Arginine and Nitrogen Storage

José L. Llacer1, Ignacio Fita2 and Vicente Rubio1

Addresses

1 Instituto de Biomedicina de Valencia–Consejo Superior de Investigaciones Científicas (IBV-CSIC) and Centro de Investigación Biomédica en Red de Enfermedades Raras (CIBERER-ISCIII), Jaime Roig 11, Valencia 46010, Spain.

2 Instituto de Biología Molecular de Barcelona–Consejo Superior de Investigaciones Científicas (IBMB-CSIC) and Institute for Research in Biomedicine, Josep Samitier 1-5, Parc Cientific, Barcelona 08028, Spain.

Corresponding authors:

Fita, Ignacio (ifrcri@ibmb.csic.es)
Rubio, Vicente (rubio@ibv.csic.es)

Summary

When nitrogen is abundant, prokaryotic and eukaryotic oxygen-producing photosynthetic organisms store nitrogen as arginine, by relieving feedback inhibition of the arginine biosynthesis controlling enzyme, N-acetylglutamate kinase (NAGK). The signalling protein PII, an ancient and widely distributed nitrogen/carbon/ADP/ATP sensor, mediates feed-back inhibition relief of NAGK by binding to this enzyme. PII phosphorylation or PII binding of ADP or 2-oxoglutarate prevent PII-NAGK complex formation. Crystal structures of NAGK and of cyanobacterial and plant PII and of the corresponding PII-NAGK complexes have been recently determined. In these complexes, two polar PII trimers sandwich one ring-like NAGK hexamer. Each PII subunit contacts one NAGK subunit, triggering a symmetry-restricted narrowing of the
NAGK ring, with concomitant adoption by the arginine sites of a low-affinity conformation.

**Introduction**

Nitrogen assimilation is a requisite for growth. The nitrogen can be supplied as ammonia, nitrate, dinitrogen or amino acids [1,2]. Ammonia incorporation is cheap in terms of energy and reducing power, being generally effected (Figure 1a) by the combined action of glutamine synthetase (GS) and glutamate synthase (GOGAT), using 2-oxoglutarate (2OG) to generate glutamate [1-3], that then provides the nitrogen for making other amino acids [4]. Pioneering studies of GS regulation [5] revealed a key controlling protein that was called PII because it was peak II in a gel filtration column [6]. PII, a very ancient signalling protein that is widely distributed throughout all domains of life but that is absent from animals [1,2,7,8], senses and integrates nitrogen and carbon abundance and energy status information, by binding 2OG and ATP/ADP and, in many organisms, by being either phosphorylated, uridylylated or adenylylated [9-11].

PII interacts with and influences many protein targets, including enzymes, channels and regulatory proteins [1,2,7,8,12,13]. Crystal structures of PII from various sources were obtained [14-19] but little was known about how PII exerts its effects. Very recently, crystal structures of complexes of PII with the ammonium channel AmtB (Figure 1b) [20,21] or with the enzyme NAGK [22,23] have been determined. In the complex with AmtB, the large and flexible T-loop of PII occludes the cytoplasmic opening of AmtB (Figure 1c) by means of its tip residue Arg47 (Figure 1d), blocking channel operation. The present review will concentrate on the other complex, with the enzyme of arginine biosynthesis N-acetyl-L-glutamate kinase (NAGK). The structure of
this complex reveals a highly sophisticated activation mechanism that renders possible massive nitrogen storage as arginine. For recent reviews on other aspects of PII function, see [1,2,7,8,24-26]

Control of nitrogen storage as arginine by the signalling protein PII

NAGK was recently recognized in cyanobacteria and plants as a PII target [27-32]. These organisms store nitrogen as arginine (Figure 1a), which is incorporated into proteins such as seed proteins [33,34] or in arginine-rich copolymers [35]. Arginine is an ideal nitrogen store because it is nitrogen-rich, its incorporation into proteins minimizes its osmotic impact, and its nitrogen is easily mobilized [36]. Since arginine synthesis is feed-back inhibited at the NAGK level [34,37,38], this enzyme should be released from inhibition for build-up of large arginine stores. In oxygenic phototrophs, PII triggers this effect on NAGK by binding to the enzyme when nitrogen is abundant [27-32]. When nitrogen is scarce, 2OG accumulates because of reduced flow through GS/GOGAT and binds to PII in the presence of ATP, promoting PII-NAGK complex dissociation [29]. ADP binding to PII, and 2OG-promoted PII phosphorylation at Ser49 also prevent PII-NAGK complex formation in cyanobacteria [27-29].

