 4). It was recently demonstrated that this response is related to the interplay between apoplastic pH and auxin signal, especially as far as it regards higher-order LR branching. In detail, the influx of NH₄⁺ facilitated by AMTs activates the PM H⁺-ATPase (Section 4.2) increasing the acidification of the root apoplast. This pH shift prompts protonated auxin to permeate across the PM into cortical and epidermal cells, finally stimulating the emergence of LR primordia [119]. Moreover, it was proposed that a signaling event mediated by AMT1;3 is required for these changes in RSA (Figure 1) [120]. Whether AMT1;3 acts as an actual transceptor or is a part of a more complex regulatory network is still to be elucidated [117]. Curiously, the observation that under excess NH₄⁺ AMT1;3 amass into clusters internalized by clathrin-mediated endocytosis [121], resembles somehow the mechanism proposed for the NRT1.1 signaling functionality. Finally, it is possible to speculate that the specific and differential phosphorylation events through which AMT1;3 goes in response to NO₃⁻ (Section 2.2) could be somehow involved in mediating the synergistic effects on RSA prompted by the co-provision of NO₃⁻ and NH₄⁺, consisting of induction of LR elongation for NO₃⁻ and of LR formation for NH₄⁺ (Figure 4).

Although root growth is inhibited by several amino acids, for most of them the effect was associated with indirect metabolic responses [12]. However, direct effects on RSA were proposed, among which that induced by Glu in Arabidopsis is the best characterized (Figure 4). Glu provision at low concentration (<50 μM) exerts an inhibitory effect on PR
growth. It was shown that this response, especially in the early phase, is dependent on the perception of Glu by the PR tip, and is also combined with minor elongation of pre-formed LRs and with stimulation of outgrowth of new LRs behind the PR apex, finally resulting in a shorter and more branched root system [122]. This response involves auxin and kinase-mediated signals and was interpreted as an adaptive strategy to enhance plant competitiveness against microorganisms in amino acid uptake into enriched soil zones [122,123]. Interestingly, the effect of Glu on Arabidopsis root growth is strongly inhibited by NO\textsuperscript{+}, but not by other forms of N. The authors suggested that the NO\textsuperscript{+} ion itself acts by an NRT1.1-mediated signal at the PR tip to repress the root architectural changes induced by Glu when inorganic N forms are abundant, conditions where a short and highly branched root system could be limiting [124]. A putative signaling role of amino acids in RSA and plant development was supported by the discovery of a family of Glutamate-like Receptors (GLRs) in plants. GLRs act as tetrameric amino-acid-gated Ca\textsuperscript{2+} channels involved in different aspects of plant physiology, including the regulation of C and N metabolism and abscisic acid signaling. However, their roles in RSA adaptation to amino acid provision still await for elucidation [12]. Moreover, other amino acids seem to act in different ways, as for instance, the auxin effect attributed to tryptophan [122]. Finally, it is important to note that crop responses are species-specific and in soil the root system is exposed to a mix of different amino acids, which probably exert combined effects [10,125].

Overall, several proteomic studies provide evidence about the involvement of hormone response to different N nutrition. For instance, the analysis of the responses induced by NO\textsuperscript{+} in maize roots suggested a down-regulation of the clathrin-mediated endocytosis that regulates the auxin signal and transport [126], as well as changes in abundance of enzymes involved in redox balance and nitric oxide (NO) signaling, were highlighted [126,127]. Moreover, a recent work-related the increase in elongation of roots occurring in rapeseed during N deficiency with a change in the abundance of an auxin-responsive protein and the reduction of peroxidase levels and activity. This RSA rearrangement was also associated with an enhanced abundance of several enzymes involved in cell wall organization and biogenesis, highlighting the importance of this metabolism in root growth [128]. Clearly, although these studies represent a good starting point, further work is needed to reach a complete description of this multifaceted process.