Arginine-sensitive NAGKs are allosteric switches operated by arginine

The first NAGK crystal structure, of the arginine-insensitive *Escherichia coli* enzyme [39,40] revealed a homodimer having as backbone a 16-stranded β-sheet spanning both subunits (Figure 2a). Each 258-residue subunit presents the amino acid kinase fold (PFam PF00696; http://pfam.sanger.ac.uk): an open α3β8α4 sandwich composed of N- and C-domains that host, respectively, on the C-edge of the central,
mainly parallel, β-sheet, the sites for the substrates N-acetyl-L-glutamate (NAG) and ATP.

Arginine-sensitive NAGKs were known to be hexameric [29,41,42]. The recent determination of the first crystal structures of these enzymes, free [43] or, more recently, PII-complexed [22",23"], has shown them to be ring-like hexameric trimers of dimers (Figure 2b) having a central hole of 25-30Å-diameter, and presenting 32 (D3) molecular symmetry [22",23",43]. The ring is formed by linking three *E. coli* NAGK-like dimers through the interlacing of a N-terminal α-helix (the N-helix) that is not found in arginine-insensitive NAGKs. Arginine binds flat and extended near the interdimeric boundary (Figures 2b and 2e) in a cavity formed in the C-domain of each subunit between the α3 layer, the β sheet, and the C-terminal part of the N-helix after the kink (see below figures 4f and 4g). In agreement with the remoteness of this site relative to the substrate sites, arginine inhibits the reaction indirectly (Figure 2c), by increasing the separation between the C- and N-domains hosting the ATP and NAG sites [43]. The changes triggered by arginine inhibition were associated with changes in the tilt of the enzyme dimers within the hexamer (Figures 2d-e), rendering the hexameric ring wider and thinner (Figures 2c, 2d and 2e) [43].

The interlaced N-helices connect in the hexamer the arginine sites from adjacent dimers [43], and thus these helices and the hexameric organization are essential features for making NAGK an arginine-operated switch exhibiting sharp sigmoid arginine inhibition kinetics [29,41,44*]. Thus, at arginine concentrations below a certain threshold, the enzyme is not inhibited, allowing arginine synthesis. However, because of the sigmoid character of the inhibition, the enzyme is inhibited quite abruptly when this threshold arginine concentration is exceeded, stopping arginine synthesis (Figure 2f).
PII structure

PII proteins are homotrimers of 12-13-kDa subunits having a $\beta\alpha\beta\alpha\beta$ subunit topology, with $\alpha$ helices looking outward and the $\beta$ sheet inward and providing the intersubunit interactions (Figures 3a and 3b) [14-19]. The trimer body is hemispheric, from which two short loops (the B- and C-loops) and a large and highly flexible loop called the T-loop emerge from each subunit. The T-loop residues S49 and Y51 are, respectively, the phosphorylation and uridylylation sites in cyanobacteria and proteobacteria. When modified, the PII proteins cannot interact with their respective NAGK or AmtB targets [8]. ADP and MgATP bind to PII in a crevice between adjacent subunits in the convex surface of the PII body, at the base of the T-loop (Figures 3a and 3b) [15,19-21,23]. MgATP binding is associated with a compact conformation of the T-loop resembling a bent finger in which the Mg interactions play a crucial role (Figures 3b-3d), whereas ADP, bound without visible Mg ions at the same site, favours an extended T-loop conformation [19] (Figures 3a and 3c). Only in *Methanococcus jannaschii* PII one 2OG molecule has been found binding to one T-loop that also had MgATP bound [19] (Figure 3d), in agreement with the well known requirement of ATP for 2OG binding [8]. This T-loop was in the compact conformation [19], and thus the site for 2OG may be formed only when the T-loop adopts its compact conformation.