4. Other Activities in the Plant Cell Plasma Membrane Related to Nitrogen Uptake

4.1. Relations among Nitrogen Uptake and Aquaporins

In plants, aquaporins (AQPs) are protein channels, mediating the transport of water and a small set of solutes and gasses across cell membranes, with key roles in the physiological adaptations to abiotic and biotic conditions, including the availability of mineral nutrients [129]. AQPs belong to the superfamily of Major Intrinsic Proteins (MIPs) and in higher plants comprise five subfamilies, consisting of the Plasma membrane Intrinsic Proteins (PIPs), the Tonoplast Intrinsic Proteins (TIPs), the Nodulin26-like Intrinsic Proteins (NIP), the Small basic Intrinsic Proteins (SIPs), and the uncategorized (X) Intrinsic Proteins (XIPs) [130]. AQPs are small proteins containing six putative transmembrane domains and fold into the membrane with the C- and N-termini inside the cytosol. Specific motifs of the protein chain, some of which diverge among the five subfamilies, play a part in determining the substrate selectivity, as well as the internal loops being engaged in the AQP gating, a composite mechanism controlled by cytosolic pH, divalent cations, and phosphorylation. AQPs assemble into tetramers, in which each subunit forms a pore, plus a putative fifth pore formed in the center of the complex. Co-expression studies in *Xenopus* oocytes suggested the formation of heterotetramers whose composition could affect the targeting, pH sensitivity, and activity of PIPs [129]. For instance, in maize, it was demonstrated that interactions between members of the two subgroups PIP1 and PIP2 participate in targeting and stabilization of the water channel into the PM [131,132]. Interestingly,
proteomic studies revealed different PTMs in PIP chains, such as multiple sites of phosphorylation detected in maize, *Brassica oleracea*, and Arabidopsis [133–135], as well as methylation events that in the last species were related to the trafficking of the protein [136].

The main AQPs in the root cell PM are PIPs, NIPs, and XIPs, among which PIPs show the highest selectivity and efficiency for water transport and play a predominant role in determining the root hydraulic conductivity (*Lp*) [129]. A significant increase in this parameter was observed in response to the renewed availability of NO\(^{-}\), which probably is involved in sustaining the recovery of plant biomass accumulation and the nutrient uptake [129,137]. The studies aimed at associating this response to an up-regulation of the transcription of PIP genes in crops provided divergent results, probably due to different treatment timings (short- or long-period responses) and plant species [138,139]. Some explanations could be found in an excellent proteomic study that highlighted that during NO\(^{-}\) deprivation the *Lp* decrease scarcely correlates with the abundance of PIPs, especially for PIP1. On the contrary, the extent of phosphorylation seems to have a predominant role [140].

Since transpiration significantly enhances *Lp*, by increasing the PIP expression in roots [129], these responses may be a secondary effect induced by the recovery of leaf metabolism, as well as the osmotic effect of NO\(^{-}\) cannot be excluded. However, a recent study conducted on Arabidopsis mutants showed that the *Lp* correlates with the shoot NO\(^{-}\) contents, and also revealed a side effect of the NRT2.1 gene, uncoupled from its transport function, in the transcriptional and translational regulation of PIPs [141]. Considering the proposed signaling role for NRT2.1 during LR development [114], this finding opens the intriguing hypothesis of the interplay between NO\(^{-}\) signaling, PIP expression, and RSA (Figure 1).