Effects of PII binding to NAGK

The effects of PII on NAGK have been characterized functionally best in the cyanobacteria *Synechococcus elongatus* strain PCC7942 and in the plant *Arabidopsis thaliana* [29,32], the same species in which the crystal structures of the complexes have been determined [22,23]. The NAGKs from both species present some functional
differences, and the effects of PII appear not to be identical, perhaps reflecting differences in the adaptive requirements in the cytoplasm of a free-living cyanobacterium and in plant chloroplasts, where PII and NAGK are found [30,45]. *S. elongatus* NAGK is virtually inactive in the absence of PII, having low $V_{\text{max}}$, high $K_m$ for NAG ($\sim$30 mM) [29] and requiring low arginine concentrations ($I_{0.5} \sim$30 µM) for inhibition [22”] (Figure 2f). *A. thaliana* NAGK is highly active, having, relative to *S. elongatus* NAGK, $\sim$3-fold higher $V_{\text{max}}$ and $\sim$4-fold lower $K_m$ for NAG, and requiring one order of magnitude higher arginine concentrations for inhibition [32]. In *S. elongatus*, PII activates NAGK up to $\sim$40-fold by increasing 2-4-fold $V_{\text{max}}$, by decreasing $\sim$10-fold the $K_m$ for NAG, and by augmenting $\sim$15-fold the arginine concentration required for inhibition [29] (Figure 2f). In contrast, in *A. thaliana*, PII activates NAGK up to 5-fold only, not affecting the $K_m$ for NAG, increasing marginally ($\sim$30%) $V_{\text{max}}$ and triggering a mere $\sim$3-fold increase in the concentrations of arginine needed for inhibition [32]. ADP and, under appropriate conditions, 2OG, dissociated the *S. elongatus* PII-NAGK complex [29], but these ligands were not found to dissociate in vitro the isolated complex from *A. thaliana* [32].

**Known structures of PII-NAGK complexes**

The reported crystal structures of the *S. elongatus* and *A. thaliana* complexes are highly similar [22”’], although only NAG was bound to the *S. elongatus* complex, whereas, in *A. thaliana*, NAG, ADP and arginine were bound to NAGK and MgATP was bound to PII. We will describe first the *S. elongatus* complex, and then we will highlight relevant differential traits of the *A. thaliana* complex. These complexes are the genuine ones formed in vivo: their stoichiometry is the same demonstrated for the *S. elongatus* complex in solution, the interacting residues include those conserved
exclusively in oxygenic phototrophs (the organisms in which this complex is formed),
and structure-directed mutagenesis of PII and NAGK triggered the expected effects in
vivo (yeast two-hybrid assays) and in functional assays of complex formation [22”].

The structure of the \textit{S.elongatus} PII-NAGK complex [22’’]

The complex (Figure 4a) consists of one NAGK hexamer sandwiched between
two PII trimers oriented with their convex surface pointing outwards. PII is not packed
tightly on NAGK, interacting with NAGK mainly through its T- and B-loops (Figures
4b and 4c). Each PII subunit interacts with one NAGK subunit only, contacting the $\alpha_4$
layer near its connection with the central $\beta$ sheet N-edge, far from both the arginine site
and the sites for the substrates (Figure 4d). The T-loop is in the compact form
resembling a bent finger (Figure 3b), explaining why ADP, by favouring an extended T-
loop conformation, disassembles the complex [29]. The T-loop tip is inserted in the
ridge between the NAGK N- and C-domains, interacting through several polar bonds
with the N-domain and with Gln258 from the C-domain (Figures 4d and 4e). An ion
pair network centred in PII reaches close to the NAG site of NAGK (Figure 4e),
favouring a high affinity form of this site, explaining the decreased $K_m$ for NAG. The
hydroxyl of Ser49 is centrally involved in the interactions with NAGK, explaining the
lack of complex formation upon Ser49 phosphorylation. If 2OG binds to PII in
\textit{S.elongatus} as in \textit{M.jannaschii} [19’’], 2OG-triggered complex disassembly must be due
to the negative charge contributed by 2OG, rather than to inappropriate T-loop structure
or to steric clashes.