It was recently proposed that in plants AQPs could be also involved in determining the toxicity symptoms induced by excess NH\(_4^{+}\) provision (Section 2.2) [57,59]. Starting from the hypothesis that in this condition an influx/exflux transport cycle of NH\(_3\)/NH\(_4^{+}\) across the PM of root cells occurs, it was recently demonstrated that the predominant form recruited in this futile cycle is the NH\(_3\) species. Moreover, the observed kinetic properties and responses to chemical treatments, such as the inhibitory effects of mercuric cation (Hg\(^{2+}\)) and intracellular acidosis on NH\(_3\) cycling (known blockers of AQP activity [130]), were consistent with the implication of AQPs. According to the proposed model, AQPs passively mediate both NH\(_3\) influx and efflux across cell membranes, without energy dissipation, but allow NH\(_3\) to span into cell compartments on the basis of the concentration gradients and pH conditions, that result in the acidic trapping of NH\(_4^{+}\) into the vacuole [59]. The hyper-accumulation of NH\(_4^{+}\) into the vacuole could account for the decrease concentration of cations, especially K\(^{+}\), Mg\(^{2+}\), and Ca\(^{2+}\), which is considered one of the major causes for the “ammonical syndrome” in plants [9,142]. Interestingly, this model could in part explain why the co-provision of NO\(^{-}\) and NH\(_4^{+}\), balancing the cationic/anionic ratio into the vacuole, is the best nutritional condition [9]. Moreover, it is well known that NH\(_4^{+}\) toxicity is rescued in plants under high K\(^{+}\) availability [142]. Although this is ascribable to a direct nutritional effect of K\(^{+}\) and competition against NH\(_4^{+}\) transport (Section 2.2), since K\(^{+}\) availability is associated with a decrease in *Lp* [129], it is intriguing to propose that a different AQP functionality could contribute to this effect (Figure 1). However, molecular aspects of the model require detailed elucidation. On the one hand it is now described that TIPs facilitate NH\(_3\) transport [143], on the other hand, to our knowledge, evidence of a significant contribution of PIPs and NIPs in the NH\(_3\) transport across the PM is lacking. Phosphoproteomics highlighted transient changes in the phosphorylation state of some PIP isoforms in Arabidopsis plants subjected to NH\(_4^{+}\) resupply or deprivation [52,67]. Moreover, we detected that the abundance of two PIP2 isoforms in maize roots differently changed in response to NO\(^{-}\) or NH\(_4^{+}\) supply. Interestingly, in co-provision, NO\(^{-}\) seemed to have a prevalent effect, opening new perspectives about the interplay between inorganic N forms and water homeostasis in roots [71].
Phylogenetic and structural analyses sorted the NIP subfamily into three main subgroups. Group I comprises members showing transport selectivity mainly for water and glycerol. Instead, NIP subgroup II shows high permeability to glycerol and large solutes, such as urea, but not to water, and, finally, group III is permeable to water and urea, but not glycerol [144,145]. Among the last two groups, NIP1 in zucchini (Cucurbita pepo L.) was the first one characterized as able to complement Dur3p deficiency (Section 2.4) in yeast [146]. In Arabidopsis roots, one of the most abundant members is NIP5;1, which is involved in boron (B) uptake, but, under B deficiency, it also participates in the high and low-affinity uptake of urea (Figure 1) [144,147]. This role for NIP was confirmed in different crops. For instance, in maize NIP2;1 and NIP2;4, both expressed in roots, were characterized as able to mediate urea transport [148]. A similar function was recently proposed for NIP2;1 in cucumber (Cucumis sativus L.), for which ectopic expression in Arabidopsis thaliana improved the growth of the wild-type genotype and rescued the growth of an Atdur3 mutant, when urea was provided as the sole N source [145]. Overall, these results sustain the hypothesis that among the AQP family, some NIPs participate in urea uptake, even if other studies are needed to decipher the actual meanings for crop N nutrition and plant physiology.

4.2. The Involvement of Plasma Membrane H+-ATPase in Nitrogen Uptake

In plants, the plasma membrane H+-ATPase is a pump that exports cytosolic protons outside the PM, coupling the transport with ATP hydrolysis. Its activity generates the transmembrane electrochemical gradient that represents the driving force for solute import into plant cells, and hence it is intrinsically connected with nutrient uptake by roots. PM H+-ATPase is a large protein (of about 100 kDa) with 10 transmembrane domains and both N- and C-terminus protruding into the cytosolic face of the PM. The C-terminus acts as an auto-inhibitory regulatory domain, the phosphorylation of which at the penultimate Thr (Thr-955 in the model AHA2 of Arabidopsis) and the binding of 14-3-3 proteins lead to the activation of the enzyme [149]. In addition, multiple phosphorylation sites in the PM H+-ATPase were identified in plants, each with a different regulatory role as recently reviewed by Falhof and co-workers [150]. Considering that the uptake of NO₃⁻, amino acids, and urea are all mediated by active solute/H⁺ symporters, the relevance of the PM H+-ATPase in the plant N nutrition is clear. Also, NH₄⁺ uptake is associated with the stimulation of the PM H+-ATPase activity, triggered by the transient depolarization of PM electrical potential induced by the influx of the cation into the cytosol (Figure 1) [151]. Furthermore, the acidification of apoplast induced by NH₄⁺ supply is involved in the typical responses in the RSA (Section 3).

Higher plants have multiple PM H+-ATPase isoforms, with tissue specificity and co-presence in a specific cell type. Phylogeny analysis of Arabidopsis PM H+-ATPase isoforms, named AHA (Autoinhibited H+-ATPase) recognizes five subfamilies. One group contains AHA4 and AHA11, the second one clusters isoforms similar to AHA1 (AHA2, AHA3, AHA5, AHA12), the third and the fourth only consist of AHA10 and AHA7, respectively, and the last group comprises AHA6, AHA8, and AHA9 [152,153]. Unexpectedly, AHA10-like isoforms target the tonoplast where they possibly assemble as heterooligomers [154,155]. Although this organization seems to be overall conserved across species, some plants were not recognized members for all the five subfamilies [27,153].

In Arabidopsis, AHA1, AHA2, and AHA7 are the predominant isoforms accumulated in root epidermal cells. Interestingly, AHA7 shows the peculiarity to be auto-inhibited by acidic extracellular pH, and, while AHA2 is fundamental in root cell expansion, AHA2 and AHA7 are required for limiting root hair length, a process in which they play different roles [153]. In the past years, phosphoproteomics highlighted multiple phosphorylated sites in the AHA1 and AHA2 isoforms in NH₄⁺-adapted Arabidopsis plants. More recently, comparative proteomic analyses revealed a major accumulation of AHA2 under nitric than ammonial nutrition [156], overall confirming its involvement in the metabolic adaptations to the N external availability [52,67].
In this case, it is possible to draw reliable similitudes between Arabidopsis and some crops. For instance, a time-resolved analysis of the PM transport systems during PNR proposed that the PM H⁺-ATPase isoforms mainly involved are those encoded by ZmHA2 and ZmHA4 genes, both belonging to the second phylogenetic group [27]. Considering that a study of the root hair proteome in this crop revealed a reduction of AHA2 and AHA11 levels during N deprivation [157], it is possible to speculate that different isoforms play different roles in specific cell types.

To conclude, it is noteworthy that the PM H⁺-ATPase activity seems to be influenced not only at transcriptional levels and by phosphorylation but also by the oligomeric state of the enzyme. In tobacco cells, BN-PAGE (Blue Native, a non-denaturing PAGE) and electron microscopy analyses showed that in the microsomal fraction the enzyme exists in dimeric form, which assembles in hexameric complexes when phosphorylated and bound to 14-3-3 proteins [158]. Later, a model for the formation of the complex was proposed. Accordingly, a PM H⁺-ATPase dimer, inactive probably due to intermolecular contacts of the C-termini, after phosphorylation becomes a target for the binding of a 14-3-3 dimer, that allows sequential interactions with two other PM H⁺-ATPase/14-3-3 dimers, finally resulting in the assembly of active hexameric complexes (Figure 5) [159]. Interestingly, it was recently shown that in maize roots the PNR is associated with an increase in the abundance of monomeric, dimeric, and hexameric states of the PM H⁺-ATPase [27]. Overall, several questions are still open, such as those regarding the functional roles of different oligomeric states and the putative involvement of different isoforms, that could be effectively addressed in the future, also thanks to the recent technical improvements in plant proteomics (Section 5).

![Figure 5](image_url)

**Figure 5.** Schematic representation of (A) H⁺-ATPase/14-3-3 hexamer activation by an unknown kinase mediated-phosphorylation (“P”, yellow colored, question mark) and binding of 14-3-3 proteins. (B) Cytosolic-side view of the active hexameric complex. Model adapted from [159]. Further details are explained in the text.