The numerous T-loop contacts with the N-domain fix this domain and the
NAGK dimers in a position [22’’] that favours a high dimer tilt and, consequently, an
arginine-free-like orientation of the N-helix [43]. This conformation differs from the
arginine-inhibited NAGK form, characterized by low-tilt dimers, a widened ring, and a characteristic N-helix orientation (Figure 2c) [43]. Thus, PII binding favours the active, low affinity form (for arginine) of the NAGK hexamer. These global NAGK ring changes fit the principle of preservation of symmetry in domain movements [46]. This principle predicts for NAGK that, in the twofold axis-related subunits of different dimers, the cycle of opening and closing of the active centres that is consubstantial with enzyme catalysis should be facilitated by coordinating the movements of the two subunits to preserve the twofold symmetry. This necessarily requires that the tilt and the distance of the dimers from the threefold axis is changed, as observed, because of the restrictions imposed by the threefold symmetry of the D3 point group.

Interactions not mediated by the T-loop, involving the B-loop and the $\beta1-\alpha1$ junction (particularly Phe11) close to the flat face of the PII body, target the C-domain of NAGK, possibly accounting for the reported [22”] C-domain displacement of ~2 Å towards PII that appears to contribute, together with the inappropriate (for arginine binding) N-helix orientation, to the observed arginine site enlargement and the low affinity for arginine (compare figures Figures 4f and 4h).

The *A.thaliana* PII-NAGK complex [23”]

Although MgATP was not added and 10 mM ADP was present in the crystallization mixture [23”], the complex contains MgATP bound to PII similarly as in *M.jannaschii* PII [19”], reflecting the previously demonstrated [47] high affinity of *A.thaliana* PII for MgATP (Figure 3b). Thus, the reported lack of complex dissociation by ADP [32] may have reflected the lack of displacement of PII-bound MgATP. The lack of effect of PII binding on the $K_m$ for NAG of *A.thaliana* NAGK [32] is explained by the exclusion of the NAG site from the ion-pair network formed upon T-loop
binding, because of the substitution by Asn of the corresponding Asp found in *S. elongatus* NAGK [22”]. Trp22 (corresponding to Phe11 of *S. elongatus* PII) was the only non-T-loop residue reported [23”] to interact with NAGK, but, actually, Lys240 interacts in one subunit of *A. thaliana* NAGK (see PDB file 2rd5) with Glu97 of PII, similarly to Arg233 of NAGK with Glu85 in the complex of *S. elongatus*. In the *A. thaliana* complex all the NAGK active centres contain bound NAG and ADP (Figure 4d), and they exhibit in three subunits a closed conformation that possibly may be equated with a catalytically active form. The movement causing this closing of the active centre is a 11º-rotation around a hinge located at the interdomain boundary in the NAGK subunit, near the site at which the T-loop contacts NAGK [23”]. Thus, the stimulatory effect of PII on the kinetics of arginine-inhibited NAGK has been attributed [23”] to PII-limitation of the degree of opening and closing of the active site cleft, in opposition to the domain-separating inhibitory effect exerted by arginine. Arginine was present at high concentration (50 mM) in the crystallization mixture and was also found binding in all the subunits, but, as expected from the overall changes triggered by the binding of PII, the N-helix is inappropriately oriented for high affinity binding of arginine, yielding a low-affinity conformation of the arginine site (Figure 4g).