5. **Nitrogen Nutrition in Plants and Root Proteomics: Goals and Pitfalls**

In the last decades, proteomics was applied to study various aspects of N nutrition in plants, analyzing different kinds of samples, from whole seedlings to specific cell types, and combining different analytical strategies. We selected some studies devoted to investigating specifically the proteomic profiles of roots, conducted in Arabidopsis and crops (Table 2).
Table 2. Proteomic studies on plant responses to different N nutrient availabilities in roots. PepF: Peptide Fractionation. NO₃⁻/NH₄⁺ indicates co-provision. The other acronyms are detailed in the text. Plant species are reported in sub-titles.

| Proteome Fraction | Analytical Methods | N Nutrient | Major Results | Ref. |
|-------------------|--------------------|------------|---------------|------|
| **Arabidopsis thaliana** | | | | |
| Total | iTRAQ / PepF + LC-MS/MS | NO₃⁻, NH₄⁺ | NO₃⁻ or NH₄⁺ availability differently affects C, N, and secondary metabolism, peroxidases, and AHA2. | [156] |
| Microsomal | PepF / TiO₂ + LC-MS/MS | NO₃⁻ | NO₃⁻ deprivation induces changes in abundance and PTMs of PIPs. | [140] |
| Microsomal | TiO₂ + LC-MS/MS | NO₃⁻, NH₄⁺ | N deprivation induces transient changes in the proteome and in phosphorylation of NRT2.1, AMT1;1, AMT1;3, DUR3, PIP2.2, PIP3, AHA1, AHA2. | [67] |
| PM | BN-PAGE + GeLC-MS/MS | NO₃⁻, NO₃⁻/NH₄⁺ | Regulative role of Ser-501 phosphorylation for the activity of NRT2.1. | [54] |
| **Brassica spp.** | | | | |
| Total | TMT / PepF + LC-MS/MS | N | N deficiency induces changes in the abundance of proteins involved in auxin and cell wall metabolism. | [128] |
| Total | 2-DE + MALDI-MS/MS | NO₃⁻, glycine | Glycine availability induces changes in the abundance of proteins involved in N metabolism and in defense. | [99] |
| **Zea mays** | | | | |
| Total (root hair) | LC-MS/MS | NO₃⁻/NH₄⁺ | N deprivation affects N, amino acid, and C metabolism and the abundance of peroxidases and AHA2 and AHA11. | [157] |
| Total | 2-DE + MALDI-MS/MS | NO₃⁻ | NUE is associated with different responses of C and N metabolism to NO₃⁻ availability. | [160] |
| Total (root transition zone) | iTRAQ + GeLC-MS/MS | NO₃⁻ | Relations of NO₃⁻ supply with the abundance of peroxidases and of proteins involved in hormone balances. | [126] |
| Total | GeLC-MS/MS | NO₃⁻, NH₄⁺, NO₃⁻/NH₄⁺ | NO₃⁻ and NH₄⁺ availability differently affects N and C metabolism, protein synthesis, K⁺ channel, peroxidases, PIP2 isomers. | [71] |
| Soluble | 2-DE + LC-MS/MS | NO₃⁻ | NO₃⁻ supply affects enzyme involved in N and C metabolism and in redox balance. | [127] |
| Soluble | 2-DE + LC-MS/MS | NO₃⁻, NH₄⁺, NO₃⁻/NH₄⁺ | NO₃⁻ and NH₄⁺ availability differently affects abundance and phosphorylation of GS (iso)forms. | [161] |
| **Hordeum vulgare** | | | | |
| Soluble | 2-DE + MALDI MS/MS | NO₃⁻, NH₄⁺ | NO₃⁻, NH₄⁺ and N deficiency differently affects C, N metabolism, peroxidases, and redox balance. | [162] |
| **Lolium perenne** | | | | |
| Soluble | 2-DE + MALDI MS | NO₃⁻/NH₄⁺, Gly | Glycine availability induces changes in abundances of enzymes involved in C, Met, and adenosine metabolism. | [98] |
| **Lycopersicon esculentum** | | | | |
| Total | TMT/PepF + LC-MS/MS | NO₃⁻, NH₄⁺ | NO₃⁻ and NH₄⁺ availability differently affects the abundance of enzymes involved in C and N metabolism and peroxidases. | [163] |
| **Solanum tuberosum** | | | | |
This selection includes studies that adopted different experimental plant growth conditions, such as availability and starvation of a specific N form, co-provision, re-supply, and depletion treatments. At a first look, it is evident that most of them deal with the plant responses to inorganic N forms, while those induced by organic N nutrients were much less frequently studied (Table 2). This trend mirrors the overall literature and indicates that the relations among plant nutrition and organic N forms represent a research field that could deserve attention in the next few years.

Despite individual topics, the studies in Table 2 are congruent in highlighting similar traits of root metabolism being affected by N availability. Some common traits include the reciprocal interplay between N and C metabolism, including protein synthesis and folding, in the relations between NH₄⁺ assimilation and mitochondrial activities as well as the effects of N availability on cell redox homeostasis and peroxidase abundance in roots.

Among those reported in Table 2, each proteomic methodological approach has its specific advantages and pitfalls.

The first steps of sample preparation have a very high relevance in determining the features of the proteome subset under investigation. In the past years, most of the proteomic studies on N nutrition in plants analyzed total or soluble proteomes, especially in crops (Table 2). This strategy is particularly suitable to highlight the interplay among different metabolic pathways and allows to obtain a widespread holistic overview of the proteome. Hence, it is advantageous when the topic of interest is addressed by proteomics in a given plant species for the first time, as often occurs in crops. However, the analysis of entire proteomes at once greatly limits the detection of the less abundant proteins that frequently play key roles in the plant responses to nutritional stimuli, such as transcription factors, kinases and phosphatases, receptors and, also, transporters [164].

On the other hand, fractionation techniques allowing the purification of microsomal or plasma membrane enriched samples, significantly help in overcoming this drawback. To our knowledge, until now this strategy was adopted to study root responses to N nutrition only in Arabidopsis [52, 67, 140]. Coupled with phosphopeptide enrichment and gel-free protein quantification, it greatly contributed to the study of the roles of PTMs in the functionality of several transporters involved in N uptake, as extensively reported above.

Similarly, the selection of the best analytical techniques to resolve and quantify a plant proteomics sample mainly depends on the aims of the research. Gel-based proteomics is very suitable to individually analyze allelic variants and isoforms [165]. For instance, a 2-DE based approach pointed out different effects of NO₃⁻ and/or NH₄⁺ availability on the abundance and PTMs of the Glutamine Synthetase (GS) (iso)forms in maize roots [161]. However, conventional 2-DE protocols are not very suitable to resolve hydrophobic proteins and hence are poorly applicable in membrane proteomics [165]. Unfortunately, although a gel-free targeted proteomic methodology was defined to analyze the abundance of members of the NRT1, NRT2, AMT1, AHA, and PIP subfamilies in Arabidopsis [166], its application to other plant species still seems to be very laborious.

At the same time, the development of LC-MS/MS-based analytical techniques to quantify the proteins in complex samples, such as isobaric/isotopic labeling or the label-free shotgun approach, has greatly increased the number of proteins quantifiable in a single experiment. However, plant samples generally contain a high amount of compounds, such as pigments, lipids, polysaccharides, and secondary metabolites, which interfere with LC-MS/MS. Therefore, LC-MS/MS often gives the best results when combined with a prior purification of the proteome by SDS-PAGE [165]. This very powerful approach, called GeLC-MS, was recently applied to analyze the time course, within the first 54 h of induction, of the responses in roots and leaves of maize plants exposed to NO₃⁻ and/or
NH₄⁺, allowing the simultaneous quantification of hundreds of proteins [71]. Another aspect that deserves attention is that overall shotgun approaches give information also about the components of the proteome not affected by the experimental treatments, that even if often overlooked, could be relevant from a biochemical point of view.

6. Conclusions and Future Trends

In the recent decades, proteomics gave its contribution in discerning the metabolic network and interactions on which N nutrition in plants relies on. However, it is evident that future work is needed to unravel this complexity in crops, and also in relation to organic N nutrition. Recently, deep proteomic profiling was applied with excellent results in several crops, including maize [167], wheat [168], and tomato [169], opening the way for their application in the study of plant responses to environmental stimuli, such as N nutrition. Similarly, considering the relevance of protein complexes in the functionality of the PM transporters (often highlighted in this review), the newly available approaches to analyze protein complexes [170] could be very effective in the future to achieve a better discerning of the role played by protein–protein interactions.

Author Contributions: Conceptualization, C.M., L.E., and B.P.; visualization, C.M.; writing—original draft preparation, C.M. and B.P.; writing—review and editing, C.M., L.E., and B.P. All authors have read and agreed to the published version of the manuscript.

Funding: This research received no external funding.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Conflicts of Interest: The authors declare no conflict of interest.

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