**Conclusions and prospects**

Despite the high similarities of the corresponding complexes, there are substantial differences in the sensitivity to PII of cyanobacterial and *A. thaliana* NAGKs. Perhaps in other photosynthetic eukaryotes expected to experience more changes in nitrogen supply than higher plants, such as unicellular algae or even multicellular algae and lower plants, the effects of PII on NAGK might be closer to those in cyanobacteria. Therefore, the PII-NAGK system should be investigated in lower photosynthetic
eukaryotes. More work is also needed on *A. thaliana* to clarify whether 2OG regulates the PII-NAGK complex, since 2OG is a low-nitrogen abundance signal in plants [48] and was found to bind to *A. thaliana* PII [47]. It is therefore surprising the lack of effect of 2OG on the PII-NAGK complex, as reported [32]. Another question deserving further work is the possibility that nitrogen accumulation as arginine could take place in other taxonomic groups, given the very widespread co-occurrence in the same organism of PII [1] and hexameric arginine-sensitive NAGK [34,37,38,41]. Although this possibility is not favoured by the exclusive conservation in photosynthetic organisms of a subset of the residues found to interact in PII-NAGK complexes [22”,23”], the possibility of the existence of different complementary sets of interacting residues in other taxonomic groups cannot be excluded. It is also important to determine the structure of novel PII complexes with other targets [49], to clarify whether the T-loop conformations observed in the PII-AmtB and PII-NAGK complexes are the only two standard conformations used in the interactions of PII or whether the T-loop conformation is tailored specifically to each target, resulting in a very diverse array of modes of regulating the functionality of the different targets.

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References and recommended reading

Papers of particular interest, published within the annual period of review, have been highlighted as:

• of special interest
•• of outstanding interest


   This review of nitrogen regulation in prokaryotes gives a good updated picture of the elements known to be involved in this regulation. The numerous illustrations provided of reactions and regulation mechanisms give clear schematics that are very helpful for understanding the regulation.


   Very clear and updated review of our present-day knowledge of PII protein functions. It integrates the existing structural knowledge at the moment of writing. It has very clear schemes to illustrate the present understanding of three different modes of negative regulation exerted by PII. It also schematizes in a very clear way the PII-NAGK system.


This is the first crystal structure of a plant PII protein. The structure, at 1.9 Å resolution, presents the characteristic PII fold and trimeric architecture and has the T-loop disordered, but it reveals some plant-specific specializations, including extensions of unknown signification at both ends of the PII polypeptide.


This study shows that the PII protein of *M. jannaschii* GlnK1 forms a complex with the channel AmtB, revealing the influence of effectors, particularly 2OG, on complex formation. It also visualizes AmtB and the GlnK1-AmtB complex by electron microscopy. Finally, it reports crystal structures of GlnK1 with MgATP or with ADP,
and it provides the first snapshot of the effector 2OG binding to a PII protein. These studies demonstrate that MgATP and ADP are associated, respectively, with an extended conformation and a compact novel conformation of the T-loop. The finding of 2OG bound to the compact conformation of the T-loop having bound MgATP indicates that the site for 2OG is formed when MgATP is bound, explaining the cooperative binding of ATP and 2OG to PII. The dissociation of the GlnK1-AmtB complex triggered by 2OG is explained on the basis of the increase in the exposed negative charge of the T-loop.


This study and [21] are the first to report the structure of a PII-target protein complex at atomic resolution. Both studies reveal that GlnK (a PII protein of E. coli) interacts with AmtB almost exclusively via the T-loop, blocking the cytoplasmic opening of the channel by inserting into it the T-loop tip residue Arg47. ADP was found binding to PII in both studies, supporting the role of this nucleotide in promoting the extended T-loop conformation [19]. The structures explain also why the uridylylation of the T-loop residue Tyr51 prevents PII-AmtB complex formation.


See annotation for [20]

This study and [23••] report the first crystal structures of PII-NAGK complexes. The cyanobacterial proteins from *S. elongatus* PCC 7942 are used in this study reporting the structure of the complex at 2.75-Å resolution and containing NAG bound to NAGK. PII presents the compact conformation (see [19••]) although no MgATP is bound. It is concluded that the compact T-loop conformation is selected by NAGK. The central participation in the interactions between PII and NAG of the T-loop residue Ser49 explains complex dissociation by Ser49 phosphorylation. The complex-dissociating effect of 2OG is attributed to negative charge repulsion, and it is predicted to be dependent on the ionic composition of the medium. The decreased $K_m$ of NAGK for NAG within the complex is attributed to PII-triggered NAG site conformational changes. NAGK relief from arginine inhibition is due to changes in the arginine site resulting from symmetry-determined changes in the overall shape of the NAGK hexamer as well as to local effects.


See also annotation for [22••]. This study reports the crystal structure at 2.5 Å-resolution of the plant PII-NAGK complex. This complex, and the cyanobacterial complex reported in [22••] are highly similar. The crystals contain MgATP bound to PII and
NAG, ADP and arginine bound to NAGK. The site for the arginine inhibitor is in a low-affinity form because of the orientation of the N-terminal helices that interconnect the three dimers in the hexamer. Both NAGK subunits in the asymmetric unit exhibit different degrees of active centre opening. The movement causing this active centre change is a rotation around a hinge located at the boundary between both domains of each NAGK subunit, near the site at which the T-loop contacts NAGK. Thus, the stimulatory effect of PII on the kinetics of arginine-inhibited NAGK is attributed to a PII-limited degree of opening and closing of the active site cleft, in opposition to a domain-separating inhibitory effect exerted by arginine.


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demonstrate the extreme importance of the hexameric organization and of the cross-talk
between the arginine sites of different NAGK subunits mediated by the N-helix, for
rendering the inhibition sigmoidal and appropriate in terms of sensitivity to the
physiological range of arginine concentrations. The results highlight the importance of
the stabilization of high or low affinity forms of the arginine site in the determination of
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Figure legends

Figure 1. Roles of PII in ammonia assimilation. (a) Scheme of ammonia assimilation as arginine in oxygenic photosynthetic organisms. Ammonia, the glutamate product of the GS/GOGAT cycle, and the final product arginine are in large type and coloured blue. The red arrow and the red negative sign within a circle symbolize arginine inhibition of NAGK. PII is also shown in large letters and is coloured green. The green arrows and the green positive and negative signs within circles symbolize, respectively, NAGK activation by PII and release of NAGK from feed-back inhibition by arginine. (b) Complex of the *Escherichia coli* PII protein GlnK with its target AmtB, which is a trimeric ammonia channel (PDB file 2nuu). (c) Zoom of the T-loop interaction with AmtB. (d) Longitudinal section of the ammonia channel of one AmtB subunit, to illustrate the blocking of the channel by the side chain of the T-loop residue Arg47.

Figure 2. NAGK structures and inhibition by arginine. (a) Dimeric, arginine-insensitive *E. coli* NAGK (PDB file 1gs5) with AMPPNP (an inert ATP analog) and NAG bound, respectively, to the C-domain (in grey) and N-domain (golden) of each subunit. (b) Hexameric, arginine-sensitive NAGK of *Thermotoga maritima* with bound arginine (PDB file 2bty). Three subunits related by threefold symmetry are shown in grey, whereas the other three subunits are coloured yellow. (c) Scheme illustrating the arginine-triggered movement of the three dimers preserving the molecular NAGK symmetry, and resulting in an increase in the distance between the substrate sites in each subunit. (d) High and (e) low tilt forms of the NAGK dimers in the respective hexameric NAGKs of *Pseudomonas aeruginosa* (PDB file 2buf) without arginine and in the arginine-containing (arginine molecules are shown in spheres representation) T. maritima enzyme. In both cases two of the three dimers forming the ring are shown, in a
view in which the threefold axis is vertical and in which the molecule is viewed along the twofold molecular axis from the side where two canonical dimers are linked by the N-helices. In each dimer one subunit is grey and the other golden. (f) Sigmoidal arginine inhibition kinetics of *Synechococcus elongatus* NAGK, illustrating the effects of addition of a saturating amount of PII on enzyme activity and on arginine inhibition. The data correspond to our own unpublished results.

Figure 3. Structure of PII to illustrate various conformations of the T-loop. (a) View of the *E. coli* PII protein GlnK containing bound ADP, taken from the complex with AmtB (PDB file 2nuu). The threefold axis is vertical and each subunit is in a different colour. A black dot in the T-loop shows the residue that is uridylylated (b) View as in (a) of *Arabidopsis thaliana* PII bound to MgATP, taken from the complex with NAGK (PDB file 2rd5). A black dot in the T-loop shows the Ser residue corresponding to the Ser that is phosphorylated in cyanobacteria (c) Superimposition of the T-loops of different PII structures. The T-loop in the structures of, (reddish) free PII from *S. elongatus* (PDB file 1qy7), GlnK from *E. coli* (violet) in the complex with AmtB (PDB file 2nuu), GlnK of *Methanococcus jannaschii* with ADP (green, PDB file 2j9d) or with MgATP and 2-oxoglutarate (golden, PDB file 2j9e), and PII of *S. elongatus* (blue, PDB file 2v5h) or *A. thaliana* (cyan, PDB file 2rd5) in their complexes with NAGK. The general orientation, although similar to that in (a), has been modified for optimal viewing of all the superimposed T-loops. (d) The binding of MgATP and 2 oxoglutarate (2OG) in the T-loop of GlnK from *M. jannaschii*. The orientation is the same as in (b).

Figure 4. The PII-NAGK complex. (a) Structure of the complex of *S. elongatus*, viewed along a twofold axis. The threefold axis is vertical. Each PII trimer has a different
shading of yellow, whereas each NAGK dimer is in a different colour. Bound NAG molecules are represented with spheres. Note the high tilt of the dimers, comparable to that in Figure 2e. (b) and (c) Surface representation of the faces that participate in the interaction of, respectively, NAGK and PII, viewed along the threefold axis. Color coding of NAGK is as in (a), and the three subunits of PII are in different shades of yellow. The surfaces for which the accessibility is restricted in the complex are depicted in blue, reddish and deep green, for the T-loop, B-loop and $\beta$1-$\alpha$1 connection of PII and their corresponding complementary surfaces in NAGK. (d) Representation of one NAGK subunit of the *A. thaliana* complex and of the parts of the PII subunit that participate in the interaction with it, to illustrate the relative positions of the active centre (shown in the more closed conformation, and occupied by NAG and ADP), of the arginine site (with bound arginine), and of the PII elements participating in the interactions between both proteins. (e) Detail of the interactions of the T-loop (backbone in yellow) with the NAGK subunit (in thin trace, with some side-chains in grey) of the PII-NAGK complex of *S. elongatus*. Bound NAG and the $\beta$3-$\beta$4 hairpin that is the mobile lid of the NAG site are shown. $\alpha$E and $\alpha$G refer to helices of the $\alpha$4 layer of the NAGK subunit. (f-h) Semitransparent surface representations of the arginine sites in the NAGKs of *T. maritima* (f) and of the PII-NAGK complexes of *A. thaliana* (g) and *S. elongatus* (h). Note the change relative to Tyr15, in the N-helix of the high-affinity arginine site of *T. maritima* NAGK (f), of Phe29 and Tyr23 in (g) and (h), respectively. The change is accompanied by a partial rearrangement of the N-helix.
Figure 1

(a) The biosynthesis of glutamate (Glu) and arginine (Arg) involves several enzymatic steps. 

1. **GS (Glutamine Synthetase)**: Converts glutamate (Glu) to glutamine (Gln) using ATP and ammonia (NH3).
2. **GOGAT (Glutamate Oxoglutarate Aminotransferase)**: Converts glutamate to 2-oxoglutarate (2OG) and glutamine.
3. **Ornithine Transcarbamoylase**: Converts ornithine to citrulline.
4. **Argininosuccinate Synthase**: Converts citrulline and aspartate to argininosuccinate.
5. **Argininosuccinate Lyase**: Converts argininosuccinate to citrulline and fumarate.
6. **Fumarase**: Converts fumarate to succinate.
7. **Guanine Nucleotide Activating Protein (PII)**: Regulates the uptake of ammonia and regulates cytoplasmic pH.

(b) PII interacts with AmtB, a major facilitator superfamily transporter. The T-loop region of AmtB is involved in the interaction with PII.

(c) The T-loop of AmtB is highlighted, showing its role in PII binding.

(d) A closer view of the T-loop region, illustrating the interaction between PII and AmtB.

Figure 1
Figure 2
Figure 3
Figure 4